

Glyphosate Issue Paper: Evaluation of Carcinogenic Potential

**EPA's Office of Pesticide Programs
September 12, 2016**



Table of Contents

List of Acronyms	7
List of Tables	10
1.0 Introduction.....	12
1.1 Background	12
1.2 Evaluation of Carcinogenic Potential.....	12
1.3 Overview of Draft “Framework for Incorporating Human Epidemiologic & Incident Data in Health Risk Assessment”	13
1.4 Summary of the Exposure Profile in the United States.....	15
1.5 Organization of this Document.....	19
2.0 Systematic Review & Data Collection.....	19
2.1 Data Collection: Methods & Sources	20
2.1.1 Open Literature Search.....	20
2.1.2 Studies Submitted to the Agency	21
2.2 Evaluation of Relevant Studies	22
3.0 Data Evaluation of Epidemiology.....	22
3.1 Introduction.....	22
3.2 Considerations for Study Quality Evaluation and Scope of Assessment.....	23
3.2.1 Study Designs.....	24
3.2.1.1 Analytical Studies	25

3.2.1.2	Descriptive Studies	27
3.2.2	Exposure Measures	27
3.2.3	Outcome Measures.....	28
3.2.4	Confounding	28
3.2.5	Statistical Analyses.....	29
3.2.6	Risk of Bias	29
3.3	Review of Quality Results	30
3.3.1	“High” Quality Group	30
3.3.2	“Moderate” Quality Group.....	31
3.3.3	“Low” Quality Group	32
3.4	Assessment of Epidemiological Studies for Relevance to Analysis	44
3.5	Summary of Relevant Epidemiological Studies	44
3.5.1	Solid Tumor Cancer Studies	45
3.5.2	Non-Solid Tumor Cancer Studies.....	53
3.6	Discussion.....	63
4.0	Data Evaluation of Animal Carcinogenicity Studies	68
4.1	Introduction.....	69
4.2	Consideration of Study Quality for Animal Carcinogenicity Studies.....	69
4.3	Assessment of Animal Carcinogenicity Studies	71

4.4	Summary of Animal Carcinogenicity Studies	73
4.5	Rat Carcinogenicity Studies with Glyphosate.....	74
4.5.1	Burnett et al., 1979 (MRID 00105164)	74
4.5.2	Lankas, 1981 (MRID 00093879)	74
4.5.3	Stout and Ruecker, 1990 (MRID 41643801)	75
4.5.4	Atkinson et al., 1993a (MRID 496317023)	79
4.5.5	Brammer, 2001 (MRID 49704601)	79
4.5.6	Pavkov and Wyand 1987 (MRIDs 40214007, 41209905, 41209907)	80
4.5.7	Suresh, 1996 (MRID 49987401)	81
4.5.8	Enemoto, 1997 (MRID 50017103-50017105)	81
4.5.9	Wood et al., 2009a (MRID 49957404).....	81
4.5.10	Summary of Rat Data.....	82
4.6	Mouse Carcinogenicity Studies with Glyphosate.....	85
4.6.1	Reyna and Gordon, 1973 (MRID 00061113)	85
4.6.2	Knezevich and Hogan, 1983 (MRID 00130406).....	85
4.6.3	Atkinson, 1993b (MRID 49631702)	87
4.6.4	Wood et al., 2009b (MRID 49957402)	88
4.6.5	Sugimoto, 1997 (MRID 50017108 - 50017109)	89
4.6.6	Pavkov and Turnier, 1987 (MRIDs 40214006, 41209907).....	90

4.6.7	Summary of Mouse Data	90
4.7	Absorption, Distribution, Metabolism, Excretion (ADME).....	94
4.8	Discussion.....	95
5.0	Data Evaluation of Genetic Toxicity	97
5.1	Introduction.....	97
5.2	Scope of the Assessment Considerations for Study Quality Evaluation.....	97
5.3	Tests for Gene Mutations for Glyphosate Technical	99
5.3.1	Bacterial Mutagenicity Assays	99
5.3.2	<i>In vitro</i> Tests for Gene Mutations in Mammalian Cells	103
5.4	<i>In vitro</i> Tests for Chromosomal Abnormalities.....	105
5.4.1	<i>In vitro</i> Mammalian Chromosomal Aberration Test	105
5.4.2	<i>In vitro</i> Mammalian Micronucleus Test	106
5.5	<i>In Vivo</i> Genetic Toxicology Tests.....	111
5.5.1	<i>In Vivo</i> Assays for Chromosomal Abnormalities	111
5.5.1.1	Mammalian Bone Marrow Chromosomal Aberration Assays.....	111
5.5.1.2	Rodent Dominant Lethal Test	111
5.5.1.3	<i>In Vivo</i> Mammalian Erythrocyte Micronucleus Assays	112
5.6	Additional Genotoxicity Assays Evaluating Primary DNA Damage	119
5.7	Summary and Discussion	126

6.0	Data Integration & Weight-of-Evidence Analysis Across Multiple Lines of Evidence	128
6.1	Background	128
6.2	Dose-Response and Temporal Concordance	128
6.3	Strength, Consistency, and Specificity	130
6.4	Biological Plausibility and Coherence.....	131
6.5	Uncertainty	133
6.6	Evaluation of Cancer Classification per the 2005 EPA Guidelines for Carcinogen Risk Assessment	135
6.6.1	Introduction	135
6.6.2	Discussion of Evidence to Support Cancer Classification Descriptors	137
6.7	Proposed Conclusions Regarding the Carcinogenic Potential of Glyphosate.....	140
7.0	Collaborative Research Plan for Glyphosate and Glyphosate Formulations	141
8.0	References	144

List of Acronyms

ADME: Absorption, Distribution, Metabolism, and Excretion
AHS: Agricultural Health Study
AOP: Adverse Outcome Pathway
AMPA: Aminomethylphosphonic Acid
BrdU: Bromodeoxyuridine
CA: Chromosomal Aberration
CARC: Cancer Assessment Review Committee
CBPI: Cytokinesis Block Proliferation Index
CHL: Chinese Hamster Lung
CHO: Chinese Hamster Ovary
CPRC: Carcinogenicity Peer Review Committee
EFSA: European Food Safety Authority
EPSPS: 5-enolpyruvylshikimate-3-phosphate synthase
FAO: Food and Agriculture Organization
FIFRA: Federal Insecticide, Fungicide, and Rodenticide Act
FISH: Fluorescence *in situ* Hybridization
GC-MS: Gas Chromatography-Mass Spectrometry
HL: Hodgkin Lymphoma
HPLC: High-Performance Liquid Chromatography
HPRT: Hypoxanthine-Guanine Phosphoribosyl Transferase
IARC: International Agency for Research on Cancer
JMPR: Joint Meeting Pesticide Residues
MGUS: Monoclonal Gammopathy of Undetermined Significance
MN: Micronuclei
MOA: Mode of Action
MPCE: Micronucleated Polychromatic Erythrocytes
MRID: Master Record Identifier (a unique number assigned to each study submitted to EPA)
MTD: Maximum Tolerated Dose
NB: Nuclear Bud
NCR: National Research Council
NHL: Non-Hodgkin Lymphoma
NPB: Nucleoplasmic Bridges
NTP: National Toxicology Program
OCSPP: Office of Chemical Safety and Pollution Prevention
OECD: Organization for Economic Cooperation and Development
OPP: Office of Pesticides Program
PCE: Polychromatic Erythrocytes
PK: Pharmacokinetic
PPE: Personal Protective Equipment
PWG: Pathology Work Group
RED: Registration Eligibility Decision
ROS: Reactive Oxygen Species
SAP: Scientific Advisory Panel
SCE: Sister Chromatid Exchange

SCGE: Single Cell Gel Electrophoresis
TAC: Total Antioxidant Capacity
TK: Thymidine Kinase
UDS: Unscheduled DNA Synthesis
USGS: United States Geological Survey
UV: Ultraviolet
WHO: World Health Organization
XPRT: Xanthine-Guanine Phosphoribosyl Transferase

List of Figures

Figure 1.1. Source to outcome pathway (adapted from NRC, 2007)

Figure 1.2. Glyphosate agricultural usage (pounds applied annually) from 1987- 2014. Boxes indicate years when glyphosate-resistant crops were introduced. Source: Proprietary Market Research Data (1987 – 2014)

Figure 1.3. Map of estimated agricultural use for glyphosate in 1994 from USGS

Figure 1.4. Map of estimated agricultural use for glyphosate in 2014 from USGS

Figure 3.1. Study evaluation process for epidemiological studies

Figure 3.2. Forest plot of effect estimates (denoted as ES for effect sizes) and associated 95% confidence intervals (CI) for non-Hodgkin lymphoma (NHL)

Figure 7.1. Results of HepG2 exposures following 24 hour incubation using different glyphosate formulations

List of Tables

Table 3.1 Study Quality Considerations

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking

Table 3.3. Summary of Findings: Solid Tumor Cancer Studies

Table 3.4. Summary of Findings: Non-Solid Tumor Cancer Studies

Table 4.1. Testicular Interstitial Cell Tumors in Male Sprague-Dawley Rats (Lankas, 1981) Cochran-Armitage Trend Test & Fisher's Exact Test Results

Table 4.2. Pancreatic Islet Cell Tumors in Male Sprague-Dawley Rats (Stout and Ruecker, 1990) Cochran-Armitage Trend Test & Fisher's Exact Test Results

Table 4.3. Historical Control Data — Pancreatic Islet Cell Adenomas in Male Sprague- Dawley Rats (MRID No. 41728701)

Table 4.4. Hepatocellular Tumors in Male Sprague-Dawley Rats (Stout and Ruecker, 1990) Cochran-Armitage Trend Test & Fisher's Exact Test Results

Table 4.6. Thyroid C-Cell Tumors in Male Sprague-Dawley Rats (Stout and Ruecker, 1990) Cochran-Armitage Trend Test & Fisher's Exact Test Results

Table 4.7. Thyroid C-Cell Tumors in Female Sprague Dawley Rats Cochran-Armitage Trend Test & Fisher's Exact Test Results (Stout and Ruecker, 1990)

Table 4.8. Thyroid Non-Neoplastic Lesions (Stout and Ruecker, 1990)

Table 4.9. Liver Adenomas in Male Wistar Rats (Brammer, 2001) Cochran-Armitage Trend Test and Fisher's Exact Test Results

Table 4.10. Mammary Gland Tumor Incidences in Female Rats (Wood et al., 2009a) Fisher's Exact Test and Cochran-Armitage Trend Test Results

Table 4.11. Summary of Rat Carcinogenicity Studies

Table 4.12. Kidney Tubular Cell Tumors in Male CD-1 Mice (Knezevich and Hogan, 1983) Cochran-Armitage Trend Test & Fisher's Exact Test Results

Table 4.13. Kidney Histopathological Alterations in Male CD-1 Mice (Knezevich and Hogan, 1983)

Table 4.14. Hemangiosarcomas in Male CD-1 Mice (Atkinson, 1993b) Cochran-Armitage Trend Test and Fisher's Exact Test Results

Table 4.15. Lung Tumors in Male CD-1 Mice (Wood et al., 2009b) Fisher's Exact Test and Cochran-Armitage Trend Test Results

Table 4.16. Malignant Lymphomas in Male CD-1 Mice (Wood et al., 2009b) Fisher's Exact Test and Cochran-Armitage Trend Test Results

Table 4.17. Hemangioma Incidences (Sugimoto, 1997) Fisher's Exact Test and Cochran-Armitage Trend Test Results

Table 4.18. Summary of Mouse Carcinogenicity Studies

Table 5.1. *In vitro* Test for Gene Mutations in Bacteria: Glyphosate Technical

Table 5.2. *In vitro* Mammalian Gene Mutation Assays: Glyphosate Technical

Table 5.3. *In vitro* Tests for Chromosome Aberrations in Mammalian Cells- Glyphosate Technical

Table 5.4. *In vitro* Tests for Micronuclei Induction in Mammalian Cells- Glyphosate Technical

Table 5.5. *In Vivo* Tests for Chromosomal Aberrations in Mammals- Glyphosate Technical

Table 5.6. *In Vivo* Tests for Micronuclei Induction in Mammals- Glyphosate Technical

Table 5.7 Assays for Detecting Primary DNA Damage- Glyphosate Technical

1.0 Introduction

1.1 Background

Glyphosate is a non-selective, phosphonomethyl amino acid herbicide registered to control weeds in various agricultural and non-agricultural settings. The herbicide acts by inhibiting the 5-enolpyruvylshikimate-3-phosphate synthase (EPSPS) enzyme, which is not present in mammalian systems. Glyphosate was initially registered in 1974. Since then, several human health analyses have been completed for glyphosate. In 1986, EPA issued the Glyphosate Registration Standard which updated the agency's toxicity database for this compound. In 1993, EPA issued the registration eligibility decision (RED) that indicated that glyphosate was eligible for re-registration.

Currently, glyphosate is undergoing Registration Review¹, a program where all registered pesticides are reviewed at least every 15 years as mandated by the Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA). The initial docket opening for glyphosate occurred in 2009 with the publication of the human health scoping document and preliminary work plan². As part of this process, the hazard and exposure of glyphosate are reevaluated to determine its potential risk to human and environmental health. Risks are assessed using current practices and policies to ensure pesticide products can still be used safely. Registration Review also allows the agency to incorporate new science. For human health risk assessment, both non-cancer and cancer effects are evaluated for glyphosate and its metabolites, aminomethylphosphonic acid (AMPA) and *N*-acetyl-glyphosate; however, this document will focus on the cancer effects only. EPA expects to complete its complete human health risk assessment in 2017 that will include an assessment of risk from anticipated exposures resulting from registered uses of glyphosate in residential and occupational settings.

1.2 Evaluation of Carcinogenic Potential

Since its registration, the carcinogenic potential of glyphosate has been evaluated by EPA several times. In 1985, the initial peer review of glyphosate was conducted in accordance with the Proposed Guidelines for Carcinogen Risk Assessment. The agency classified glyphosate as a Group C chemical (Possible Human Carcinogen), based on the presence of kidney tumors in male mice. In 1986, the agency requested that the FIFRA Scientific Advisory Panel (SAP) evaluate the carcinogenic potential of glyphosate. The panel determined that the data on renal tumors in male mice were equivocal (only an increase in adenomas was observed and the increase did not reach statistical significance). As a result, the panel recommended a Group D chemical classification (Not Classifiable as to Human Carcinogenicity) for glyphosate and advised the agency to issue a data call-in notice for further studies in rats and/or mice to clarify the unresolved questions (FIFRA SAP Report, 1986)³.

¹ Additional information on the Registration Review process can be found at: <https://www.epa.gov/pesticide-reevaluation/registration-review-process>

² Documents of the Registration Review can be found in the public docket at: EPA-HQ-OPP-2009-0361, accessible at www.regulations.gov.

³ Review available at: http://www.epa.gov/pesticides/chem_search/cleared_reviews/csr_PC-103601_24-Feb-86_209.pdf

With the submission of two rat carcinogenicity studies following this data call-in, a second peer review was conducted in 1991 by the agency's Carcinogenicity Peer Review Committee (CPRC) to incorporate the new data. In accordance with the agency's 1986 Draft Guidelines for Carcinogen Risk Assessment, the CPRC classified glyphosate as a Group E Chemical: "Evidence of Non-Carcinogenicity for Humans" based upon lack of evidence for carcinogenicity in mice and rats and the lack of concern for mutagenicity (TXR# 0008897).

Most recently, in September 2015, a third review was done by the Cancer Assessment Review Committee (CARC). Relevant glyphosate data available to EPA at that time for glyphosate were reevaluated, including studies submitted by the registrant and studies published in the open literature. The agency performed this evaluation in support of Registration Review in accordance with the 2005 Guidelines for Carcinogen Risk Assessment, classified glyphosate as "Not Likely to be Carcinogenic to Humans" (CARC, 2015; TXR #0057299).

Recently, several international agencies have evaluated the carcinogenic potential of glyphosate. In March 2015, the International Agency for Research on Cancer (IARC), a subdivision of the World Health Organization (WHO), determined that glyphosate was a probable carcinogen (group 2A) (IARC, 2015). Later, in November 2015, the European Food Safety Authority (EFSA) determined that glyphosate was unlikely to pose a carcinogenic hazard to humans (EFSA, 2015). In May 2016, the Joint Food and Agriculture Organization (FAO)/WHO Meeting on Pesticide Residues (JMPR), another subdivision of the WHO, concluded that glyphosate was unlikely to pose a carcinogenic risk to humans from exposure through the diet (JMPR, 2016). Some individual countries (e.g., France, Sweden) have been moving to ban glyphosate based on the IARC decision, while other countries (e.g., Japan, Canada) have continued to support their conclusion that glyphosate is unlikely to pose a carcinogenic risk to humans.

The recent peer review performed by CARC served as an initial analysis to update the data evaluation for glyphosate at that time. Based on an evaluation of the studies included in the recent analyses by IARC, JMPR, and EFSA, the agency then became aware of additional relevant studies not available to EPA. As a result, EPA also requested information from registrants about studies that existed, but had never been submitted to the agency. The current evaluation incorporates these additional studies. In addition, the Agency conducted a systematic review of the open literature and toxicological databases for glyphosate by using a draft "Framework for Incorporating Human Epidemiologic & Incident Data in Health Risk Assessment". As such, the current evaluation also provides a more thorough evaluation than the 2015 CARC review.

1.3 Overview of Draft "Framework for Incorporating Human Epidemiologic & Incident Data in Health Risk Assessment"

In 2010, OPP developed a draft "Framework for Incorporating Human Epidemiologic & Incident Data in Health Risk Assessment" which provides the foundation for evaluating multiple lines of scientific evidence in the context of understanding of the mode of action (MOA)/adverse outcome pathway (AOP) (U.S. EPA, 2010). The draft framework, which includes two key components, problem formulation and use of the MOA/AOP pathway frameworks, was reviewed

favorably by the FIFRA SAP in 2010 (FIFRA SAP, 2010). Recently, EPA has applied this framework to the evaluation of atrazine and chlorpyrifos⁴.

OPP's draft framework is consistent with updates to the World Health Organization/International Programme on Chemical Safety MOA/human relevance framework, which highlights the importance of problem formulation and the need to integrate information at different levels of biological organization (Meek *et al.*, 2014). Consistent with recommendations by the National Research Council (NRC) in its 2009 report on *Science and Decisions*, OPP's draft framework describes the importance of using problem formulation at the beginning of a complex scientific analysis. The problem formulation stage starts with planning dialogue with risk managers to identify goals for the analysis and possible risk management strategies. This initial dialogue provides the regulatory context for the scientific analysis and helps define the scope of such an analysis. The problem formulation stage also involves consideration of the available information regarding the pesticide use/usage, toxicological effects of concern, and exposure pathways and duration along with key gaps in data or scientific information. Specific to glyphosate, the scoping document prepared for Registration Review (J. Langsdale *et al.*, 2009) along with the review conducted by the CARC (CARC, 2015) represent the problem formulation analyses for the weight-of-evidence evaluation for carcinogenic potential. A summary of the US exposure profile is provided in Section 1.4 below to provide context for interpreting the various lines of evidence.

One of the key components of the agency's draft framework is the use of the MOA framework/AOP concept (Figure 1.1) as a tool for organizing and integrating information from different sources to inform the causal nature of links observed in both experimental and observational studies. Specifically, the modified Bradford Hill Criteria (Hill, 1965) are used to evaluate strength, consistency, dose response, temporal concordance and biological plausibility of multiple lines of evidence in a weight-of-evidence analysis.

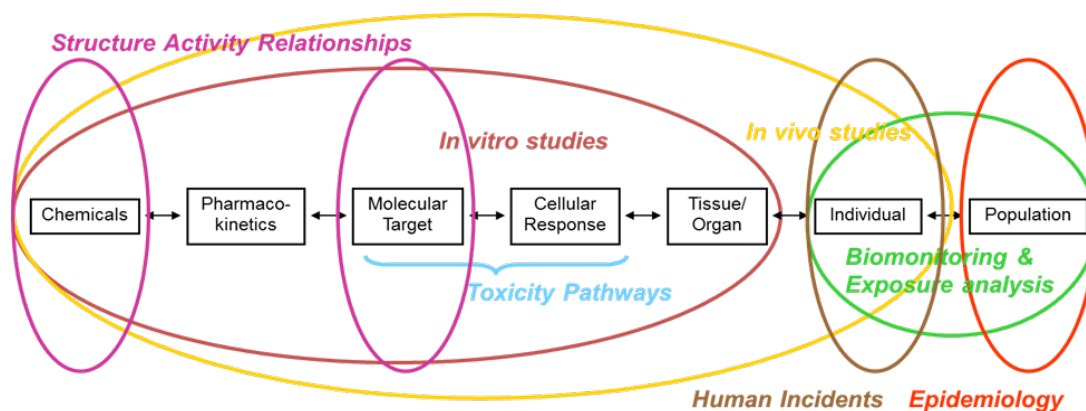


Figure 1.1. Source to outcome pathway (adapted from NRC, 2007).

⁴ Chlorpyrifos Revised Human Health Risk Assessment for Registration Review; 29-DEC-2014; D424485. U.S. EPA 2010 SAP Background White Paper – Re-evaluation of Human Health Effects of Atrazine: Review of Experimental Animal and In Vitro Studies and Drinking Water Monitoring Frequency. EPA-HQ-OPP-2010-0125. U.S. EPA 2011 SAP Issue Paper – Re-evaluation of Human Health Effects of Atrazine: Review of Cancer Epidemiology, Non-cancer Experimental Animal and In Vitro Studies and Drinking Water Monitoring Frequency. EPA-HQ-OPP-2011-0399.

1.4 Summary of the Exposure Profile in the United States

All pesticide products provide critical information about how to safely and legally handle and use pesticide products. Pesticide labels are legally enforceable and all carry the statement “it is a violation of Federal law to use this product in a manner inconsistent with its labeling.” In other words, the label is law. As a result, a key function of the pesticide product label is to manage the potential risk from pesticides.

Labeled uses of glyphosate include over 100 terrestrial food crops as well as other non-agricultural sites, such as greenhouses, aquatic areas, and residential areas. It is also registered for use on glyphosate-resistant (transgenic) crop varieties such as corn, soybean, canola, cotton, sugar beets, and wheat. Registered tolerances in the United States include residues of the parent compound glyphosate and *N*-acetyl-glyphosate, a metabolite found in/on glyphosate-tolerant crops⁵.

Dietary (food and water) exposures are anticipated from applications to crops. Since there are registered uses of glyphosate that may be used in residential settings, residential handlers may be exposed to glyphosate during applications. Exposures may also occur from entering non-occupational areas that have been previously treated with glyphosate. Occupational/commercial workers may be exposed to glyphosate while handling the pesticide prior to application (mixing and/or loading), during application, or when entering treated sites. The agency considers all of the anticipated exposure pathways as part of their evaluation for human health.

Oral exposure is considered the primary route of concern for glyphosate. Oral absorption has been shown to be relatively low for glyphosate (~30% of administered doses) with negligible accumulation in tissues and rapid excretion (primarily unchanged parent) via the urine. Due to its low vapor pressure, inhalation exposure to glyphosate is expected to be minimal. Dermal penetration has also been shown to be relatively low for human skin (<1%) indicating dermal exposure will only contribute slightly to a systemic biological dose. Furthermore, in route-specific inhalation and dermal toxicity studies, no adverse effects were observed. This all suggests that there is low potential for a sustainable biological dose following glyphosate exposure.

In residential/non-occupational settings, children 1-2 years old are considered the most highly exposed subpopulation with oral exposures from dietary (food and water) ingestion and incidental oral ingestion (e.g., hand-to-mouth activities) in treated areas. There is also potential for dermal exposures in previously treated areas. Using HED’s standard exposure assessment methodologies which are based on peer-reviewed and validated exposure data and models⁶, a high-end estimate of combined exposure for children 1-2 years old is 0.47 mg/kg/day (see Appendix E).

⁵ All currently registered tolerances for residues of glyphosate can be found in the Code of Federal Regulations (40 CFR §180.364).

⁶ Available: <http://www2.epa.gov/pesticide-science-and-assessing-pesticide-risks/standard-operating-procedures-residential-pesticide>

At the time of initial registration (1974), total use of glyphosate in the United States was approximately 1.4 million pounds (Benbrook, 2016). In 1995, total use of glyphosate increased to approximately 40 million pounds with agriculture accounting for 70% of use. With the introduction of transgenic crop varieties in the United States circa 1996, (such as soybean, cotton, and corn) use of glyphosate increased dramatically (Green and Owen, 2011), and in 2000 the total use of glyphosate in the United States was approximately 98.5 million pounds. By 2014, total annual use of glyphosate was approximately 280-290 million pounds (based on Benbrook, 2016 and industry proprietary data accessible to EPA) with agriculture accounting for 90% of use. Although glyphosate use has continuously increased up to 2012, the stabilization of glyphosate usage in recent years is due to the increase in a number of glyphosate-resistant weed species, starting with rigid ryegrass identified in California in 1998 and currently totaling 16 different weed species in the United States as of March 2016. Figure 1.2 below provides a visual representation of the increased agricultural use of glyphosate in the United States using proprietary market research data from 1987-2014.

The increased use of glyphosate may be partly attributed to an increase in the number of farmers using glyphosate; however, it is more likely that individuals already using glyphosate increased their use and subsequent exposure. With the introduction of transgenic crop varieties, glyphosate use shifted from pre-emergent to a combination of pre- and post-emergent applications. Additionally, application rates increased in some instances and more applications were allowed per year (2-3 times/year). Maps from the United States Geological Survey (USGS) displaying glyphosate use in the United States indicate that although use has drastically increased since 1994, areas treated with glyphosate for agricultural purposes appear to be approximately the same over time (Figures 1.3-1.4). The introduction of transgenic crops in some cases led to a shift in crops grown on individual farms, such that more acreage within the farm would be dedicated to growing the glyphosate-tolerant crops replacing other crops. In addition, during the 2000s there was also an increase in growing corn for ethanol production, which could also have resulted in increased acreage dedicated glyphosate-tolerant corn.

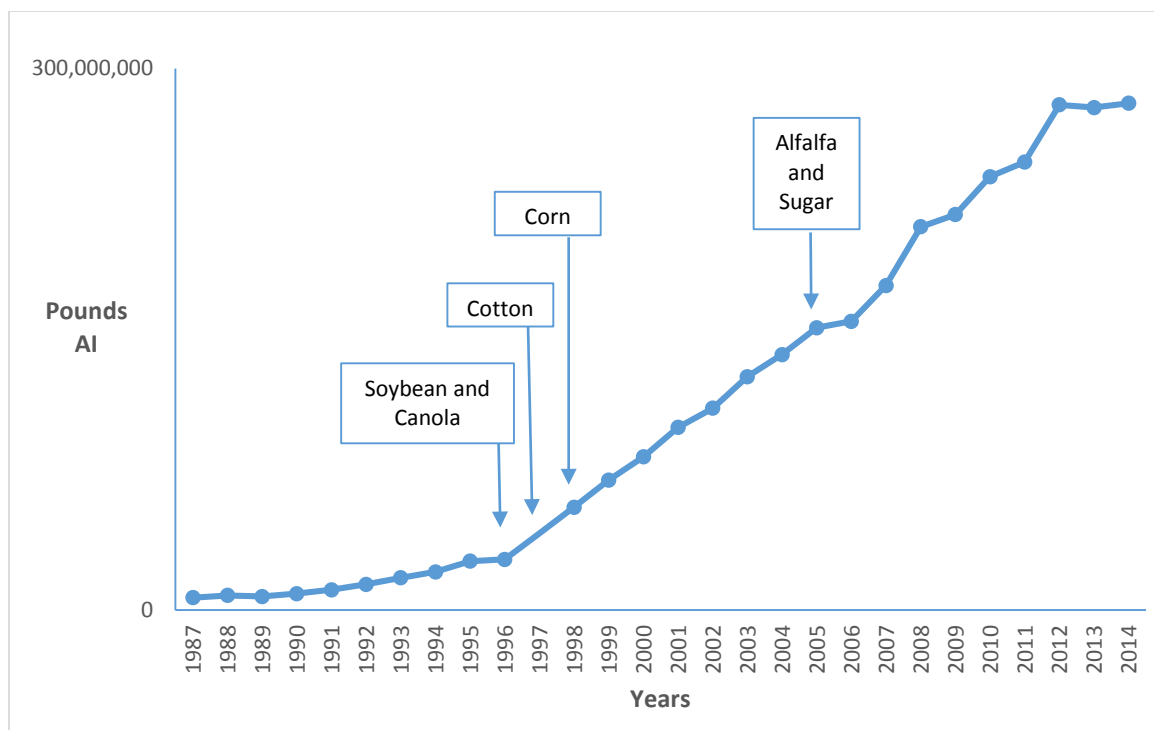


Figure 1.2. Glyphosate agricultural usage (pounds applied annually) from 1987- 2014. Boxes indicate years when glyphosate-resistant crops were introduced. Source: Proprietary Market Research Data (1987 – 2014).

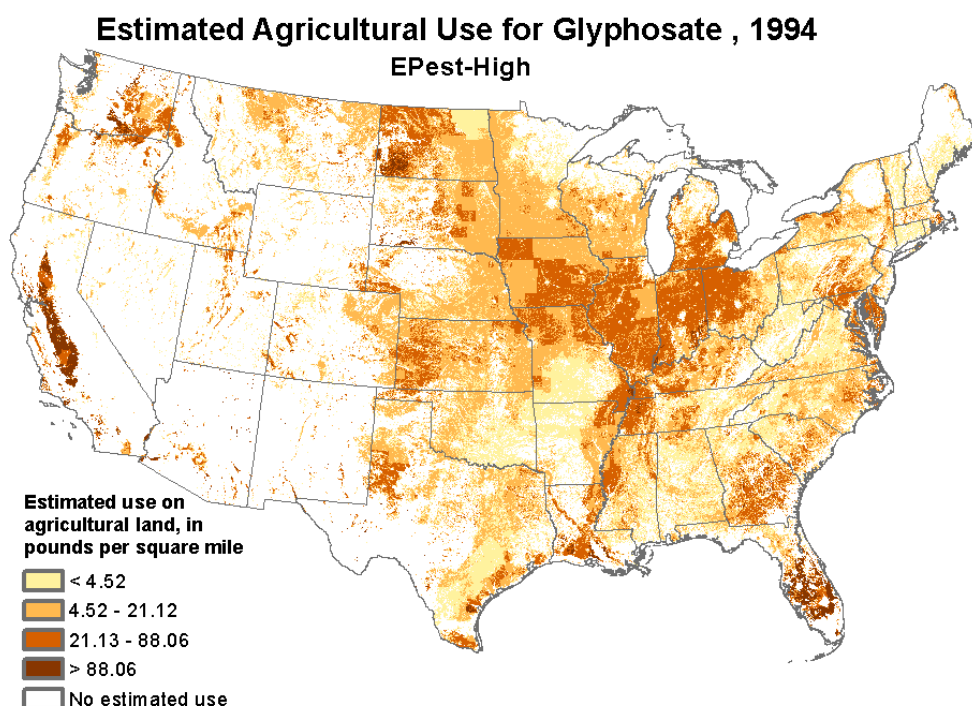


Figure 1.3. Map of estimated agricultural use for glyphosate in 1994 from USGS (http://water.usgs.gov/nawqa/pnsp/usage/maps/show_map.php?year=1994&map=GLYPHOSATE&hilo=H)

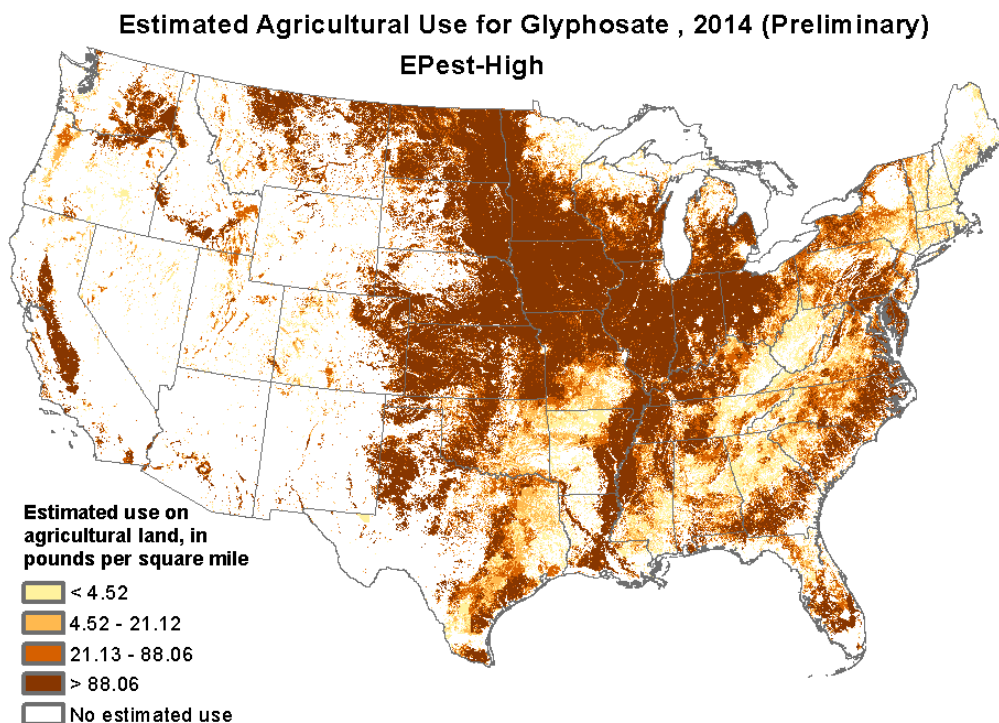


Figure 1.4. Map of estimated agricultural use for glyphosate in 2014 from USGS
(http://water.usgs.gov/nawqa/pnsp/usage/maps/show_map.php?year=2014&map=GLYPHOSATE&hilo=H)

The potential exposure to occupational handlers is dependent on the formulation, specific task (mixer, loader, and/or applicator), rate of application, and acreage treated. Using HED's standard occupational exposure assessment methodologies which are based on peer-reviewed and validated exposure data and models⁷, mixer/loaders result in the highest potential exposure estimates. Assuming no personal protective equipment (PPE), exposure estimates for mixer/loaders range from 0.03-7 mg/kg/day using the maximum application rate for high acreage agricultural crops (6 lb ai/acre)⁸. For applicators, exposure would be lower with estimates ranging from 0.02-0.03 mg/kg/day using the same application rate and acreage.

The maximum potential exposures from currently registered uses of glyphosate in residential and occupational settings in the United States are used in the current evaluation to aid in the determination of whether findings in laboratory studies are relevant for human health risk assessment. In Sections 4.0 and 5.0, descriptions are provided for animal carcinogenicity and genotoxicity studies, respectively. Results from these studies, particularly those administering high doses, are put into context with the human exposure potential in the United States.

⁷ Available: <https://www.epa.gov/pesticide-science-and-assessing-pesticide-risks/occupational-pesticide-handler-exposure-data>

⁸ Based on use information provided by the Joint Glyphosate Task Force for the following end-use products: EPA Registration Nos.: 100-1182, 228-713, 524-343, 524-475, 524-537, 524-549, 524-579, 4787-23, and 62719-556.

1.5 Organization of this Document

In this analysis of the human carcinogenic potential of the active ingredient glyphosate, the agency has performed a comprehensive analysis of available data from submitted guideline studies and the open literature. This includes epidemiological, animal carcinogenicity, and genotoxicity studies. Consistent with the 2010 draft framework, the agency has evaluated these multiple lines of evidence and conducted a weight-of-evidence analysis. Although there are studies available on glyphosate-based pesticide formulations, the agency is soliciting advice from the FIFRA SAP on this evaluation of human carcinogenic potential for the active ingredient glyphosate only at this time. The remainder of this document is organized by the following:

- *Section 2.0 Systematic Review & Data Collection Methods* provides a description of methods used to compile all relevant studies used in the current evaluation.
- *Section 3.0 Data Evaluation of Epidemiology* describes the available epidemiological studies, evaluates relevant studies for study quality, and discusses reported effect estimates.
- *Section 4.0 Data Evaluation of Animal Carcinogenicity Studies* provides a description and evaluation of the available animal carcinogenicity studies for glyphosate.
- *Section 5.0 Data Evaluation of Genetic Toxicity* summarizes and discusses the various genotoxicity assays that have been tested with glyphosate.
- *Section 6.0 Data Integration & Weight of Evidence Analysis Across Multiple Lines of Evidence* integrates available data discussed in Sections 3.0-5.0 to consider concepts, such as strength, consistency, dose response, temporal concordance and biological plausibility in a weight-of-evidence analysis. This section also provides discussion of the data in the context of cancer descriptors provided in the 2005 Guidelines for Carcinogen Risk Assessment.
- *Section 7.0 Collaborative Research Plan for Glyphosate and Glyphosate Formulations* provides a discussion of planned research that is intended to evaluate the role of glyphosate in product formulations and the differences in formulation toxicity.

2.0 Systematic Review & Data Collection

In recent years, the National Academy of Sciences National Research Council (NRC) has encouraged the agency to move towards systematic review processes to enhance the transparency of scientific literature reviews that support chemical-specific risk assessments to inform regulatory decision making (NRC, 2011). The NRC defines systematic review as “a scientific investigation that focuses on a specific question and uses explicit, pre-specified scientific methods to identify, select, assess, and summarize the findings of similar but separate studies” (NRC, 2014). Consistent with NRC’s recommendations, EPA’s Office of Chemical Safety and Pollution Prevention (OCSPP) is currently developing systematic review policies and procedures. In short, OCSPP employs “fit for purpose” systematic reviews that rely on standard methods for **collecting, evaluating, and integrating** the scientific data supporting the agency’s decisions. The concept of fit for purpose implies that a particular activity or method is suitable for its intended use. Inherent in this definition is the concept that one size does not fit all situations and thus flexibility is allowed. However, it is notable that with flexibility comes the

importance of transparency of documented processes; including the importance of transparency and clarity in approaches to data collection, evaluation, and integration. These are described throughout the document with data collection in Sections 2.1.1-2.1.2, evaluation in Sections 3-5, and integration in Section 6.

As a result, more recent evaluations are starting to reflect this progression in the agency's process. Similar to the draft framework for incorporating human epidemiologic and incident data, systematic review begins with a problem formulation to determine the scope and purpose of the search. Studies are considered based on their relevance to answer specific questions and those studies deemed relevant are then further considered for their usefulness in risk assessment.

The agency strives to use high-quality studies when evaluating the hazard potential of pesticidal chemicals and considers a broad set of data during this process. This includes registrant generated studies required under FIFRA, as well as peer-reviewed scientific journals and other sources, such as other governments and academia. A wide range of potential adverse effects are assessed using acute, subchronic, chronic, and route-specific studies; predominately from studies with laboratory animals, in addition to epidemiologic and human incident data. All studies are thoroughly reviewed to ensure appropriate conduct and methodologies are utilized, and that sufficient data and details are provided. In this way, hazards are identified and potential risks characterized to ensure that decisions are informed by the best science available.

2.1 Data Collection: Methods & Sources

Data were collected by searching the open literature (Section 2.1.1) and other publicly available sources (e.g., recent internal reviews, evaluations by other organizations) (Section 2.1.2). Internal databases were also searched for submitted studies conducted according to Organization for Economic Cooperation and Development (OECD) test guidelines, OCSPP harmonized test guidelines, and other pesticide test guidelines (OPP guidelines) (Section 2.1.2).

It should be noted that glyphosate is primarily manufactured as various salts with cations, such as isopropylamine, ammonium, or sodium. These salts are derivatives of the active substance glyphosate and increase the solubility of technical-grade glyphosate acid in water. All of these forms were considered for the current evaluation.

2.1.1 Open Literature Search

As part of the evaluation of the human carcinogenic potential of glyphosate, the literature review described here uses concepts consistent with fit for purpose systematic review, such as detailed tracking of search terms and which literature have been included or excluded. The primary goal of the literature search was to identify relevant and appropriate open literature studies that had the potential to inform the agency on the human carcinogenic potential of glyphosate. Therefore, non-mammalian studies were not considered, and several terms were used in the search string in an attempt to exclude non-mammalian studies.

To obtain literature studies, OPP worked with EPA librarians to conduct searches in PubMed, Web of Science, and Science Direct. A search was conducted on May 6, 2016 in PubMed and Web of Science using the following search string to yield 141 and 225 results, respectively:

((glyphosate OR "1071-83-6" OR roundup OR "N-(Phosphonomethyl)glycine") AND (aneuploid* OR chromosom* OR clastogenic* OR "DNA damag*" OR "DNA adduct*" OR genome* OR genotoxic* OR micronucle* OR cancer* OR carcinogen* OR oncogenic* OR mutagen* OR cytotoxic* OR tumor* OR tumour* OR malignanc* OR neoplasm* OR *oma)) NOT (fish* OR frog* OR tadpole* OR insect* OR eco* OR amphibian* OR reptil* OR invertebrate* OR fly OR flies OR aquatic OR bird* OR aqueous OR water OR yeast* OR worm* OR earthworm* OR bacteria* OR lichen OR resist* OR "herbicide resist")

Due to differences with using Science Direct, the search string was slightly changed. This search was also conducted on May 6, 2016 and yielded 459 results:

((glyphosate OR "1071-83-6" OR roundup OR "N-(Phosphonomethyl)glycine") AND (aneuploid* OR chromosom* OR clastogenic* OR (DNA pre/2 (damag* OR adduct*)) OR genome* OR genotoxic* OR micronucle* OR cancer* OR carcinogen* OR oncogenic* OR mutagen* OR cytotoxic* OR tumor* OR tumour* OR malignanc* OR neoplasm* OR *oma)) AND NOT (eco* OR fish* OR frog* OR tadpole* OR invertebrate* OR bird* OR insect* OR fly OR flies OR amphibian* OR reptil* OR yeast* OR aquatic OR aqueous OR water OR worm* OR earthworm* OR bacteria* OR lichen OR resist* OR "herbicide resist")

After cross-referencing the results obtained from the three open literature searches for duplicates, a total of 735 individual articles were obtained (Appendix A) and one additional study (Alvarez-Moya et al., 2014) not identified in the search was added to this list for a total of 736 individual articles. All of the studies were evaluated to determine if the study would be considered relevant to the issue of concern (i.e., human carcinogenic potential of glyphosate). Many of the articles were not considered to be within the scope of the search or not considered relevant in general (658 articles). Additionally, 27 articles were not appropriate due to the type of article (i.e., correspondence, abstract only, not available in English, retraction). Of the 51 relevant articles, 42 were used in the current evaluation (31 genotoxicity, 9 epidemiological, and 2 animal carcinogenicity). Three articles also reported on the potential of glyphosate and its metabolites to be developed into therapeutic drugs for cancer treatment. The remaining 6 articles evaluated effects on glyphosate or glyphosate formulations on cellular processes, mostly focusing on epidermal cells, and were not considered informative for the current evaluation.

2.1.2 Studies Submitted to the Agency

For all pesticides, there are toxicology data requirements that must be submitted to the agency for registration. These studies, defined under the 40 CFR Part 158 Toxicology Data Requirements, provide information on a wide range of adverse health outcomes, routes of exposure, exposure durations, species, and lifestages. They typically follow OECD, OCSPP, or OPP accepted protocols and guidelines, which ease comparisons across studies and chemicals. The toxicological databases for glyphosate⁹ were reviewed and all relevant animal, genotoxicity, and metabolism studies were collected for consideration.

⁹ Glyphosate pesticide chemical (PC) codes: 103601, 103603, 103604, 103605, 103607, 103608, 103613, 128501, and 417300.

Several resources were used to ensure all relevant studies were included in the current evaluation. The list of studies obtained from the toxicological database and the open literature search were cross-referenced with recent internal reviews (CARC, 2015; S. Recore *et al.*, 2014). This list was also cross-referenced with review articles from the open literature [Chang and Delzell (2016), Greim *et al.* (2015), Kier and Kirkland (2013), Kier (2015), Mink *et al.* (2012), Schinasi and Leon (2014), and Williams *et al.* (2000)]¹⁰. EPA requested studies from registrants that were not previously available to the EPA. As a result, numerous studies were subsequently submitted to the agency. Study reports for one animal carcinogenicity study and 17 genotoxicity studies, were not available to the agency and have been noted in the relevant sections below. For these studies, data and study summaries provided in Greim *et al.* (2015) and Kier and Kirkland (2013) were relied upon for the current evaluation.

2.2 Evaluation of Relevant Studies

Studies submitted to the agency are evaluating based on OECD, OCSPP, or OPP test guideline requirements to determine whether studies are acceptable for use in risk assessment. In the current evaluation, animal carcinogenicity, genotoxicity, and metabolism studies located in the internal databases with access to full study reports were evaluated in this manner. Those classified as unacceptable were noted and subsequently excluded from the current evaluation.

In order to evaluate open literature studies, criteria described in the OPP guidance for considering and using open literature toxicity studies to support human health risk assessment was utilized (U.S. EPA, 2012). This guidance assists OPP scientists in their judgement of the scientific quality of open literature publications. More specifically, the document discusses how to screen open literature studies for journal articles/publications that are relevant to risk assessment, how to review potentially useful journal articles/publications and categorize them as to their usefulness in risk assessment, and how the studies may be used in the risk assessment. As with submitted studies, those deemed unacceptable were noted and subsequently excluded from the current evaluation.

3.0 Data Evaluation of Epidemiology

3.1 Introduction

Epidemiological studies are valuable for risk assessment since they may provide direct evidence on whether human exposure to a chemical may cause cancer. Studies of high quality and adequate statistical power are preferable and remove the need to account for extrapolation from animals to humans or extrapolation from high to low doses. Epidemiological studies can also be integrated with experimental evidence when determining or clarifying the carcinogenic potential of a chemical for risk assessment. The key considerations in evaluating epidemiologic studies are study design, exposure assessment, outcome assessment, confounding control, statistical analyses, and risk of other bias.

¹⁰ All review articles, except Schinasi and Leon (2014), were funded and/or linked to Monsanto Co. or other registrants.

OPP routinely evaluates the available epidemiological literature. As part of Registration Review of glyphosate, an evaluation was initially conducted in 2014 (S. Recore *et al.*, 2014) and subsequently another evaluation was performed in 2015 (CARC, 2015). The 2015 evaluation began with the epidemiological studies previously identified in the 2014 evaluation and included three additional studies that were not included in the 2014 evaluation. These studies were identified in review articles, included in the evaluation by IARC (2015), or were published since the 2014 OPP evaluation. Both the 2014 and 2015 OPP evaluations considered the design and overall quality of the epidemiological studies; however, formal study quality evaluations and rankings were not conducted. In the current review, all of the studies in the 2015 report, as well as additional epidemiological articles identified from a comprehensive search and cross-referencing with available resources as described under Section 2.0, were considered in the current evaluation, which totaled 58 epidemiological studies. The following sections provide a description of how epidemiological studies were evaluated for study quality and subsequent overall rankings, a summary of relevant studies, and a discussion of the overall results.

3.2 Considerations for Study Quality Evaluation and Scope of Assessment

This section summarizes how specific study characteristics were factored into the determination of a study's overall quality category. It should be noted that these study quality considerations are specific to the issue of concern (i.e., carcinogenic potential of glyphosate). These considerations are considered 'fit-for-purpose' under this context and could differ in another regulatory or scientific context. Although the basic concepts apply broadly, the study quality considerations are tailored specifically to studies investigating the association between glyphosate exposure and cancer outcomes. As with all research studies, the design elements of an epidemiological study have potential impacts on study quality and relevance to the research question under investigation. Each study was, therefore, judged to be of high, moderate, or low quality in each of the following six domains affecting study quality: study design, exposure assessment, outcome assessment, confounder control, statistical analysis, and susceptibility to bias (See Section 3.2.1 for general considerations under each domain). A similar approach was recently used by OPP for the evaluation of epidemiological studies for organophosphate pesticides (A. Lowit *et al.*, 2015).

Primary literature and associated meta-analyses evaluating the association between glyphosate exposure and a cancer outcome were the focus of this analysis. Reviews were only used to identify individual studies that should be considered for study evaluation. Commentaries, correspondence, and letters to the editor without original data were excluded. Of the relevant studies identified, studies with the most complete analyses utilizing the greatest number of cases and controls (e.g., pooled case-control studies) were evaluated for ranking (see Appendix B for visual representation of these studies). If studies did not collect exposure information on glyphosate from individual subjects, did not assess an outcome (e.g., biomonitoring studies), and/or did not provide a quantitative measure of an association between glyphosate and a cancer outcome, then these studies were assigned a low quality ranking and were not further evaluated in detail (see Figure 3.1). A similar process was used by JMPR for their identification of epidemiological studies for evaluating the carcinogenic potential of glyphosate and two other pesticides (JMPR, 2016).

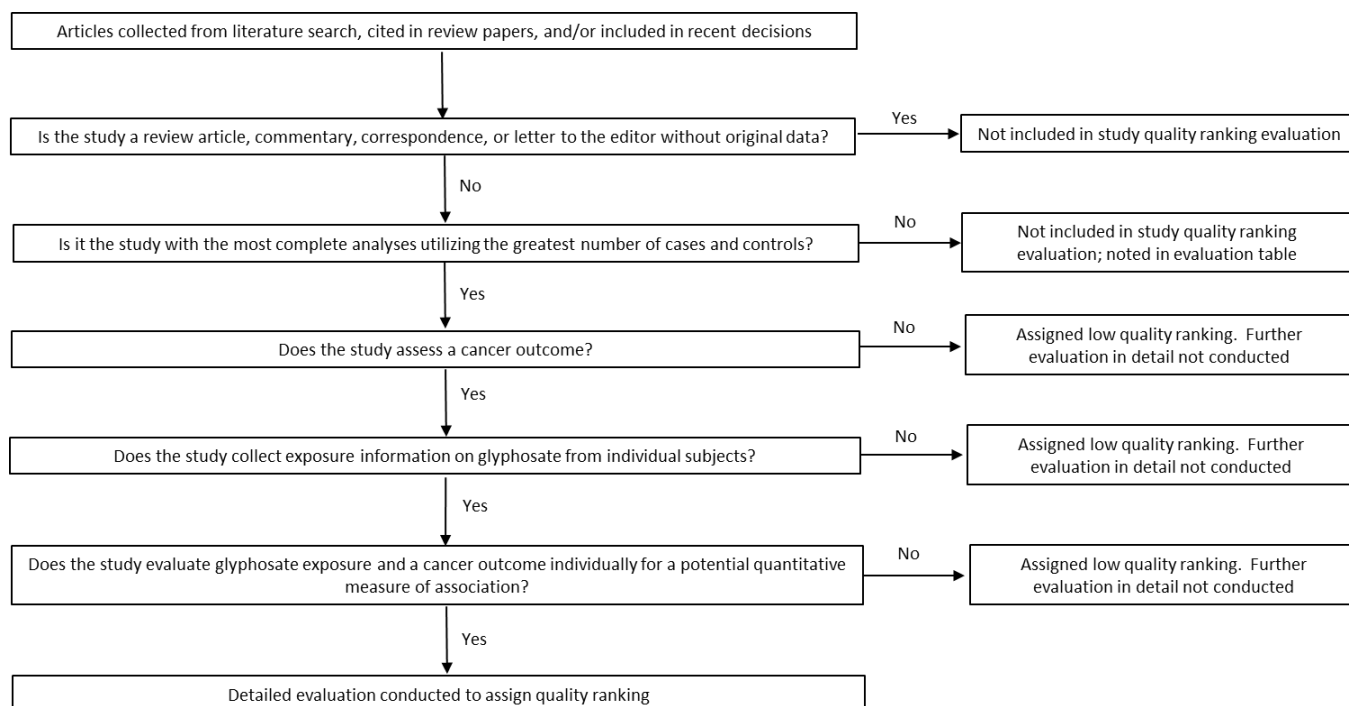


Figure 3.1. Study evaluation process for epidemiological studies.

3.2.1 Study Designs

In judging an individual study's contribution to the strength of evidence in the epidemiologic literature base, the following general hierarchy of observational study designs was considered (from most to least preferred): prospective cohort study (including nested case-control studies), case-control study, and cross-sectional study. It is important to note, however, that this hierarchy of study designs reflects the *potential* for the collection of high quality information (related to exposure, outcome, confounders, and effect modifiers) and *potential* for efficient and valid estimation of the true association. Thus, in deliberating on quality, care has been taken to consider the circumstances and particulars of each individual study to consider whether the study was well conducted independent of the type of study design.

The study designs used in the epidemiological literature reviewed were analytical and descriptive studies. Cohort and case-control study designs are analytical studies used to evaluate relative incidence of health and disease outcomes by exposure status. Cross-sectional and ecological studies are generally considered descriptive or hypothesis-generating study designs; however, they can also be used to test hypotheses regarding prevalence of health outcomes and, under certain conditions, incidence as well.

Table 3.1. Epidemiological Study Quality Considerations^a.			
Parameter	High Score	Moderate Score	Low Score
Study Design	Cohort	Case-control	Cross-sectional/Ecological

Table 3.1. Epidemiological Study Quality Considerations^a.			
Parameter	High Score	Moderate Score	Low Score
Exposure Assessment	Questionnaire and/or interview answered by subjects for chemical-specific exposure	Questionnaire and/or interview for chemical-specific exposure answered by subjects or proxy individuals	Low-quality questionnaire and/or interview; information collected for groups of chemicals rather than chemical-specific; no chemical-specific exposure information collected; ever/never use of pesticides in general evaluated
Outcome Assessment	State or National registries, physicians, and/or special surveillance programs with cases verified by histopathological evaluation for tumors; appropriate consideration of prevalent vs. incident cases; analysis by valid method specific for biomarkers	State or National registries, physicians, and/or special surveillance programs without histopathological verification for tumors; analysis by assays that are less specific for biomarkers of interest	No outcome evaluated; unclear/no consideration for whether prevalent or incident cases are appropriate; biomarker methods not validated
Confounder Control	Good control for important confounders related to cancer, standard confounders, and known confounders for glyphosate and cancer outcomes (e.g., exposure to multiple pesticides) through study design or analytic control with well measured co-exposures (i.e., cumulative exposure)	Moderately good control for confounders related to cancer; standard variables accounted for and; attempt to control for known confounders via a less efficient measure of co-exposure (e.g., ever/never use)	No adjustments for confounders
Statistical Analyses	Appropriate to study question and design, supported by relatively adequate sample size, maximal use of data, reported well	Acceptable methods, lower/questionable study power	Minimal attention to statistical analyses, sample size evidently low, comparison not performed or described clearly
Risk of (Other) Bias	Major sources of other potential biases not likely present, present but analyzed, unlikely to influence magnitude and direction of effect estimate, no/low potential of selection bias	Other sources of bias present, acknowledged but not addressed in study, may influence magnitude but not direction of estimate, evidence of potential selection bias with low impact on effect estimate	Major study biases present, unacknowledged or unaddressed in study, cannot exclude other explanation for study findings, evidence of selection bias with high potential to impact effect estimate

^a Overall study quality ranking based on comprehensive assessment across the parameters.

3.2.1.1 Analytical Studies

(1) Cohort Study

In a typical cohort study, such as the Agricultural Health Study (AHS), individuals are classified according to exposure status (i.e., presence, absence, or magnitude of exposure) and then followed over time to quantify and compare the development (i.e., incidence) of the health outcome of interest by exposure group. Conceptually, the non-exposed comparison group in a cohort study provides an estimate of the incidence of the outcome among the exposed, had they, counter-to-fact, not been exposed. Apart from chance variations, a valid cohort study comparing exposed individuals to non-exposed individuals provides an estimate of the relative risk (or rate) of the disease associated with exposure. Ideally, the exposed and non-exposed groups are exchangeable, in the sense that switching the exposed to non-exposed, and non-exposed to exposed would yield the same measure of association (e.g., relative risk). If this were the case then, apart from chance, a cohort study would yield a measure of association equivalent to that produced in a corresponding (intervention) study where exposure status was randomly assigned.

The chief advantage of the cohort study design is that it affords the investigator the opportunity to avoid and/or adjust for potential biases (i.e., selection bias, information bias, and confounding); however, these biases may also be avoided in other well-designed study designs, such as a case-control study. Cohort studies also allow for discernment of the chronological relationship between exposure and outcome, and can be particularly efficient for studying uncommon exposures. The primary disadvantage of the cohort study design is logistical inefficiency with respect to the necessary time, expense, and other resources needed to conduct them. Cohort studies are particularly inefficient for evaluating associations with rare outcomes and diseases with long induction or latency periods. Case-control studies that are nested within a cohort study (nested case-control studies) share the attributes of the cohort study and may be more efficient. However, when follow-up throughout the study period is incomplete, the potential for selection bias is increased, especially if follow-up rates are related to exposure status.

Two sub-categories of cohort studies – prospective and retrospective – are often applied to distinguish between studies in which the health outcome has occurred (retrospective study), or has not occurred (prospective study) at the time the investigators initiate the study. This distinction is important primarily as it pertains to the potential differences in the quality (e.g., completeness, accuracy, and precision) of information that can be ascertained by the investigators, and also as it relates to potential sources of bias. Although not always true, the prospective study design is considered the preferable of the two, as investigators can potentially have more choices in determining how exposure, outcome, and covariate information is collected. In a retrospective study conducted to evaluate the same hypothesis, by contrast, the investigators would have to rely on exposure information based on self-reporting or historical records. Such reporting is subject to (human) errors in recall, however when such errors are uncorrelated with disease state, there can be a bias towards the null due to random exposure measurement error (information bias) and only when such errors are correlated with the disease state can there be bias away from the null.

(2) Case-Control Study

In a typical case-control study, individuals are classified according to their outcome status (i.e., cases who have developed the outcome of interest, and controls who represent the population

from which the cases arise). The relative odds of exposure are then compared between cases and controls. The primary advantage of case-control studies is that they are logistically efficient relative to cohort studies, often being conducted at a fraction of the cost and in a fraction of the time as a corresponding cohort study. Case-control studies can be used to examine associations between multiple exposures and a given health outcome. They are particularly efficient for evaluating rare outcomes, but are inefficient for studying uncommon exposures. An important point to evaluate in each case-control study is the potential for selection bias, which arises if the exposure distribution among the control subjects is not representative of the exposure distribution among the population that gave rise to the cases. When participation rates between cases and controls are low or distinctly imbalanced, the potential for selection bias is increased, especially if participation rates are related to exposure status. Case-control studies that rely on self-reported exposure measures are also potentially susceptible to information bias which could result in bias towards the null or away from the null.

3.2.1.2 Descriptive Studies

Cross-sectional studies are used to evaluate associations between exposure and outcome prevalence in a population at a single point in (or period of) time. The primary advantage of a cross-sectional study is logistical efficiency. They are relatively quick and inexpensive to conduct, as a long period of follow-up is not required, and exposure and outcome assessments occur simultaneously. Cross-sectional studies have three primary *potential* disadvantages: 1) potential difficulty in discerning the temporal relationships (i.e., whether the exposure precedes the outcome); 2) estimating outcome prevalence rather than incidence of the outcome; and 3) the possible overrepresentation of cases of the outcome with long duration relative to the average in the population, and often with a better prognosis.

Ecological studies are used to evaluate associations between exposures and outcomes using population-level rather than individual-level data. The primary advantages of ecological studies are related to logistical efficiency, as they often rely on pre-existing data sources and require no individual-level exposure, outcome, or covariate assessments. The primary weakness of the ecologic study is the potential for confounding and resultant inappropriate extrapolation of associations observed on the aggregate-level to associations on an individual level. The discrepancy that associations observed at the population level are not observed at the individual level is referred to as the ecological fallacy. Semi-ecological studies are less susceptible to the ecological fallacy due to incorporation of individual-level data on outcomes and/or confounders. The quality of these studies depends on the ability of the group exposure data to represent individual exposure and the research question of interest.

3.2.2 Exposure Measures

As described in Section 3.2 and Figure 3.1, studies assigned a low quality ranking based on an initial evaluation were not further evaluated in detail. In all of the studies included in the analysis that were reviewed and ranked for study quality, exposure information was collected from subjects and/or proxy individuals via questionnaires and/or interviews. These exposure assessments typically include questions to determine the amount of direct pesticide use or to collect information on behaviors and conditions associated with pesticide use (e.g., occupation, tasks). This type of reporting likely misclassifies actual pesticide exposure. If conducted as part

of a prospective exposure assessment, these errors are likely to be non-differential with respect to the outcome(s) of interest. In a retrospective assessment, the subject or proxy has knowledge of the outcome; therefore, these errors may be differential or non-differential. Studies that exclusively used subjects rather than including proxy individuals were considered more reliable and given a higher weight given that the subjects would have a more accurate recollection of their own exposure.

3.2.3 Outcome Measures

All of the studies evaluated in detail, except one, utilized state or national cancer registries, physicians, and/or special surveillance programs to determine outcome status (i.e., subjects with or without a cancer of interest). In several studies, the cases were also verified by histopathological evaluation. Overall, outcome measures were relatively consistent across studies and these assessments are likely to have minimal errors. The remaining study evaluated in detail (Koureas et al., 2014) assessed oxidative DNA damage rather than a type of cancer. For this evaluation, the oxidation by-product 8-hydroxydeoxyguanosine (8-OHdG) was measured by enzyme immunoassay. This type of assay generally exhibits low specificity. More sensitive quantitative methods are available to analyze genomic DNA for 8-OHdG by high-performance liquid chromatography (HPLC) with electrochemical detection, gas chromatography-mass spectrometry (GC-MS), and HPLC tandem mass spectrometry. Consideration of incident or prevalent cases should also be carried out. By using only incident cases, there is greater confidence that exposures occurred prior to the development of the outcomes. Inclusion of prevalent cases can lead to an over-representation of cases with a long course of disease.

3.2.4 Confounding

The degree to which confounders were controlled varied across studies. Some studies adjusted for particular medical variables, while others did not. Some standard variables, such as age, geographical location, and sex, were either adjusted for analytically or by matching in case-control studies. Several studies collected information on potential confounders; however, not all of these variables were evaluated or results of the evaluation were not reported. The direction and magnitude for confounders are, in general, difficult to determine because they are dependent upon the relationship of each confounding factor with glyphosate and the cancer under investigation. Several studies considered the potential for confounding from co-exposure to other pesticides; however, only a few reported effect estimates between glyphosate exposure and cancer risk adjusted for the use of other pesticides. Given most people in the epidemiological studies who use pesticides occupationally will be exposed to multiple pesticides and, in some instances, those other pesticides were observed to be risk factors for the same cancer, this is a particularly important concern to address in either the study design or in the statistical analyses. Across numerous studies, co-exposures to other pesticides was found to be positively correlated with exposure to glyphosate and exposure to those other pesticides appear to increase the risk of some cancers. As a result, the direction of confounding would be to inflate any true effect of glyphosate in the absence of statistical control. This underlines the importance of controlling for co-exposures to other pesticides.

For NHL, other potential confounders, such as exposure to diesel exhaust fumes, solvents, ultraviolet radiation, livestock, and viruses, have been identified. Some of these are more

plausible than others. For example, occupational exposure to diesel exhaust fumes (e.g., McDuffie et al., 2002; Karunanayake et al. 2008; Baris et al. 2001; Maizlish et al. 1998) and solvents (Wang et al., 2009; Kato et al., 2005; Olsson and Brandt, 1988) are considered likely to increase the risk of NHL. Agricultural workers are exposed to diesel fumes when using agricultural vehicles when applying pesticides, such as glyphosate, and when using heavy equipment during mixing, loading, and/or applying pesticides. Agricultural workers are also exposed to solvents. Solvents are often used in pesticide products to aid the delivery of the active ingredient and enhance efficacy. Solvents are also used for cleaning and maintenance/repair of agricultural equipment used for mixing, loading, and/or applying pesticides. With an association between exposure and outcome of interest, it is reasonable to consider diesel exhaust fumes and solvents as probable confounders; however, neither of these factors were accounted for in any of the studies evaluated in detail. There is also evidence that ultraviolet (UV) radiation may increase the risk of NHL (Karipidis et al., 2007; Zhang et al., 2007). As a result, there is a support that UV radiation is also a potential confounder given the extended amount of time agricultural workers spend outside performing activities, including those associated with pesticide use. This was also not accounted for in any of the studies evaluated in detail.

3.2.5 Statistical Analyses

Statistical analyses that were appropriate to the study question and study design, supported by adequate sample size, maximized the use of available data, and were well characterized in the report were weighted most highly. Acceptable statistical methods, questionable study power, and analytical choices that resulted in the loss of information were given moderate weight. Reports with only minimal attention paid to the conduct and reporting of the statistical analyses were given the lowest weight.

3.2.6 Risk of Bias

The internal validity of the studies reviewed was judged by noting the design strategies and analytic methods used in each study to constrain or eliminate selection bias, information bias, and confounding. Selection bias can occur when the sampling of the population by the investigator yields a study population that is not representative of the exposure and outcome distributions in the population sampled. Put simply, selection bias occurs if selection of the study sample yields a different estimate of the measure of association than that which would have been obtained had the entire target population been evaluated. Although there are numerous sources of selection bias, there are several mechanisms that may have induced selection bias in the studies reviewed: low participation rates of eligible individuals due to non-responsiveness or refusal (self-selection bias); loss to follow-up (i.e., failure to retain all study participants initially enrolled in the study); and, in a case-control study, control selection bias arising because the exposure distribution in the control sample does not represent the exposure distribution of the study base (i.e., the population that gave rise to the cases or more formally, the person-time experience of that population).

Information bias (also referred to as observation bias) arises when study participants are incorrectly categorized with respect to their exposure or outcome status, or when errors arise in the measurement of exposure or outcome, in the case of continuously distributed measures.

Epidemiologists often distinguish between two mechanisms or types of misclassification – those that are non-differential (or random) and those that are differential (non-random). Non-differential misclassification of exposure (or non-differential exposure measurement error) occurs when the probability or magnitude of error in the classification or measurement of exposure is independent of the outcome status of the study participants. Non-differential exposure measurement error typically results in a bias towards the null which may obscure any true effect of the exposure of interest. Similarly, non-differential misclassification of outcome (or outcome measurement error) occurs when the probability or magnitude of error in the assignment of outcome status or level is independent of exposure status. Non-differential outcome measurement error typically does not cause bias but does decrease the precision of effect estimates and therein inflates the width of confidence intervals. In contrast, differential exposure misclassification (or measurement error) occurs when the error in the exposure assignment is not independent of the outcome status. The mechanisms that cause non-differential misclassification in the currently reviewed literature include random errors in exposure recall from subjects or proxy respondents. The mechanisms that could induce differential misclassification include recall bias and interviewer/observer bias. Note that mismeasurement of confounders can result in residual confounding of the association of interest, even when adjustment for that confounder has been conducted in the analysis.

Studies in which major sources of potential biases were not likely to be present, studies in which potential sources of bias were present, but effectively addressed and analyzed to maximize the study validity, and studies in which sources of bias were unlikely to influence the magnitude and direction of the effect estimate were given more weight than studies where sources of bias may be present, but not addressed in the study.

3.3 Review of Quality Results

Each study was judged to be of high, moderate, or low quality in each of the six domains affecting study quality, as discussed above and in Table 3.1. The results of the quality assessment are presented separately for each group below. The quality rankings presented are specific to the current evaluation of the carcinogenic potential of glyphosate. As noted above and in Table 3.2, several studies were not included in the ranking evaluation because they did not represent the most complete analysis. Rather, the subjects were included in a larger analysis (e.g., pooled case-control study) to produce a greater number of cases and controls (see Appendix B for visual representation of these studies). For example, Cantor *et al.* (1992) was not individually evaluated for ranking because the data from this study were pooled with data from other studies in De Roos *et al.* (2003), which was included.

3.3.1 “High” Quality Group

Three studies were given a high quality ranking: De Roos *et al.* (2005), Eriksson *et al.* (2008), and Koutros *et al.* (2013).

De Roos *et al.* (2005) was the only cohort study available for ranking. This prospective cohort study evaluated associations between various pesticide exposures, including glyphosate, and cancer incidence for numerous solid and non-solid tumors in the AHS. The aim of the AHS is to evaluate the role of agricultural exposures in the development of cancer and other diseases in the

farming community. AHS recruited 52,934 licensed private pesticide applicators along with 32,345 of their spouses between 1993 and 1997. In the first two phases of the study, the cohort also included 4,916 commercial pesticide applicators from Iowa. As a prospective analysis of the AHS cohort, information was obtained from exposed subjects at enrollment and no proxies were necessary. Exposure was evaluated as ever/never use, cumulative lifetime exposure, and intensity-weighted cumulative exposure. Due to the study design, the potential for many biases were reduced. Additionally, the study adjusted and/or considered numerous factors, including use of other pesticides. Study participants provided detailed pesticide exposure information prior to enrollment in the study and this information has been incorporated into the study evaluation by determining tertile cut points and calculating effect estimates by comparing to the lowest tertile. Additional evaluations with quartiles and quintiles were performed for cancers with elevated effect estimates in the study and for NHL.

Eriksson *et al.* (2008) was a population-based case-control study that recruited a consecutive series of incident cases of NHL in several regions of Sweden from physicians treating lymphoma within specified health service areas. Cases were verified pathologically and matched to randomly selected controls from the national population registry by age, sex and health service area. Exposure information was collected from exposed individuals (i.e., no use of proxy respondents) using a comprehensive questionnaire including a total work history with in depth questions about exposures to pesticides, solvents, and other chemicals. Interviewers were blinded to case/control status. The study only reported minimal demographic information on subjects (age and sex) and a table with subject characteristics (e.g., smoking status, alcohol intake, physical activity, education) that could potentially be used to adjust effect estimates was not provided. Glyphosate exposure was reported in 29 cases and 18 controls during the study period. Multivariate analyses were adjusted for co-exposure to different agents, including MCPA, “2,4,5-Y and/or 2,4-D”, mercurial seed dressing, arsenic, creosote, and tar. An analysis for a potential exposure-response relationship was also conducted; however, it was not clear whether this analysis controlled for co-exposure to other pesticides based on the statistical methods description. The number of cases and controls were also not reported for this analysis.

Koutros *et al.* (2013) was a nested case-control study within the AHS that evaluated the association between pesticide use and prostate cancer. Exposure information was collected from exposed subjects (no proxies necessary) through the enrollment questionnaires, as well as in a follow-up questionnaire administered 5 years after enrollment. This study evaluated the association between glyphosate and prostate cancer diagnoses from enrollment (1993-1997) through 2007 resulting in a longer follow-up time than many of the other case-control studies that utilized AHS subjects. The study used lifetime cumulative exposure and intensity-weighted cumulative exposure metrics. Analyses were also conducted using unlagged exposure and 15-year lagged exposure, which excluded the most recent 15 years of exposure for both exposure metrics. Although the effect estimate reported for glyphosate in this study was not adjusted for co-exposure to other pesticides, additional analyses were not considered necessary since there was no association observed.

3.3.2 “Moderate” Quality Group

Twenty-one case-control studies were assigned a moderate quality rating (Table 3.2). In general, these studies share many study design characteristics. Exposure information was collected from subjects and/or proxy individuals, the outcome measurement(s) utilized state/national registries and surveillance programs, appropriate statistical analyses were performed, some covariates but maybe not all relevant covariates were evaluated and/or considered, and risks of bias were minimized to some extent. Sample sizes varied across studies. Case-control studies investigating solid tumors included study populations in the United States and Canada. For non-solid tumors, study populations were located in the United States, Canada, Sweden, France, Germany, Italy, Ireland, Spain, and the Czech Republic. Although several nested case-control studies shared most of the characteristics of the AHS cohort study, these studies were primarily given a moderate quality ranking since co-exposure to other pesticides was not accounted for in the analyses.

3.3.3 “Low” Quality Group

Seven case-control and 27 cross-sectional/ecological studies were assigned a low quality ranking. All of these studies, except one case-control study (Cocco *et al.*, 2013) and one descriptive study (Koureas *et al.*, 2014), were not subjected to a detailed evaluation because they did not report a quantitative measure of an association between glyphosate exposure and a cancer outcome, did not collect information on glyphosate exposure from all subjects, and/or did not evaluate risk to a cancer outcome (Appendix D). In many instances, effect estimates were reported only for total pesticide exposure. Additionally, exposure was assumed and glyphosate-specific exposure information was not collected. In other studies, the aim of the study was to assess exposure methods for epidemiological studies and/or to evaluate the impact of exposure misclassification; therefore, there was no evaluation of a cancer outcome.

It should be noted that some of the studies assigned a low quality ranking in the current evaluation were included in the recent evaluation by IARC. There were a number of descriptive studies that evaluated the genotoxicity in human populations; however, these studies did not meet the criteria for inclusion in the ranking as described in Section 3.2 and Figure 3.1. In most instances, these studies reported effect estimates for total pesticide exposure and/or assumed glyphosate exposure without collecting glyphosate-specific exposure information. For case-control studies, Cocco *et al.* (2013), Dennis *et al.* (2010) and Ruder *et al.* (2004) were included in the 2015 IARC evaluation, but were not considered informative in the current evaluation.

Detailed evaluations were not performed in the current evaluation for Dennis *et al.* (2010) and Ruder *et al.* (2004) because a quantitative measure of an association between glyphosate and a cancer outcome was not reported. Cocco *et al.* (2013) received a detailed evaluation and was assigned a low quality ranking. This case-control study, which evaluated lymphoma risk across six European countries, was not considered informative due to a combination of numerous limitations in the study. The power of the study was low with only four cases and two controls exposed to glyphosate. Control ascertainment was not consistent across countries, with a mix of hospital- and population-based controls used. The overall participation rate for population-based controls was found to be much lower than the overall participation rates of the cases or hospital-based controls. Lastly, the study was limited to ever/never use of glyphosate and did not control for confounders, in particular exposure to other pesticides. Although this study was included in

the IARC evaluation, IARC also stated that the study had very limited power to assess the effects of glyphosate on risk of NHL.

The other study subjected to a detailed evaluation and assigned a low quality ranking was Koureas *et al.* (2014). This cross-sectional study evaluated the association between glyphosate exposure and oxidative DNA damage in 80 Greek pesticide sprayers. Although the study reported a non-statistically significant effect estimate for glyphosate, it is limited in its ability to contribute to the overall evaluation of the carcinogenic potential of glyphosate. The effect estimate was not adjusted for any standard covariates or potential confounders, including co-exposure to other pesticides. The power of the study was questionable. There were 80 subjects, but the number exposed to glyphosate was not reported. The outcome is measured using an immunoassay that is less specific for measuring the biomarker of interest than other available analytical methods. Lastly, the study evaluates primary DNA damage, but does not measure the consequence of genetic damage. An increase in oxidative DNA damage may lead to cell death or initiate DNA repair rather than lead to a mutation.

Due to the limitations in the studies assigned a low quality ranking, they do not provide reliable information to evaluate associations between glyphosate exposure and cancer outcomes. Therefore, the remaining sections of this document do not further discuss these studies except to note when a study is included in meta-analyses.

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
Alavanja <i>et al.</i> (2003)	This study was not included in the study quality ranking because the data were used in the updated analysis by Koutros <i>et al.</i> (2013).						
Andreotti <i>et al.</i> (2009)	Nested Case-control	Questionnaire answered by subjects at study enrollment followed by take-home questionnaire; examined exposure for glyphosate as ever/never, and intensity-weighted cumulative exposure days; spouses either self-administered questionnaire (81%) or telephone interview (19%)	State cancer registries without histopathological verification; exclusion of subjects with prevalent cancer at enrollment; follow-up ~ 9 years	Adjusted for age, smoking, and diabetes for both exposure metrics as well as applicator type forever/never exposure metric No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Unconditional logistic regression to obtain OR and 95% CI	Exposure misclassification particularly for spouses, low response rate to take-home questionnaire (40%) but unclear if affected cases and controls differently, insufficient power for pesticide exposure interactions	Moderate
Band <i>et al.</i> (2011)	Population-based case-control Males only	Self-administered questionnaire answered by subjects or proxies for deceased subjects requesting work history and demographic information; use of a job exposure matrix to estimate exposure to pesticides	Cancer registry with histopathological verification; excluded farmers that worked all outside of British Columbia; included prostate cancer cases prior to the PSA era	Adjustment for alcohol consumption, cigarette years, education level, pipe years, and respondent type. Marital status and ethnicity not significant No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Conditional logistic regression to obtain ORs and 95% CIs	Recall bias, use of proxy for deceased, exposure misclassification, participation rates cited from another study, use of cancer patients as controls (excluding lung and unknown cancer)	Moderate
Brown <i>et al.</i> (1990)	Pooled population-based case-control Males only	In-person interviews using standardized questionnaire with subjects or proxies for deceased/incapacitated; supplementary questionnaire administered by telephone for Iowa subjects to obtain more	State cancer registry (Iowa) and special surveillance network including hospitals and pathology laboratories (Minnesota); cases ascertained retrospectively and prospectively (2 years after start of study);	Adjusted for vital status, age, state, ever used tobacco daily, close relative with lymphopoietic cancer, nonfarming job related to risk of leukemia in the study, exposure to substances related to risk in this study	Unconditional logistic models to obtain OR and 95% CI; questionable power (15 cases)	Recall bias; exposure misclassification, use of proxy respondents	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
		detailed information from those indicating pesticide use	~26% of cases deceased or too ill when identified and ~15% deceased or too ill at time of interview; histopathological verification by pathologists	(benzene, naphthalene, hair dyes) No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)			
Brown <i>et al.</i> (1993)	Population-based case-control Males only	In person interviews with standardized questionnaire to obtain detailed information on farm activities and use of pesticides from subjects or proxies	State cancer registry (Iowa) ascertained retrospectively and prospectively (2 years after start of study); ~26% of cases deceased or too ill when identified and ~15% deceased or too ill at time of interview; histopathological verification by pathologists	Adjusted for vital status and age; smoking and education evaluated and not found to be significant No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Logistic models to obtain OR and 95% CI; questionable power (11 cases)	Recall bias; exposure misclassification, use of proxy respondents	Moderate
Cantor <i>et al.</i> (1992)	This study was not included in the study quality ranking because the data were used in the pooled analysis conducted by De Roos <i>et al.</i> (2003).						
Carreon <i>et al.</i> (2005)	This study was not included in the study quality ranking because the data were used in the pooled analysis conducted by Yiin <i>et al.</i> (2012).						
Cocco <i>et al.</i> (2013)	European multi-center case-control Hospital-based and population-based (mixed for 2 countries, only hospital-based for the rest)	Trained interviewers conducted in person interviews using structured questionnaire answered by subjects; those identified as agricultural worker on questionnaire given subsequent questions about pesticide use, crops, etc.	Surveillance centers, 20% of slides from each center reviewed by pathologist	Adjustment for age, sex, education, and center. No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Unconditional logistic regression to obtain ORs and 95% CIs; Low power (4 cases, 2 controls)	Recall bias, selection bias (low response rate for population-based controls and differed from cases), exposure misclassification, mix of hospital- and population-based controls,	Low
De Roos <i>et al.</i> (2003)	Population-based case-control Males only Pooled analysis of	Interviews with subjects or proxy for deceased subjects. Different interview techniques across states. One study collected information on	State cancer registries (one state chose a random sample, other states chose all cases), surveillance programs, and hospitals without	Adjustment for age, study site, and other pesticides. First degree relative with haematopoietic	Logistic regression and hierarchical regression to obtain ORs and 95% CIs	Recall bias, exposure misclassification, , use of proxy for deceased, , varying quality of questionnaire/interview techniques across studies	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.

Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
	Cantor <i>et al.</i> , 1992; Hoar <i>et al.</i> , 1986; Zahm <i>et al.</i> , 1990	pesticide use and then followed-up with questions on selected specific pesticides, another study had a direct question about a selected list of specific pesticides, and the last study used an open ended question without prompting for specific pesticides	histopathological verification	cancer, education, and smoking not found to be important confounders. No adjustment for other potential confounders (e.g., solvents, diesel fumes, UV radiation)			
De Roos <i>et al.</i> (2005)	Prospective cohort (licensed pesticide applicators)	Questionnaire answered by subjects at enrollment and with subsequent take-home questionnaire; examined exposure as ever/never, cumulative lifetime days, and intensity-weighted cumulative exposure days	State cancer registries without histopathological verification; follow-up ~7 years	Adjustment for state of residence, age, education, smoking history, alcohol consumption, family history of cancer, use of other common pesticides No adjustment for other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Poisson regression to obtain RRs and 95% CIs	Major sources of potential biases unlikely, potential exposure misclassification due to any changes in exposure since enrollment, follow-up period may be limited	High
Engel <i>et al.</i> (2005)	Nested case-control Females only	Take-home questionnaire from spouses of enrolled applicators used to obtain farm exposures, general health information, and reproductive health history; Information obtained from applicators used as measure of possible indirect exposure to spouses	State cancer registries identifying malignant breast cancer; ~5 years average follow-up time	Adjusted for age, race and state. Evaluated BMI, age at menarche, parity, age at first birth, menopausal status, age at menopause, family history of breast cancer, physical activity, smoking, alcohol consumption, fruit and vegetable consumption and education but none	Poisson regression to obtain RRs and 95% CIs	Exposure misclassification, exposure to other pesticides (however no association observed), lack of information on length of marriage could result in overestimating exposure based on husband	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
				found to be significant No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)			
Eriksson <i>et al.</i> (2008)	Population-based case-control	Questionnaire answered by subjects; follow-up by phone if incomplete answers; excluded exposures that occurred during the same calendar year and year before diagnosis (cases) or enrollment (controls); minimal demographic information reported	Physicians treating lymphoma within specified health service areas and verified by pathologists	Adjustment for age, sex, year of diagnosis/enrollment, as well as exposure to other pesticides in multivariate analyses. Not stated what adjustments were made for other pesticides in latency analyses. No adjustment other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Unconditional logistic regression and multivariate analyses to obtain ORs and 95% CIs; not clear how multivariate was performed; questionable power (29 cases, 18 controls); also included analysis of ≤ 10 vs. >10 years exposure	Recall bias, exposure misclassification, lack of subject demographics/ characteristics (e.g., smoking, alcohol consumption, race, etc)	High
Flower <i>et al.</i> (2004)	Nested case-control	Questionnaire answered by applicators at enrollment; spouses enrolled through a questionnaire brought home by applicator; females (applicators and spouses) were asked to complete a questionnaire on female and family health that collected information on children born during or after 1975	State cancer registry to identify childhood cancer cases (diagnosed from birth through 19 yrs of age) for children of parents enrolled; hybrid prospective/retrospective ascertainment; excluded female applicators	Child's age at parent's enrollment was included in model; parental age at child's birth, child's sex, child's birth weight, history of parental smoking, paternal history of cancer, and maternal history of miscarriage were evaluated but not found to be significant and not included in model No adjustment for co-	Logistic regression to obtain OR and 95% CI; calculated standardized incidence ratios to compare observed number of childhood cancer cases identified to the expected number; low/questionable power (6 parental cases, 13 maternal cases)	Exposure misclassification, lack of timing data to determine if exposure occurred prior to conception or during pregnancy, exposure to other pesticides (however no association observed and lack of power for adjustment)	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
				exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)			
Hardell and Eriksson (1999)	This study was not included in the study quality ranking because the data were used in the pooled analysis conducted by Hardell <i>et al.</i> (2002).						
Hardell <i>et al.</i> (2002)	Population-based case-control Males only Pooled analysis of Hardell and Eriksson 1999 and Nordstrom <i>et al.</i> , 1998	Questionnaire answered by subjects or proxy for deceased subjects to obtain complete working history and exposure to different chemicals; follow-up with interview for clarification	Registries with histopathological verification	Adjustment for age, vital status, and county (by matching). Exposure to other pesticides in multivariate analysis. No adjustment for other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Conditional logistic regression to obtain OR and 95% CI (univariate and multivariate analyses). Questionable power (8 cases/8 controls)	Recall bias, exposure misclassification, use of proxy for deceased	Moderate
Hohenadel <i>et al.</i> (2011)	This study was not included in the study quality ranking because a more complete analysis was conducted by McDuffie <i>et al.</i> (2001).						
Kachuri <i>et al.</i> (2013)	Population-based case-control Males only	Questionnaire answered by subjects or proxies; pesticide use collected via detailed telephone interview on all participants with 10+ hours of pesticide use during lifetime and 15% random sample of those who did not; exposure based on lifetime exposure to glyphosate	Cancer registries or hospitals in 6 Canadian provinces with histopathological verification for 36.55% of samples	Adjustment for age, province, selected medical conditions, family history of cancer, use of proxy respondent, smoking status No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Unconditional logistic regression to obtain OR and 95% CI; trends examined using multiple logistic regression	Recall bias, exposure misclassification, control selection based on three different sources depending on province of residence, low participation rates among controls, use of proxy respondents	Moderate
Karunanayake <i>et al.</i> (2012)	Population-based case-control Males only	Questionnaire answered by subjects; pesticide use collected via detailed telephone interview on all participants with 10+ hours of pesticide use during lifetime and 15%	Cancer registries or hospital in 6 Canadian provinces with histopathological verification for 49% of samples; difficulty recruiting control	Adjusted for age, province of residence, and significant medical history variables No adjustment for co-	Conditional logistic regression to obtain OR and 95% CI	Recall bias, exposure misclassification, control selection based on three different sources depending on province of residence, low participation rates among	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
		random sample of those who did not; exposure based on lifetime exposure to glyphosate	participants for older age groups	exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)		controls, unable to evaluate Epstein-barr virus exposure	
Koureas <i>et al.</i> (2014)	Cross-sectional	Questionnaire answered by pesticide sprayers	Genomic DNA extracted from peripheral blood samples and oxidation by-product 8-hydroxydeoxyguanosine (8-OHdG) was determined by enzyme immunoassay; more specific methods (HPLC, GC-MS) are available for measurement	No adjustments. In univariate, occupational exposure, sex and alcohol consumption were statistically significant while DAP concentrations and smoking were not.	For univariate, chi-square test used to obtain RR and 95% CI; 8-OHdG levels transformed into binary variables (categorized as high and low using the 75 th percentile cut-off); unknown number of exposed and unexposed cases (questionable power possible given total number of subjects is only 80)	Recall bias, did not control for risk factors identified as statistically significant for univariate analysis, does not measure the consequence of genetic damage	Low
Koutros <i>et al.</i> (2013)	Nested case-control Males only	Questionnaire answered by subjects at study enrollment; examined exposure as cumulative lifetime days and intensity-weighted cumulative exposure days	State cancer registries with histopathological verification; total and aggressive prostate cancers evaluated	Adjustment for age, state, race, smoking, fruit servings, family history of prostate cancer, and leisure time physical activity in the winter. No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Poisson regression to obtain RRs and 95% CIs; also included unlagged vs. lagged analysis	Exposure misclassification	High
Landgren <i>et al.</i> (2009)	Nested case-control ^a Males only	Questionnaire answered by subjects at enrollment in AHS cohort and subsequent take-home questionnaire to collect	Venous blood collected from antecubital vein and analyzed for MGUS; same method as used for controls group in	Adjusted for age and education level Association with other pesticides examined	Logistic regression models to obtain OR and 95% CI comparing to population-based	Exposure misclassification, control group not from geographical area (used control group with	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
		information on 50 pesticides; occupational exposures, medical histories, and lifestyle factors updated with 5-year follow-up interview; subjects with prior history of lymphoproliferative malignancy excluded	Minnesota	and not found to be significant so no adjustment performed No adjustment for other potential confounders (e.g., solvents, diesel fumes, UV radiation)	screening study in Olmsted County, Minnesota; questionable power (27 cases; 11 controls)	similar demographics from Minnesota)	
Lee <i>et al.</i> (2004a)	This study was not included in the study quality ranking because the data were used in the pooled analysis conducted by De Roos <i>et al.</i> (2003).						
Lee <i>et al.</i> (2004b)	Population-based case-control White males and females only	Subjects or proxies were interviewed by telephone; those living/working on a farm asked for detailed history of pesticide use and farming information	State cancer registry or review of discharge diagnosis and pathology records at 14 hospitals; only newly diagnosed cases with confirmed adenocarcinoma of stomach or esophagus retained; controls randomly selected from a prior study conducted in geographical area	Adjusted for age and sex; evaluated BMI, smoking, alcohol consumption, educational level, family history of stomach or esophageal cancer, respondent type, dietary intake of particular vitamins and minerals, protein, and carbohydrates (included in model if changed value of OR by more than 10%) No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Unconditional logistic regression to obtain OR and 95% CI; questionable power (12 cases for stomach; 12 cases for esophagus)	Recall bias, exposure misclassification, use of proxy respondents, control selection	Moderate
Lee <i>et al.</i> (2005)	Population-based case-control	Questionnaire and/or interview with subject or proxy individuals to collect information on use of specific pesticides; telephone follow-up for unclear responses	Referral by hospitals or through state cancer registries with histopathological verification; controls selected from a previous study	Adjusted for age and respondent type; evaluated history of head injury, marital status, education level, alcohol consumption, medical history of diabetes mellitus,	Unconditional logistic regression to obtain OR and 95% CI	Recall bias, exposure misclassification, large number of proxy respondents, control selection (historical control group from another cancer evaluation, differences in	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
				dietary intake of α- and β-carotene, and dietary fiber (included in model if changed value of OR by more than 10%) No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)		exposure time period evaluated, needed to add younger controls, exposure information collected for different time periods for cases vs. controls)	
Lee <i>et al.</i> (2007)	Nested case-control	Questionnaire answered by subjects at enrollment in AHS cohort and subsequent take-home questionnaire to collect information on 50 pesticides	State cancer registries without histopathological verification; follow-up ~ 7 years	Adjustment for age, smoking, state, total days of pesticide application No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Unconditional multivariate logistic regression to obtain OR and 95% CI	Exposure misclassification, limited data on dietary factors, NSAID drug use and family cancer history	Moderate
McDuffie <i>et al.</i> , 2001	Population based case-control Males only	Questionnaire answered by subjects; pesticide use collected via detailed telephone interview on all participants with 10+ hours of pesticide use during lifetime and 15% random sample of those who did not; exposure based on lifetime exposure to glyphosate	Cancer registries or hospital in 6 Canadian provinces with histopathological verification for 84% of samples; ascertainment of cases stopped in each province once target numbers were reached	Adjustment for age, province, and significant medical variables (including history of cancer in study participants and family history). No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Conditional logistic regression to obtain OR and 95% CI	Recall bias, exposure misclassification, control selection based on three different sources depending on province of residence, relatively low participation rates	Moderate
Nordstrom <i>et al.</i> , 1998	This study was not included in the study quality ranking because the data were used in the pooled analysis conducted by Hardell <i>et al.</i> (2002).						
Orsi <i>et al.</i> , 2009	Hospital-based case-control	Data collection in 2 stages: 1) self-	Hospital catchment area with histopathological/	Adjustment for age, center, and	Unconditional logistic regression	Recall bias, exposure misclassification,	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.

Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
	Males only (occupationally exposed)	administered questionnaire on socioeconomic characteristics, family medical history, and lifelong residential and occupational histories and more specific information for each job held for at least 6 months, and 2) face-to-face interview with trained staff (blinded) using standardized questionnaire	cytological verification Controls were hospital based with no prior history of lymphoid neoplasms, excluding patients with cancer or a disease directly related to occupation, smoking or alcohol abuse (but history of any of these did not prevent selection as a control)	socioeconomic category. Education and housing not found to impact results. Flu immunization, previous history of mononucleosis, skin type, smoking, and drinking did not change results. Evaluated particular crops and animal husbandry as well. No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	to obtain OR and 95% CI. Questionable power (12 cases/24 controls)	hospital-based controls	
Pahwa <i>et al.</i> (2011)	Population-based case-control Males only	Questionnaire answered by subjects; pesticide use collected via detailed telephone interview on all participants with 10+ hours of pesticide use during lifetime and 15% random sample of those who did not; exposure based on lifetime exposure to glyphosate	Cancer registries or hospitals in 6 Canadian provinces with histopathological verification for 30% of samples	Adjustment for age group, province of residence, and statistically significant medical history variables No adjustment for co-exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Conditional logistic regression to obtain OR and 95% CI; trends examined using multiple logistic regression	Recall bias, exposure misclassification, control selection based on three different sources depending on province of residence, low participation rates among controls	Moderate
Pahwa <i>et al.</i> (2012)	Population-based case-control Males only	Questionnaire answered by subjects; pesticide use collected via detailed telephone interview on all participants with 10+ hours of pesticide use during lifetime and 15% random sample of those	Cancer registries or hospitals in 6 Canadian provinces with histopathological verification for 36.5% of samples	Adjustment for age group, province of residence, and statistically significant medical history variables No adjustment for co-	Conditional logistic regression to obtain OR and 95% CI; trends examined using multiple logistic regression	Recall bias, exposure misclassification, control selection based on three different sources depending on province of residence, low participation rates among controls	Moderate

Table 3.2. Summary of Study Design Elements Impacting Study Quality Assignment and Overall Ranking.							
Journal Article	Study Design	Exposure Assessment	Outcome Assessment	Confounder Control	Statistical Analyses	Risk of (Other) Bias	Overall Ranking
		who did not; exposure based on lifetime exposure to glyphosate		exposure to other pesticides or other potential confounders (e.g., solvents, diesel fumes, UV radiation)			
Yiin <i>et al.</i> (2012)	Population-based case-control Pooled analysis of men with women analyzed in Carreon <i>et al.</i> (2005)	Questionnaire and/or interview for chemical-specific exposure answered by subjects or proxy individuals	Cases referred by physicians or through state cancer registries with histopathological verification; controls matched within state, but not county of residence	Adjustment for age, education, sex, and , sex, and farm pesticide exposure (yes/no) No adjustment for other potential confounders (e.g., solvents, diesel fumes, UV radiation)	Unconditional logistic regression to obtain ORs and 95% CIs	Acknowledge other sources of bias. Recall bias, exposure misclassification, control selection (low number of deceased controls obtained)	Moderate

^a Mixed methods used in the Landgren et al (2009) study, with cross-sectional study design used to calculate prevalence rates comparing the AHS to a reference population MN. Pesticide risk estimates (including glyphosate) calculated using nested case-control approach, comparing AHS exposed/unexposed (ever/never) study participants.

3.4 Assessment of Epidemiological Studies for Relevance to Analysis

Using the criteria summarized in Section 3.2, a total of 58 individual literature studies were identified in the literature review and were judged as high, moderate, or low quality. Overall, 3 studies, 21 studies, and 34 studies were assigned high, moderate, or low rankings, respectively. All of the high and moderate quality studies were considered relevant to the current evaluation.

The majority of the studies were case-control studies evaluating a wide-range of cancers in the United States and Canada. There were several case-control studies from Canada that utilized the same study population (Kachuri *et al.*, 2013; Karunanayake *et al.*, 2012; McDuffie *et al.*, 2001; Pahwa *et al.*, 2011; Pahwa *et al.*, 2012). In a similar fashion, numerous studies in the United States were nested case-control studies, where the AHS cohort served as the source population for selecting cases and controls (Andreotti *et al.*, 2009; Engel *et al.*, 2005; Flower *et al.*, 2004; Koutros *et al.*, 2013; Landgren *et al.*, 2009; Lee *et al.*, 2007). In these studies, a subset of the AHS cohort were selected based on their outcome status for a particular cancer and exposure information was used from the AHS enrollment questionnaire and/or during follow-up interviews. Nested case-control studies allow for testing of hypotheses not anticipated when the cohort was initially assembled. In the AHS prospective cohort study (De Roos *et al.*, 2005), exposure and demographic information were also obtained from the questionnaires at enrollment; however, subjects were enrolled prior to developing cancer outcomes of interest. Subjects were then followed from enrollment to a subsequent time point to determine if subjects developed cancer outcomes of interest. As such, all available subjects in the cohort are included in the evaluation of whether there was an association between a risk factor (e.g., glyphosate exposure) and outcome.

The moderate studies included a varying degree of control for confounding and biases across studies. As moderate studies, they encompass a combination of strengths and limitations. In particular, important factors that impacted the quality assessment for these studies included whether there was control for known confounders, identification of control selection issues, study power issues, and length of follow-up. As noted previously, most people in these epidemiological studies used pesticides occupationally and were exposed to multiple pesticides over their working lifetime. Therefore, exposure to other pesticides is a particularly important factor to control for and studies that made this adjustment were given more weight than those that did not. Similarly, control selection issues were noted in a few studies and were given less weighting than those without control selection issues. The issues ranged from concerns using hospital-based controls, using different population sources to ascertain controls within the same study, and appropriateness of using controls ascertained for another research question. Numerous studies had limited power due to small sample size, which results in large confidence intervals and reduces the reliability of the results to demonstrate a true association. Studies demonstrating low or questionable power were therefore given less weighting. Lastly, the length of follow-up time varied across studies.

3.5 Summary of Relevant Epidemiological Studies

A summary of the relevant studies evaluating the association between glyphosate exposure and cancer are discussed below. Results of the studies reporting data on glyphosate exposure and

solid tumors (non-lymphohematopoietic) at various anatomical sites are presented in Table 3.3. Results of the studies reporting data on glyphosate exposure and non-solid tumors (lymphohematopoietic) are presented in Table 3.4. For study details, see Table 3.2 above and Appendix C.

3.5.1 Solid Tumor Cancer Studies

(1) Cancer at Multiple Sites from the AHS Cohort

De Roos *et al.*, (2005) evaluated associations between glyphosate exposure and cancer incidence of all cancers combined in the AHS cohort study and did not find an association [ever/never use relative risk ratio (RR) =1.0 with 95% confidence interval (CI) of 0.90–1.2) when adjusting for age, demographic and lifestyle factors, and exposure to other pesticides]. In addition, De Roos *et al.*, 2005 evaluated cancer at specific anatomical sites. Along with several nested case-control studies, no statistical evidence of an association with glyphosate was observed at any specific anatomical site (Table 3.3). Specifically, AHS researchers reported no evidence of an association between glyphosate use and cancers of the oral cavity (De Roos *et al.*, 2005), colon (De Roos *et al.*, 2005; Lee *et al.*, 2007), rectum (De Roos *et al.*, 2005; Lee *et al.*, 2007), lung (De Roos *et al.*, 2005), kidney (De Roos *et al.*, 2005), bladder (De Roos *et al.*, 2005), pancreas (De Roos *et al.*, 2005; Andreotti *et al.*, 2009), breast (Engel *et al.*, 2005), prostate (De Roos *et al.*, 2005; Koutros *et al.*, 2013) or melanoma (De Roos *et al.*, 2005). The adjusted RR or odds ratio (OR) and 95% CI for these studies are provided in Table 3.3.

(2) Prostate Cancer

In a Canadian population-based study (Band *et al.*, 2011), researchers reported non-statistically significant elevated odds of prostate cancer in relation to glyphosate use (OR=1.36; 95% CI=0.83–2.25). There was no adjustment made for exposure to other pesticides. This study included prostate cancer cases from 1983-1990, prior to the prostate-specific antigen (PSA) era. Consequently, the study included more advanced tumors before diagnosis. The AHS related studies (De Roos *et al.*, 2005; Koutros *et al.*, 2013), reflect PSA-era cases (i.e., cases which are typically identified at an earlier stage in the progression of the disease) and also did not identify an association with prostate cancer.

(3) Brain (Glioma) Cancer

Lee *et al.* (2005) investigated the association between brain cancer with farming and agricultural pesticide use. Matching for age, sex, vital status, and region, study authors reported a non-significant elevated odds of glioma (OR=1.5; 95% CI=0.7–3.1) in relation to glyphosate use by male farmers; however, the results were significantly different between those who self-reported pesticide use (OR=0.4; 95% CI=0.1–1.6), and for those for whom a proxy respondent was used (OR=3.1; 95% CI=1.2–8.2), indicating recall bias was a potential factor in this study. Furthermore, there was no adjustment for co-exposure to other pesticides and issues noted with control selection.

A population-based case-control study evaluated the risk of brain cancer, specifically, glioma risk, among men and women participating in the Upper Midwest Health Study (Yiin *et al.*,

2012). Using a quantitative measure of pesticide exposure (in contrast to an ever-use metric), Yiin *et al.* (2012) observed no statistical evidence of an association with glyphosate with effect estimates roughly equal to the null value following adjustment for age, education, sex, and use of other pesticides (home and garden use: OR=0.98; 95% CI=0.67–1.43; non-farm jobs: OR=0.83; 95% CI=0.39–1.73).

(4) Stomach and Esophageal Cancer

In a population-based case-control study in eastern Nebraska, Lee *et al.* (2004b) investigated pesticide use and stomach and esophageal adenocarcinomas. There was no association observed between glyphosate exposure and either stomach cancer (OR=0.8; 95% CI=0.4–1.5) or esophageal cancer (OR=0.7; 95% CI=0.3–1.4) after adjustment for age and sex. No adjustment was made for exposure to other pesticides.

(5) Soft Tissue Sarcoma

A Canadian case-control study (Pahwa *et al.*, 2011) examined exposure to pesticides and soft tissue sarcoma and found no relation with the use of glyphosate after adjustment for age, province of residence, and medical history variables (OR=0.90; 95% CI= 0.58–1.40); however, control selection issues were noted, including low response rate and selection from three different sources depending on the province of residence.

(6) Total Childhood Cancer

Flower *et al.* (2004), a nested case-control study in the AHS cohort, examined the relation between parental pesticide use and all pediatric cancers reported to state registries among children of AHS participants and did not observe a significant association with maternal use exposure to glyphosate (OR=0.61; 95% CI= 0.32–1.16) or paternal (prenatal) exposure to glyphosate (OR=0.84; 95% CI= 0.35–2.54). The models adjusted for the child's age at the time of parents' enrollment. There was no adjustment for exposure to other pesticides.

Table 3.3. Summary of Findings: Solid Tumor Cancer Studies					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
All Cancers Combined					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.0 (0.9-1.2)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.0 (0.9-1.1) 1.0 (0.9-1.1)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 0.9 (0.8-1.0) 0.9 (0.8-1.1)	Age, demographic and lifestyle factors, and other pesticides ^b
			Lung		
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	0.9 (0.6-1.3)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 0.9 (0.5-1.5) 0.7 (0.4-1.2)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 1.1 (0.7-1.9) 0.6 (0.3-1.0)	Age, demographic and lifestyle factors, and other pesticides ^b
			Oral Cavity		
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.0 (0.5-1.8)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 0.8 (0.4-1.7) 0.8 (0.4-1.7)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5	1.0 1.1 (0.5-2.5)	Age, demographic and lifestyle factors, and other pesticides ^b

Table 3.3. Summary of Findings: Solid Tumor Cancer Studies					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
			79.6-337.1 337.2-18,241	1.0 (0.5-2.3)	
Kidney					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.6 (0.7-3.8)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 0.6 (0.3-1.4) 0.7 (0.3-1.6)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 0.3 (0.1-0.7) 0.5 (0.2-1.0)	Age, demographic and lifestyle factors, and other pesticides ^b
Bladder					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.5 (0.7-3.2)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.0 (0.5-1.9) 1.2 (0.6-2.2)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 0.5 (0.2-1.3) 0.8 (0.3-1.8)	Age, demographic and lifestyle factors, and other pesticides ^b
Melanoma					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.6 (0.8-3.0)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.2 (0.7-2.3) 0.9 (0.5-1.8)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days		Age, demographic and lifestyle factors, and other pesticides ^b

Table 3.3. Summary of Findings: Solid Tumor Cancer Studies					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
			(by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 0.6 (0.3-1.1) 0.7 (0.3-1.2)	
<i>Colon</i>					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.4 (0.8-2.2)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.4 (0.9-2.4) 0.9 (0.4-1.7)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 0.8 (0.5-1.5) 1.4 (0.8-2.5)	Age, demographic and lifestyle factors, and other pesticides ^b
Lee <i>et al.</i> (2007)	Nested Case-Control	USA: Iowa and North Carolina	Ever/never	1.0 (0.7-1.5)	Age, smoking, state, total days of pesticide application
<i>Rectum</i>					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.3 (0.7-2.3)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.3 (0.7-2.5) 1.1 (0.6-2.3)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 1.0 (0.5-2.0) 0.9 (0.5-1.9)	Age, demographic and lifestyle factors, and other pesticides ^b
Lee <i>et al.</i> (2007)	Nested Case-Control	USA: Iowa and North Carolina	Ever/never	1.6 (0.9-2.9)	Age, smoking, state, total days of pesticide application
<i>Colorectal</i>					
Lee <i>et al.</i> (2007)	Nested Case-Control	USA: Iowa and North Carolina	Ever/never	1.2 (0.9-1.6)	Age, smoking, state, total days of pesticide application
<i>Pancreas</i>					

Table 3.3. Summary of Findings: Solid Tumor Cancer Studies					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	0.7 (0.3-2.0)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.6 (0.6-4.1) 1.3 (0.5-3.6)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 2.5 (1.0-6.3) 0.5 (0.1-1.9)	Age, demographic and lifestyle factors, and other pesticides ^b
Andreotti <i>et al.</i> (2009)	Nested Case-Control	USA: Iowa and North Carolina	Ever/never	1.1 (0.6-1.7)	Age group, cigarette smoking, diabetes, and applicator type
			Intensity-Weighted Exposure Days (by control median): ≤184 ≥185	1.4 (0.9-3.8) 0.5 (0.2-1.3)	Age group, cigarette smoking, and diabetes
Prostate					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.1 (0.9-1.3)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 0.9 (0.7-1.1) 1.1 (0.9-1.3)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 1.0 (0.8-1.2) 1.1 (0.9-1.3)	Age, demographic and lifestyle factors, and other pesticides ^b
Koutros <i>et al.</i> (2013) ^c	Nested Case-Control	USA: Iowa and North Carolina	Intensity-Weighted Cumulative Exposure Days (by quartile): Q1 Q2 Q3 Q4	Total prostate cancer: 0.91 (0.79-1.06) 0.96 (0.83-1.12) 1.01 (0.87-1.17) 0.99 (0.86-1.15)	Age, state, race, smoking, fruit servings, family history of prostate cancer, and leisure time physical activity in the winter

Table 3.3. Summary of Findings: Solid Tumor Cancer Studies					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
			Intensity-Weighted Cumulative Exposure Days (by quartile): Q1 Q2 Q3 Q4	Aggressive prostate cancer: 0.93 (0.74-1.16) 0.91 (0.73-1.13) 1.01 (0.82-1.25) 0.94 (0.75-1.18)	Age, state, race, smoking, fruit servings, family history of prostate cancer, and leisure time physical activity in the winter
Band <i>et al.</i> (2011)	Case-Control	Canada: British Columbia	Ever/never	1.36 (0.83-2.25)	Alcohol consumption, cigarette years, education level, pipe years, and respondent type
<i>Esophagus</i>					
Lee <i>et al.</i> (2004b)	Case-Control	USA: Nebraska	Ever/never	0.7 (0.3-1.4)	Age and sex
<i>Stomach</i>					
Lee <i>et al.</i> (2004b)	Case-Control	USA: Nebraska	Ever/never	0.8 (0.4-1.5)	Age and sex
<i>Breast</i>					
Engel <i>et al.</i> (2005)	Nested Case-Control	USA: Iowa and North Carolina	Ever/never	Wives who apply pesticides: 0.9 (0.7-1.1) Wives who never used pesticides: 1.3 (0.8-1.9)	Age, race, and state of residence
<i>Soft Tissue Sarcoma</i>					
Pahwa <i>et al.</i> (2011)	Case-Control	Canada	Ever/never	0.90 (0.58-1.40)	Age group, province of residence, and statistically significant medical history variables
<i>Brain (glioma)</i>					
Lee <i>et al.</i> (2005)	Case-Control	USA: Nebraska	Ever/never	Overall: 1.5 (0.7-3.1) Self-reported: 0.4 (0.1-1.6) Proxy respondents: 3.1 (1.2-8.2)	Age for overall analysis; age and respondent type for other analyses
Yiin <i>et al.</i> (2012)	Case-Control	USA: Iowa, Michigan, Minnesota, and Wisconsin	Ever/never	House/garden use: 0.98 (0.67-1.43) Non-farm jobs: 0.83 (0.39-1.73)	Age, education, sex, and use of other pesticides
<i>Total Childhood</i>					

Table 3.3. Summary of Findings: Solid Tumor Cancer Studies					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
Flower <i>et al.</i> (2004)	Nested Case-Control	USA: Iowa and North Carolina	Ever/never	Maternal use: 0.61 (0.32-1.16) Paternal use: 0.84 (0.35-2.34)	Child's age at enrollment

^a Some studies report multiple quantitative risk measurements. This table reports the most highly adjusted quantitative measurements.

^b De Roos *et al.* (2005) excluded subjects missing covariate data for demographic and lifestyle factors and exposure to other pesticides; therefore, the number of subjects included in each analysis varies.

^c Effect estimates for glyphosate reported in the supplemental web material for Koutros *et al.* (2013).

3.5.2 Non-Solid Tumor Cancer Studies

(1) Leukemia

De Roos *et al.* (2005) reported no association between leukemia and glyphosate-exposed (ever/never used) pesticide applicators in the AHS cohort. For applicators with the full data set (54,315), the RR was 1.1 (95% CI=0.6–2.4) with only adjustment for age. In the fully adjusted model, the RR was similar (RR=1.0; 95% CI=0.5–1.9). The number of participants included in the adjusted analysis was lower (n=40,716) due to the exclusion of subjects with missing covariate data. Effect estimates using cumulative lifetime exposure and intensity-weighted cumulative exposure were also found to be non-statistically significant and did not demonstrate a trend with increasing exposure.

In a population-based case-control study in Iowa and Minnesota, Brown *et al.* (1990) did not observe an association with the ever-use of glyphosate (OR=0.9; 95% CI=0.5–1.6). A limitation in the study was the low number of cases exposed to glyphosate (n=15). Adjustments were made for several covariates, including vital status, age, tobacco use, family history of lymphopoietic cancer, high risk occupations, and high risk exposures; however, no adjustment was made for exposure to other pesticides.

Chang and Delzell (2016) conducted a meta-analysis exploring glyphosate exposure and leukemia using 3 studies (De Roos *et al.*, 2005; Brown *et al.*, 1990; and Kaufman *et al.*, 2009). I^2 values were reported, which represented the percentage of the total variance explained by study heterogeneity and measure inconsistency in results. Larger I^2 values indicate greater inconsistency. A meta-risk ratio of 1.0 (95% CI=0.6-1.5) was obtained with an I^2 value of 0.0%, indicating consistency across the data sets. It should be noted that this analysis included data from Kaufman *et al.* (2009), which is not considered in the current evaluation because it was assigned a low quality ranking because a quantitative measure of an association between glyphosate and a cancer outcome was not reported for that study.

(2) Multiple Myeloma

In a follow-up analysis of the study population from Iowa and Minnesota used in Brown *et al.* (1990), Brown *et al.* (1993) investigated whether pesticide use was related to multiple myeloma. Among men in Iowa, the authors observed a non-statistically significant elevated association with glyphosate use (OR=1.7; 95% CI=0.8–3.6; 11 exposed cases); however, no adjustment was made for exposure to other pesticides. The authors cautioned that while the study may lend support to the role of pesticides in general, the study limitations preclude use of the evidence as a definitive finding for any one compound.

De Roos *et al.* (2005) reported a suggestive association between multiple myeloma and glyphosate-exposed pesticide applicators based on 32 multiple myeloma cases observed in the AHS cohort. For applicators with the full data set, the RR was 1.1 (95% CI=0.5–2.4) with only adjustment for age. In the fully adjusted model excluding subjects with missing covariate data, there was a non-statistically significant elevated risk following adjustment for age, demographic and lifestyle factors, and exposure to other pesticides (RR=2.6; 95% CI=0.7–9.4). The authors

postulated that the increased myeloma risk could be due to bias resulting from a selection of subjects in adjusted analyses that differed from subjects included in unadjusted analyses or may be due to a confounder or effect modifier that is prevalent among the subgroup and has not been accounted for in the analyses. When exposure data were also stratified by tertiles with the lowest tertile of exposure as the referent category, trend analyses were not statistically significant. Non-statistically significant elevated RRs of 1.9 (95% CI: 0.6-6.3) and 2.1 (95% CI: 0.6-7.0) were estimated for the highest tertile of both cumulative and intensity-weighted exposure days, respectively. The study authors did note that small sample size precluded precise estimation (n=19 for adjusted analyses). When using never exposed as the referent category, the trend analysis was again non-statistically significant, but the RRs ranged from 2.3 (95% CI: 0.6-8.9) to 4.4 (95% CI: 1.0-20.2) from the lowest tertile to the highest tertile, respectively. When stratified by quartiles, a statistically significant trend is achieved and the RR increased to 6.6 (95% CI: 1.4-30.6); however, the authors noted that the cases were sparsely distributed for these analyses.

Sorahan (2015)¹¹ re-analyzed the AHS data reported by De Roos *et al.* (2005) to examine the reason for the disparate findings in relation to the use of a full data set versus the restricted data set. Using Poisson regression, risk ratios were calculated without excluding subjects with missing covariate data. When adjusted for age and sex, the RR for ever-use of glyphosate was 1.12 (95% CI of 0.5–2.49). Additional adjustment for lifestyle factors and use of other pesticides did not have a large impact (RR=1.24; 95% CI=0.52–2.94). The authors concluded that the disparate findings in De Roos *et al.* (2005) could be attributed to the use of a restricted dataset that was unrepresentative.

Landgren *et al.* (2009), within the AHS study population, also investigated the association between pesticide use and prevalence of monoclonal gammopathy of undetermined significance (MGUS). MGUS is considered a pre-clinical marker of multiple myeloma progression. The authors did not observe an association with glyphosate use and MGUS using subjects from the AHS cohort (OR=0.50; 95% CI=0.20–1.0). No adjustment was made for exposure to other pesticides.

In a population-based case-control study (Pahwa *et al.*, 2012) among men in six Canadian provinces, a non-statistically significant elevated odds of multiple myeloma was reported in relation to glyphosate use (OR=1.22; 95% CI = 0.77–1.93), based upon 32 glyphosate exposed multiple myeloma cases and 133 controls. There was no adjustment for exposure to other pesticides. Kachuri *et al.* (2013), using the same Canadian study population, further explored multiple myeloma in relation to days per year that glyphosate was used. Adjustment for exposure to other pesticides was also not performed in this study. For ever-use, there was a slight non-statistically significant increased odds ratio (OR=1.19; 95% CI=0.76–1.87). For light users (>0 and ≤2 days/year), there was no association (OR=0.72; 95% CI = 0.39–1.32; 15 exposed cases); whereas, for heavy users (>2 days/ year), there was a non-statistically significant increased odds ratio (OR=2.04; 95% CI=0.98–4.23; 12 exposed cases). Similar results were obtained when proxy respondents were excluded from the analysis. The low number of cases and controls exposed to glyphosate, particularly when exposed subjects were divided into light and heavy users, was a limitation of the study. It would be expected that effect estimates would be reduced if adjustment for co-exposure to other pesticides had been performed.

¹¹ Funded by Monsanto

In a hospital-based case-control study conducted by Orsi *et al.* (2009) in France, 56 multiple myeloma cases and 313 age- and sex-matched controls were identified. A non-statistically significant elevated risk was observed (OR=2.4; 95% CI=0.8–7.3; 5 exposed cases and 18 exposed controls). The wide CI range can primarily be attributed to the low number of exposed cases indicating the analysis is underpowered. Additionally, the study did not adjust for exposure to multiple pesticides.

Chang and Delzell (2016) conducted a meta-analysis exploring glyphosate exposure and multiple myeloma using data from the 6 studies described above (Brown *et al.*, 1993; De Roos *et al.*, 2005; Sorahan, 2015; Pahwa *et al.*, 2012; Kachuri *et al.*, 2013; Orsi *et al.*, 2009). Meta-risk ratios were obtained using data from each of the 4 independent study populations, such that if a study population was already represented in the analysis by one study, then the same population analyzed by another study would not be included (e.g., Sorahan, 2015 and De Roos *et al.*, 2005 could not be used simultaneously in a meta-analysis). The combined meta-risk ratio based on data from prioritized studies (Brown *et al.*, 1993; Kachuri *et al.*, 2013; Orsi *et al.*, 2009; and Sorahan, 2015) was 1.4 (95% CI=1.0-1.9) using random-effects and fixed-effects models and the I^2 value = 0.0% indicating consistency across data sets. There was relatively no impact on the meta-risk ratio and associated 95% CI when secondary analyses were conducted using alternative estimates for a study population (e.g., substituting the data from Sorahan, 2015 for De Roos *et al.*, 2005).

(3) Hodgkin Lymphoma

In a Canadian case-control study, Karunanayake *et al.*, (2012) evaluated Hodgkin lymphoma (HL) and observed no association with glyphosate exposure following adjustment for age, province of residence, and medical history variables (OR=0.99; 95% CI=0.62-1.56; 38 cases). No adjustment was made for exposure to other pesticides.

In a hospital-based case-control study conducted by Orsi *et al.* (2009) in France, authors identified 87 HL cases and 265 age-and sex-matched controls. There was a non-statistically significant elevated odds ratio observed (OR=1.7; 95% CI=0.6–5.0; 6 exposed cases and 15 exposed controls). The wide CI range can primarily be attributed to the low number of exposed cases indicating the analysis is underpowered. Also, as noted earlier, this study did not adjust for exposure to multiple pesticides.

Chang and Delzell (2016) conducted a meta-analysis exploring glyphosate exposure and HL using data from both of these studies. A meta-risk ratio of 1.1 (95% CI=0.7-1.6) was obtained with a I^2 value of 0.0%, indicating consistency across the data sets.

(4) Non-Hodgkin Lymphoma

NHL has about 60 subtypes classified by the WHO, which may have etiological differences (Morton *et al.*, 2014). There are analyses available for particular subtypes of NHL; however, these are particularly limited by the small sample sizes. As a result, this evaluation only presents results for total NHL.

There were six studies available that investigated the association between glyphosate exposure and NHL, which was the most for any type of cancer. As discussed in Section 3.4, these studies encompass a combination of strengths and limitations. These studies are therefore discussed in more detail in this section as compared to discussions of other cancer types in order to highlight the strengths and identify the limitations for each study.

De Roos *et al.* (2005) was the only prospective cohort study available; therefore, subjects were enrolled prior to developing cancer outcomes. Disease status was determined through state cancer registries. Exposure information was obtained from a large number of licensed pesticide applicators and no proxies were used. Exposure was evaluated as ever/never use, cumulative lifetime exposure, and intensity-weighted cumulative exposure. Due to the study design, the potential for many biases were reduced. Additionally, the study adjusted and/or considered numerous factors, including use of other pesticides. Median follow-up time was approximately 7 years and a longer follow-up would increase the ability of the study to detect subjects developing cancer outcomes; however, as discussed in Section 3.3.1, study participants provided exposure information prior to enrollment and this information was incorporated into the cumulative lifetime and intensity-weighted cumulative exposure metrics. As a result, the amount of time exposed was longer than just the follow-up time since enrollment. For applicators with the full data set, the RR for ever/never use was 1.2 (95% CI=0.7–1.9; 92 cases) with only adjustment for age. In the fully adjusted model excluding subjects with missing covariate data, the RR was similar following adjustment for age, demographic and lifestyle factors, and exposure to other pesticides (RR=1.1; 95% CI=0.7-1.9). Effect estimates obtained using cumulative lifetime exposure and intensity-weighted cumulative exposure were below 1 (RR = 0.6-0.9 when comparing to the lowest tertile).

De Roos *et al.* (2003) used pooled data from three case-controls studies evaluating NHL in white males from Nebraska, Kansas, and in Iowa and Minnesota (Cantor *et al.*, 1992; Hoar *et al.*, 1986; Zahm *et al.*, 1990; Appendix B). Exposure information was obtained from exposed individuals or their next of kin (i.e., proxy respondents) if the subjects were dead or incapacitated; however, techniques varied across the three studies. There is potential for selection bias due to exclusion of observations with missing covariate data, but only if the lack of the covariate data was associated with glyphosate exposure. The effect estimates for the association between glyphosate exposure and NHL was significant (OR=2.1; 95% CI=1.1–4.0) in the logistic regression analyses controlling for co-exposure to other pesticides. However, utilizing alternative hierarchical regression techniques to adjust for co-exposure to other pesticide exposures, the odds ratio was still elevated, but the increase was not statistically significant (OR=1.6; 95% CI=0.90–2.8).

Eriksson *et al.* (2008) is a Swedish case-control study that used detailed exposure information from exposed individuals (i.e., no use of proxy respondents), but only minimal demographic information was provided on subjects (age and sex) and a table with subject characteristics (e.g., smoking status, alcohol intake, physical activity, education) was not provided. Cases were identified through physicians and verified histopathologically. Glyphosate exposure, which was reported in 29 cases and 18 controls between 1999 and 2003, produced a statistically significant increased OR in the univariate analysis (OR=2.02; 95% CI=1.10–3.71); however, in the

multivariate analysis adjustments were conducted for co-exposure to different agents including MCPA, “2,4,5-Y and/or 2,4-D”, mercurial seed dressing, arsenic, creosote, and tar and the OR reduced to 1.51 (95% CI=0.77–2.94) and was not statistically significant. Additional analyses were conducted to investigate the impact of various exposure times. When exposure was for more than 10 cumulative days (the median number of days among exposed controls), the OR was 2.36 (95% CI=1.04–5.37; 17 exposed cases) and for exposure less than 10 cumulative days, the OR was 1.69 (95% CI=0.7–4.07; 12 exposed cases). By dividing the exposed cases and controls using this exposure metric, wider CIs were observed indicating reduced power from the smaller sample sizes. Additionally, these analyses did not account for co-exposure to other pesticides. Similarly, wider CIs were also observed when exposed cases and controls were divided by a longer exposure metric. ORs of 1.11 (95% CI=0.24-5.08) and 2.26 (95% CI=1.16-4.40) were obtained for 1-10 years and >10 years, respectively. It was not clear whether this analysis controlled for co-exposure to other pesticides based on the statistical methods description and the subjects for each exposure group were not reported. This finding, while limited to a single study, suggests that cohort studies without sufficient follow-up time or other case-control studies which did not stratify by time since first exposure may be less sensitive in detecting risk.

Hardell *et al.* (2002) used pooled data from two case-control studies in Sweden (Hardell and Eriksson, 1999; Nordstrom *et al.*, 1998; Appendix B) that examined hairy cell leukemia, a subtype of NHL, and NHL (not including hairy cell leukemia). Exposure information was collected from individuals or proxy respondents based on a working history with specific questions on exposures to different chemicals. Cases were identified from regional cancer registries and verified histopathologically. In the univariate analysis, risk of NHL associated with glyphosate exposure was found to be significantly increased (OR=3.04; 95% CI=1.08–8.52), but when study site, vital status, and co-exposure to other pesticides were considered in the multivariate analysis, the OR noticeably attenuated and was found to be non-statistically significant (OR=1.85; 95% CI=0.55–6.20). The wide range of the CI suggests that the analysis is underpowered (only 8 glyphosate-exposed cases and 8 glyphosate-controls).

McDuffie *et al.* (2001) is a multicenter population-based study among men of six Canadian provinces. This case-control study utilized a well-conducted exposure assessment and cases were ascertained from cancer registries or hospitals in six provinces with histopathological verification for 84% of the samples. There are concerns with control selection. There was low control participation (48%) and different sources were used for selecting controls depending on the province of residence. Effect estimates were obtained using a considerable number of exposed cases and controls (51 cases and 133 controls); however, the study did not assess co-exposure to other pesticides. There was a non-statistically significant increased risk of NHL from glyphosate exposure when adjusting for age and province (OR=1.26; 95% CI=0.87–1.80) and when adjusting for age, province and medical variables (OR=1.20; 95% CI=0.83–1.74). Medical variables found to be statistically significant included history of measles, mumps, previous cancer, skin-prick allergy tests, allergy desensitization shots, and a positive family history of cancer in a first-degree relative. It would be expected that effect estimates would attenuate if control for co-exposure to other pesticides had been performed. Additional analyses were conducted to investigate differences in exposure time. When exposure was for more than 2 days/year, the OR was 2.12 (95% CI=1.20-3.73; 23 exposed cases and 36 exposed controls) compared to unexposed subjects and for exposure more than 0 and ≤ 2 days/year, the OR was

1.00 (95% CI=0.63–1.57; 28 exposed cases and 97 exposed controls) compared to unexposed subjects.

Orsi *et al.* (2009) is a French hospital-based case-control study that obtained exposure information from subjects (no proxies used) using a detailed questionnaire with lifelong residential and occupational histories followed by a discussion with a trained interviewer who was blinded to case status. No issues regarding exposure or outcome assessment were identified; however, there is potential for selection bias given the study utilized hospital-based controls. The study evaluated several potential confounders; however, it did not assess co-exposure to other pesticides. There was no association observed between NHL and glyphosate use (OR=1.0; 95% CI=0.5-2.2; 12 exposed cases and 24 exposed controls). The low number of cases and controls exposed to glyphosate and lack of adjustment for exposure to multiple pesticides were limitations of the study.

Schinasi and Leon (2014) conducted a meta-analysis exploring occupational glyphosate exposure and NHL using data from six of the above mentioned studies (McDuffie *et al.*, 2001; Hardell *et al.*, 2002; De Roos *et al.*, 2003; De Roos *et al.*, 2005; Eriksson *et al.*, 2008; and Orsi *et al.*, 2009). Since the authors identified a variety of sources of heterogeneity between publications, they decided a priori to calculate meta-risk ratio estimates and 95% CIs using random effect models, allowing between study heterogeneity to contribute to the variance. I^2 values were reported as a measure of inconsistency in results. For glyphosate, the meta-risk ratio was 1.5 with a 95% CI of 1.1–2.0 and the I^2 value was 32.7% indicating relatively low levels of heterogeneity among these studies. This study combined multiple smaller studies that on their own were very limited in statistical power.

The 2015 IARC evaluation noted that fully adjusted effect estimates in two of the Swedish studies (Hardell *et al.*, 2002 and Eriksson *et al.*, 2008) were not used in the analysis conducted by Schinasi and Leon (2014). Consequently, the IARC Working Group conducted a reexamination of the results of these studies (IARC 2015). For an association between glyphosate exposure and NHL, the IARC estimated a meta-risk ratio of 1.3 (95% CI=1.03–1.65, $I^2=0\%$; $p=0.589$ for heterogeneity).

Chang and Delzell (2016) conducted their own meta-analysis exploring glyphosate exposure and NHL using six independent studies (De Roos *et al.*, 2003; De Roos *et al.*, 2005; Eriksson *et al.*, 2008; Hardell *et al.*, 2002; McDuffie *et al.*, 2001; and Orsi *et al.*, 2009). A meta-risk ratio of 1.3 (95% CI=1.0-1.6) was obtained with an I^2 value of 0.0%. In a secondary analysis, the De Roos *et al.* (2003) OR using hierarchical regression was replaced by the logistic regression OR. This change had no impact on the meta-risk ratio and associated confidence interval (meta-risk ratio=1.3; 95% CI=1.0-1.6). In another secondary analysis, the OR from McDuffie *et al.* (2001) was replaced by the OR from Hohenadel *et al.* (2011), which evaluated the same study population (minus four previously misclassified NHL cases). This analysis also yielded similar results (meta-risk ratio=1.3; 95% CI=1.0-1.7). A final analysis was performed with the replacements for both secondary analyses [i.e., logistic regression OR from De Roos *et al.* (2003) and OR from Hohenadel *et al.* (2011)]. The results were relatively the same as the other meta-analyses (meta-risk ratio=1.4; 95% CI=1.0-1.8). Chang and Delzell (2016) also tested for publication bias using Egger's linear regression approach to evaluating funnel plot asymmetry,

and found no significant asymmetry indicating little evidence of publication bias; however, given the small sample size ($n=6$), this analysis would lack power and the results are not considered meaningful.

Table 3.4. Summary of Findings: Non-Solid Tumor Cancer Studies.					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
<i>Leukemia</i>					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.0 (0.5-1.9)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.9 (0.8-4.5) 1.0 (0.4-2.9)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 1.9 (0.8-4.7) 0.7 (0.2-2.1)	Age, demographic and lifestyle factors, and other pesticides ^b
Brown <i>et al.</i> (1990)	Case-Control	USA: Iowa and Minnesota	Ever/never	0.9 (0.5-1.6)	Vital status, age, tobacco use, family history of lymphopoietic cancer, high occupations, and high risk exposures
<i>Multiple Myeloma</i>					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	2.6 (0.7-9.4)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 1.1 (0.4-3.5) 1.9 (0.6-6.3)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 1.2 (0.4-3.8) 2.1 (0.6-7.0)	Age, demographic and lifestyle factors, and other pesticides ^b
Brown <i>et al.</i> (1993)	Case-Control	USA: Iowa	Ever/never	1.7 (0.8-3.6)	Age and vital status
Kachuri <i>et al.</i> (2013)	Case-Control	Canada	Ever/never	1.19 (0.76-1.87)	Age, province of residence, smoking status, selected medical conditions, family history of cancer, and use of a proxy respondent
			Days per year of use: 0 to ≤2 days/year >2 days/year	0.72 (0.39-1.32) 2.04 (0.98-4.23)	Age, province of residence, smoking status, selected medical conditions, family history of cancer, and use of a proxy respondent
Pahwa <i>et al.</i> (2012)	Case-Control	Canada	Ever/never	1.22 (0.77-1.93)	Age group, province of residence, and statistically significant medical history variables

Table 3.4. Summary of Findings: Non-Solid Tumor Cancer Studies.					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
Orsi <i>et al.</i> (2009)	Case-Control	France	Ever/never	2.4 (0.8-7.3)	Age, centre, and socioeconomic category
<i>Monoclonal Gammopathy of Undetermined Significance (MGUS)</i>					
Landgren <i>et al.</i> (2009)	Nested Case-Control	USA: Iowa and North Carolina	Ever/never	0.5 (0.2-1.0)	Age and education
<i>Hodgkin Lymphoma (HL)</i>					
Karunanayake <i>et al.</i> (2012)	Case-Control	Canada	Ever/never	0.99 (0.62-1.56)	Age group, province of residence, and statistically significant medical history variables
Orsi <i>et al.</i> (2009)	Case-Control	France	Ever/never	1.7 (0.6-5.0)	Age, centre, and socioeconomic category
<i>Non-Hodgkin Lymphoma (NHL)</i>					
De Roos <i>et al.</i> (2005)	Prospective Cohort	USA: Iowa and North Carolina	Ever/never	1.1 (0.7-1.9)	Age, demographic and lifestyle factors, and other pesticides ^b
			Cumulative Exposure Days (by tertile cut points): 1-20 21-56 57-2,678	1.0 0.7 (0.4-1.4) 0.9 (0.5-1.6)	Age, demographic and lifestyle factors, and other pesticides ^b
			Intensity-Weighted Cumulative Exposure Days (by tertile cut points): 0.1-79.5 79.6-337.1 337.2-18,241	1.0 0.6 (0.3-1.1) 0.8 (0.5-1.4)	Age, demographic and lifestyle factors, and other pesticides ^b
De Roos <i>et al.</i> (2003)	Case-Control	USA: Iowa, Nebraska, Minnesota, and Kansas	Ever/never	1.6 (0.9-2.8)	Age, study site, and use of other pesticides
Eriksson <i>et al.</i> (2008)	Case-Control	Sweden	Ever/never	Multivariate: 1.51 (0.77-2.94)	Age, sex, year of diagnosis or enrollment, and exposure to other pesticides
			Days per year of use: ≤ 10 days >10 days	1.69 (0.70-4.07) 2.36 (1.04-5.37)	Age, sex, and year of diagnosis or enrollment
			Years of use: 1-10 years >10 years	1.11 (0.24-5.08) 2.26 (1.16-4.40)	Unknown
Hardell <i>et al.</i> (2002)	Case-Control	Sweden	Ever/never	Multivariate: 1.85 (0.55-6.20)	Study, study area, vital status, and exposure to other pesticides
McDuffie <i>et al.</i> (2001)	Case-Control	Canada	Ever/never	1.20 (0.83-1.74)	Age, province of residence, and statistically significant medical variables
			Days per year of use: >0 and ≤ 2 days	1.00 (0.63-1.57)	Age and province of residence

Table 3.4. Summary of Findings: Non-Solid Tumor Cancer Studies.					
Study	Study Design	Study Location	Exposure Metric	Adjusted Effect Estimate: RR or OR (95% CI) ^a	Covariate Adjustments in Analyses
			>2 days	2.12 (1.20 -3.73)	
Orsi <i>et al.</i> (2009)	Case-Control	France	Ever/never	1.0 (0.5-2.2)	Age, centre, and socioeconomic category

^a Some studies report multiple quantitative risk measurements. This table reports the most highly adjusted quantitative measurements.

^b De Roos *et al.* (2005) excluded subjects missing covariate data for demographic and lifestyle factors and exposure to other pesticides; therefore, the number of subjects included in each analysis varies.

3.6 Discussion

A total of 24 epidemiological studies from the open literature were identified as appropriate for detailed evaluation. Of these, 23 studies were considered informative with regard to the carcinogenic potential of glyphosate. There was no evidence of an association between glyphosate exposure and solid tumors. There was also no evidence of an association between glyphosate exposure and leukemia, or HL. This conclusion is consistent with those recently conducted by IARC, EFSA, and JMPR who also concluded there is no evidence of an association for these tumors at this time. The data should be considered limited though with only one or two studies available for almost all of the cancer types investigated. Additionally, with the increased use of glyphosate following the introduction of glyphosate-tolerant crops in 1996, there is a need for more recent studies since a large number of studies were conducted prior to 1996. As described in Section 1.1, the use pattern changed following the introduction of transgenic crops, which may impact overall effect estimates. The remainder of this discussion focuses on multiple myeloma and NHL. Study elements for the available studies and their potential to impact effect estimates are examined; however, the discussion is applicable in most cases to all of the epidemiological studies used in this evaluation.

Multiple Myeloma

Five studies were available evaluating the association between glyphosate exposure and risk of multiple myeloma (Brown et al., 1993; De Roos et al., 2005; Kachuri et al., 2013; Orsi et al., 2009; Pahwa et al., 2012). The effect estimates for ever/never use ranged from 1.19 to 2.6 although none were found to be statistically significant. Only one study (De Roos et al., 2005) controlled for co-exposures to other pesticides; therefore, potential confounding was not addressed in the other studies. There was an indication of a possible exposure-response relationship; however, this was the only study that evaluated the exposure-response relationship for multiple myeloma. Furthermore, reanalysis of the full dataset by Sorahan (2015) raised concerns about whether the restricted dataset used for these analyses was representative of the whole cohort. There was a single study of MGUS, a precursor to multiple myeloma, which showed decreased risk with exposure to glyphosate; however, the study did not control for exposure to other pesticides. Overall, the available epidemiologic evidence for an association between glyphosate and risk of multiple myeloma is inadequate to assess the carcinogenic potential at this time due to the potential for confounding in three of the four studies, the limited observation of a possible exposure-response relationship in a single study, and concerns whether restricted datasets were representative of the whole cohort.

NHL

Six studies were available evaluating the association between glyphosate exposure and risk of NHL. Effect estimates for ever/never use ranged from 1.0-1.85 in adjusted analyses with none reaching statistical significance (Figure 3.2). Two of these studies did not adjust for co-exposures to other pesticides (McDuffie et al., 2001; Orsi et al., 2009). Many of the evaluated studies had limited power due to small sample sizes, which resulted in large confidence intervals and reduced the reliability of the results to demonstrate a true association. Meta-analyses were performed by IARC (2015) and Chang and Delzell (2016) using these results for the ever/never

use metric. Both analyses reported similar meta-risk ratios ranging from 1.3-1.5, depending on the effect estimates and studies included in the analyses. All meta-analysis estimates reported were non-statistically significant except the meta-risk ratio reported by IARC (2015), which was borderline significant with the lower limit of the 95% CI at 1.03. It should also be noted that publication bias may play a role in this evaluation given there is a tendency to only publish positive results and potential concerns regarding glyphosate have only been raised in recent years.

With respect to meta-analyses, caution should be taken when interpreting results. Meta-analyses are a systematic way to combine data from several studies to estimate a summary effect. Analyses were performed with 6 studies, which many would consider small for performing meta-analyses. Rarely will meta-analyses synthesize data from studies with identical study designs and methods. In the meta-analyses performed by IARC (2015) and Chang and Delzell (2016), inclusion was primarily based on whether a study addressed the broader question regarding the association between glyphosate exposure and risk of NHL. For meaningful results, careful consideration of whether studies are similar and should be combined in the analysis. Furthermore, the bias and confounding issues inherent for each individual study are carried over into the meta-analyses. Across the NHL studies, study characteristics varied, such as overall study design (i.e., cohort and case-control), source population, proxy respondent use, covariate adjustments, and confounding control. Even if these differences are not detected statistically, the meta-analysis estimate should be considered in the context of the data that are used to generate it.

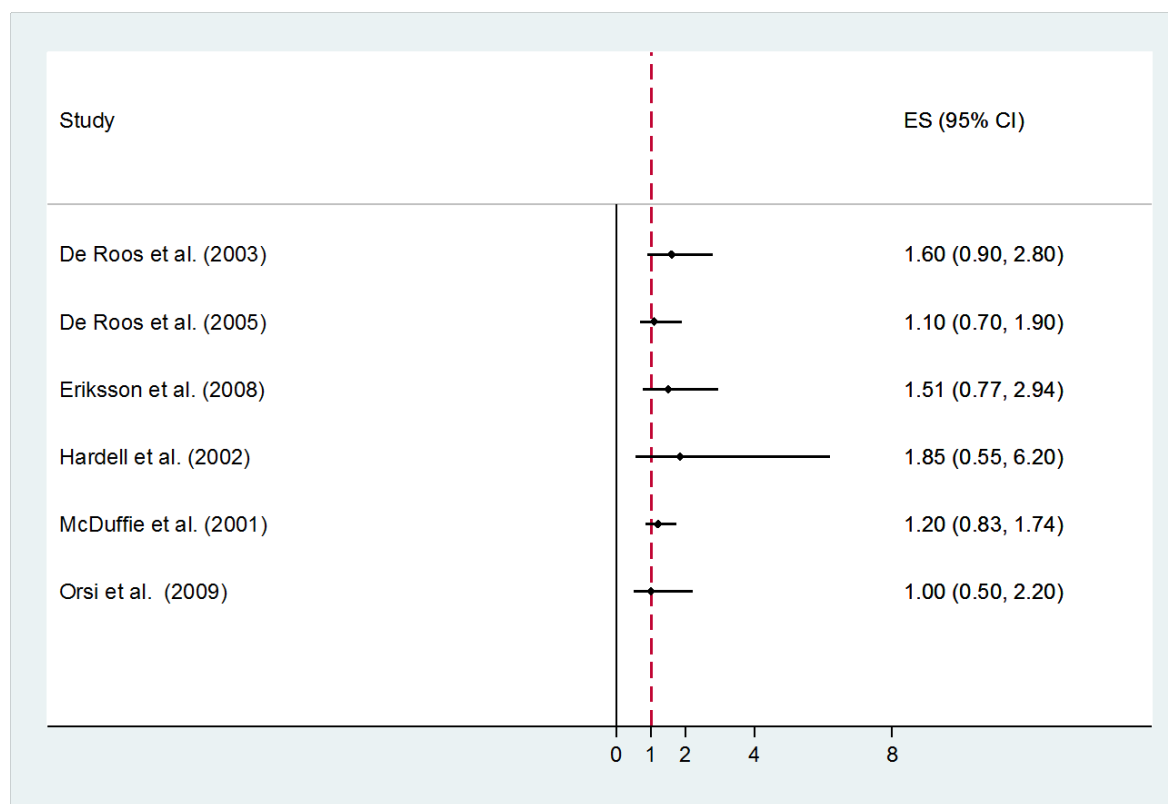


Figure 3.2. Forest plot of effect estimates (denoted as ES for effect sizes) and associated 95% confidence intervals (CI) for Non-Hodgkin lymphoma (NHL).

Using cumulative lifetime and intensity-weighted cumulative exposure metrics, all effect estimates were less than 1 (OR = 0.6-0.9 when comparing to the lowest tertile) in the AHS cohort study (De Roos *et al.*, 2005). Two case-control studies (Eriksson *et al.*, 2008; McDuffie *et al.*, 2001) evaluated the association of glyphosate exposure and NHL stratifying exposure by days per year of use. These studies obtained effect estimates greater than 1, which conflicted with the results in the prospective cohort study; however, these estimates from the case-control studies do not appear to be adjusted for co-exposures to other pesticides. As mentioned previously (and will be discussed further below), there was clearly strong potential for confounding from exposure to other pesticides. In each instance where a study controlled for co-exposure to other pesticides, the adjusted effect estimate decreased in magnitude, including other analyses performed in one of these case-control studies. Consequently, lack of adjustment for co-exposure to other pesticides in these analyses could partially explain the conflicting results between the cohort and case-control studies.

The possible effect of confounding factors, which are related to both the exposure of interest and the risk of disease, may make it difficult to interpret the results. Control for confounding varied considerably across studies (Table 3.2). Studies primarily adjusted for standard variables, such as age, gender, and residency location. Co-exposure to other pesticides was considered for several of the NHL studies for ever/never use (De Roos *et al.*, 2003; De Roos *et al.*, 2005; Eriksson *et al.*, 2008; Hardell *et al.*, 2002); however, analyses of exposure-response and latency effects did not appear to control for these co-exposures.

There is clearly a strong potential for confounding by co-exposures to other pesticides since many are highly correlated and have been reported to be risk factors for NHL. In the studies that did report a quantitative measure adjusted for the use of other pesticides, the risk was always found to be closer to the null than the risk calculated prior to this adjustment. For examples, Eriksson *et al.* (2008) reported unadjusted and adjusted effect estimates of 2.02 (95% CI: 1.10-3.71) and 1.51 (95% CI: 0.77-2.94), respectively. Comparing the magnitude of those effect sizes on the natural log scale, the unadjusted effect was $\beta=0.70$ (95% CI: 0.10, 1.31) while the adjusted effect was $\beta=0.41$ (95% CI: -0.26, 1.08), suggesting a difference compatible with a degree of confounding by those herbicide co-exposures which appeared to have inflated the unadjusted effect upwards by 70% on the natural log scale (or by 46% on the OR scale). This demonstrates the profound effect this adjustment has on effect estimates and the concern for residual confounding by other pesticides that cause NHL themselves. As discussed in Section 3.2.4, other potential confounders have also been identified. With an association between glyphosate exposure and the outcome of interest, occupational exposure to diesel exhaust fumes, solvents, and UV radiation are highly likely confounders in the NHL studies; however, none of these studies accounted for these potential confounders.

Recall bias and missing data are also limitations in most of the studies. In epidemiologic studies, the quality of the exposure assessment is a major concern since the validity of the evaluations depends in large part on the ability to correctly quantify and classify an individual's exposure. Variation in the quality of exposure assessment, study design and methods, as well as available information concerning potential confounding variables could also explain discrepancies in study findings. During their lifetime, farmers are typically exposed to multiple pesticides and often

several may be used together posing a challenge for identifying specific risk factors. Moreover, there is no direct information on pesticide exposure or absorbed dose because analyses are based on self-reported pesticide use. The studies included in this epidemiology assessment relied primarily on questionnaires and interviews to describe participants' past and/or current exposure to glyphosate. Since the questionnaires are commonly used to account for exposure and capture self-reporting, the results can be subject to misclassification and recall bias.

Furthermore, the use of proxy respondents has the potential to increase recall bias and thus may increase exposure misclassification, especially for proxy respondents not directly involved in farming operations that may be more prone to inaccurate responses than directly interviewed subjects. In some of the NHL studies, the study participants were interviewed directly to assess exposure (De Roos *et al.*, 2005; Eriksson *et al.*, 2008; McDuffie *et al.*, 2001; Orsi *et al.*, 2009), making proxy respondent use a non-issue for these studies. In other studies, however, study participants or proxy respondents were interviewed to assess exposure (Hardell *et al.*, 2002, De Roos *et al.*, 2003). De Roos *et al.* (2003) did not find type of respondent to be statistically significant, but Hardell *et al.* (2002) did not conduct analyses to evaluate the impact of proxy use. In non-NHL studies, proxy analyses were conducted in a small subset (Kachuri *et al.*, 2013; Lee *et al.*, 2004b; Lee *et al.*, 2005; Yiin *et al.*, 2012) and differences in effect estimates were often observed. In a few studies, respondent type was used as an adjustment variable when calculating effect estimates (Band *et al.*, 2011; Kachuri *et al.*, 2013; Lee *et al.*, 2005). As with all study design elements of case-control studies, one concern is whether or not the use of proxy respondents had a differential impact on the cases and controls included in the study because any differential impact may result in differential exposure misclassification. When use of proxy respondents was comparable for cases and controls in the full study population, it could be assumed that there is less concern for potential recall bias from the use of proxy respondents. In Hardell *et al.*, (2002), the percentage of cases and controls with proxy respondents was not fully reported for cases and controls though and this adds a potential source of uncertainty for the study. Moreover, when proxy respondents were used in a study, the percentages were usually reported only for the full study population and were not reported for the specific cases and controls exposed to glyphosate. This lack of information makes it difficult to assess the degree to which recall bias may have occurred due to the use of proxy respondents.

The highest risk measures were reported in studies with subjects developing NHL during a period of relatively low use of glyphosate. For example, Hardell *et al.* (2002) and De Roos *et al.* (2003) acquired cases from 1987-1990 and 1979-1986, respectively. These studies reported the largest adjusted ORs for glyphosate exposure and NHL (1.6 and 1.85); however, these studies investigated subjects prior to the introduction of genetically engineered glyphosate-tolerant crops. As discussed in Section 1.4, glyphosate use dramatically increased following the introduction of genetically engineered glyphosate-tolerant crops in 1996. Prevalence alone would not be expected to result in a corresponding increase in outcomes associated with glyphosate; however, the use pattern changed following the introduction of transgenic crops, such that in addition to new users, individuals already using glyphosate would have a corresponding increase in glyphosate exposure. As a result, if a true association exists between glyphosate exposure and NHL, then a corresponding increase in effect estimates would also be expected during this time. The currently available studies do not display this trend. In more recent years, including the AHS prospective cohort study (De Roos *et al.*, 2005), reported

adjusted risk measures were lower (1.0-1.51). Furthermore, if a true association exists, it would also be expected that the higher effect estimates would be reported in countries where individuals are more exposed to glyphosate, such as the United States and Canada, as compared to countries that exhibit less use¹². Once again, the expected trend was not observed, such that effect estimates for studies conducted in Sweden (Eriksson *et al.*, 2008; Hardell *et al.*, 2002), where glyphosate-tolerant crops are sparsely grown, were similar or higher than those reported in the United States (De Roos *et al.*, 2003; De Roos *et al.*, 2005) and Canada (McDuffie *et al.*, 2001). These counterintuitive results highlight the need for additional studies to determine the true association between glyphosate exposure and NHL, as well as further elucidate the exposure-response relationship.

Some have argued that the follow-up period (median = 7 years) in De Roos *et al.* (2005) is not sufficiently long to account for the latency of NHL (Portier *et al.*, 2016); however, the latency period for NHL following environmental exposures is relatively unknown and estimates have ranged from 1-25 years (Fontana *et al.*, 1998; Kato *et al.*, 2005; Weisenburger, 1992). Eriksson *et al.*, (2008) evaluated the impact of time since first exposure. This study found an increased effect estimate for subjects with more than 10 years of glyphosate exposure prior to diagnosis of NHL. This finding suggests a potential for a longer latency for NHL than the follow-up period in De Roos *et al.* (2005); however, this analysis did not appear to account for co-exposures to other pesticides and the number of subjects in the analysis were not reported. It should be noted that the follow-up time in De Roos *et al.* (2005) does not represent the amount of time subjects have been exposed. In this study, prior pesticide exposure was provided at time of enrollment and used to evaluate subjects that contribute person-time from enrollment until the point of diagnosis, death, movement from the catchment area, or loss to follow-up. As such, estimated exposure for each subject did not continue to accrue during follow-up. Additionally, subjects were not checked against state registries for inclusion in the cohort. Rather, cancer analyses were restricted to those who are cancer-free at the time of enrollment to remove any issues related to treatment that might impact subsequent cancer risk. At the time of enrollment, the average and median times of exposure 7.5 years and 8 years, respectively, with a standard deviation of 5.3¹³. These values were calculated using the midpoint of exposure categories provided in the questionnaire; therefore, these values represent a range of subject exposure time. Given the majority of the subjects were at least 40 years old at the time of analysis and the recognition that these workers generally start in their profession at a much earlier age and stay in that profession over their lifetime, time of exposure for many of these subjects would be greater than the average and median times. All of this information indicates that subjects within the cohort have ample amount of time for the outcome of interest to develop and be detected during the study. Furthermore, NHL has about 60 subtypes classified by the WHO, which may have etiological differences (Morton *et al.*, 2014). In this evaluation, the analysis of effect estimates was restricted to total NHL due to the small sample sizes in the few instances where NHL subtypes were analyzed. There are concerns with grouping the subtypes together despite etiological differences and the latency period for each NHL subtype may vary due to these etiological differences. Given the latency analysis was limited to Eriksson *et al.* (2008) and lack of NHL latency understanding in general, further analyses are needed to determine the true

¹² Components in glyphosate formulations in the United States and abroad are similar according to personal communication with Monsanto.

¹³ Information provided by email from NIEHS.

latency time of NHL and NHL subtypes. The next update to the AHS cohort study with a longer follow-up would also aid in alleviating any concerns regarding the ability of De Roos et al. (2005) to detect subjects developing NHL.

There are conflicting views on how to interpret the overall results for NHL. Some believe that the data are indicative of a potential association between glyphosate exposure and risk of NHL. This is primarily based on reported effect estimates across studies and the associated meta-analyses greater than 1 despite lack of statistical significance. Additionally, the analysis conducted by Eriksson et al. (2008) observed a slightly statistically significant increase for those with more than 10 years of exposure prior to diagnosis. There were also two case-control studies that investigated the association of glyphosate exposure and NHL by stratifying exposure by days per year of use that reported effect estimates greater than 1 for groups with the highest exposure.

Conversely, others have viewed the effect estimates as relatively small in magnitude and observed associations could be explained by chance and/or bias. All of the effect estimates for ever/never use were non-statistically significant. Sample sizes were small or questionable in some of the studies. Half of the studies reported effect estimates approximately equal to 1, while the other half of the studies reported effect estimates clustered from 1.5-1.85, with the largest effect estimate having the widest confidence interval indicating the estimate was less reliable. As such, the higher effect estimates were contradicted by the results from studies at least equal quality. Meta-analyses were based on studies with varying study characteristics. Given the limitations and concerns discussed above for the studies included in this evaluation, chance and/or bias cannot be excluded as an explanation for the relatively small increase observed in the meta-risk ratios. Meanwhile, analyses performed by De Roos et al. (2005) reported effect estimates less than 1 for cumulative lifetime exposure and intensity-weighted cumulative exposure and these extensive analyses did not detect any exposure-response relationship, which conflicts with the two case-control studies that indicate potential for an exposure-response relationship comparing two groups stratified by days per year of use. Although increased effect estimates were observed in one case-control study (Eriksson et al., 2008) for subjects exposed more than 10 years prior to diagnosis and in two case-control studies (McDuffie et al., 2001; Eriksson et al., 2008) that stratified exposure by days per year of use, none of these analyses appeared to adjust for exposures to other pesticides, which has been found to be particularly important for these analyses and would attenuate these estimates towards the null. Furthermore, none of the studies in this evaluation of glyphosate exposure and risk of NHL accounted for other potential confounders, such as diesel exhaust fumes, solvents, and UV radiation. These adjustments would also be expected to reduce effect estimates towards the null.

Based on the weight-of-evidence, the agency cannot exclude chance and/or bias as an explanation for observed associations in the database. Due to study limitations and contradictory results across studies of at least equal quality, a conclusion regarding the association between glyphosate exposure and risk of NHL cannot be determined based on the available data. The agency will continue to monitor the literature for studies and any updates to the AHS will be considered when available.

4.0 Data Evaluation of Animal Carcinogenicity Studies

4.1 Introduction

Cancer bioassays in animals have historically been the primary studies available to evaluate cancer hazard in humans, since until recently epidemiological evidence was limited. The results of these bioassays, as well as results from screening assays for genotoxicity, are considered in a weight-of-evidence approach to determine the potential of a chemical to induce cancer in humans. Carcinogenicity studies in two rodent species are required for the registration of food use pesticides or when the use of a pesticide is likely to result in repeated human exposure over a considerable portion of the human lifespan (40 CFR Part 158.500). Rodent carcinogenicity studies identified from the data collection phase of the systematic review were evaluated for study quality and acceptable studies were evaluated in the context of the 2005 EPA Guidelines for Carcinogen Risk Assessment as described in Sections 4.2 and 4.3 below, respectively.

4.2 Consideration of Study Quality for Animal Carcinogenicity Studies

The agency has published test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200) and combined chronic/carcinogenicity studies (OCSPP 870.4300) in rodents which have been harmonized with OECD guidelines (Test Nos. 451 and 453). Test substances are typically administered in animal carcinogenicity studies by the oral route for food use pesticides. The studies are generally conducted in mice and rats with exposure durations of 18-24 months for mice and 24 months for rats, which represent exposures of the majority of the expected lifespan in these animals. Guideline carcinogenicity studies are designed to test three or more doses in both sexes (with at least 50 animals/sex/dose) with adequate dose spacing to characterize tumor dose-response relationships. Key considerations when evaluating carcinogenicity studies for cancer hazard assessment include identification of target organs of carcinogenicity, increased incidence of tumors or proportion of malignant neoplasms, and reduction in the time to appearance of tumors relative to the concurrent control group (OECD TG 451).

There are a number of criteria the agency uses when evaluating the technical adequacy of animal carcinogenicity studies. A primary criterion is the determination of the adequacy of dosing. The 2005 EPA Guidelines for Carcinogen Risk Assessment recommends that the highest dose level selected should elicit signs of toxicity without substantially altering the normal life span due to effects other than tumors; or without inducing inappropriate toxicokinetics (e.g., overwhelming absorption or detoxification mechanisms); however, the high dose need not exceed 1,000 mg/kg/day (i.e., limit dose) (OCSPP 870.4200; OCSPP 870.4300). Additional criteria to judge the technical adequacy and acceptability of animal carcinogenicity studies are provided in the test guidelines as well as other published sources (NTP, 1984; OSTP, 1985; Chhabra et al., 1990). As stated in the 2005 EPA Guidelines for Carcinogen Risk Assessment, studies that are judged to be wholly inadequate in protocol, conduct or results, should be discarded from analysis. Studies the agency consider acceptable are further evaluated for potential tumor effects.

Following study quality evaluation, a total of 9 chronic/carcinogenicity studies in the rat and 6 carcinogenicity studies in the mouse were considered acceptable for use in the current evaluation for the active ingredient glyphosate and were subsequently evaluated in the context of the 2005 EPA Guidelines for Carcinogen Risk Assessment as described in Section 4.3. A number of studies were judged to be inadequate in protocol, conduct or reporting and were not considered in the analysis of glyphosate. These studies and the justification for not including them in the analysis are listed below:

1. A two-year chronic oral toxicity study in Albino rats by Reyna (1974)¹⁴. The study was considered inadequate to assess carcinogenicity due to insufficient reporting on the histopathology findings in the control and treatment groups. Approximately 70 animals were unaccounted for across the study.
2. A two-year drinking water study in Wistar rats with a formulated product (13.6% ammonium salt) by Chruscielska et al., (2000). In addition to deficiencies including inadequate reporting of water consumption and body weight data, this study was conducted with a glyphosate formulated product and not the active ingredient glyphosate, which is the focus of this review. Glyphosate formulations contain various components other than glyphosate and it has been hypothesized these components are more toxic than glyphosate alone. The agency is collaborating with NTP to systematically investigate the mechanism(s) of toxicity for glyphosate and glyphosate formulations. This project is discussed in more detail in Section 7.0 of this document.
3. An initiation-promotion study (George et al., 2010) in male Swiss mice that tested a commercial formulation of glyphosate (41%) on the skin. Study deficiencies included small number (20) of animals, tested only males, and lack of histopathological examination.
4. A carcinogenicity study in Swiss albino mice (Kumar, 2001)¹⁵. This study was not included due to the presence of a viral infection within the colony, which confounded the interpretation of the study findings. Malignant lymphomas were reported in this study in all dose groups. However, lymphomas are one of the most common types of spontaneous neoplastic lesions in aging mice (Brayton et al., 2012). Murine leukemia viruses (MuLVs) are also a common cause of lymphoma in many different strains of mice (Ward, 2006). For example, Tadesse-Heath et al. (2000) reported 50% lymphoma (mostly B-cell origin) incidence in a colony of Swiss mice infected with MuLVs. Although the lymphoma incidences in Kumar (2001) were within or near normal background variation, it is not clear whether or not the viral infection may have contributed to the lymphoma incidence reported or the lower survival seen at the high dose in this study.

¹⁴ MRID 00062507.

¹⁵ MRID 49987403. In Greim et al. (2015), the same study is cited as Feinchemie Schwebda (2001).

5. A two year feeding study in Sprague-Dawley rats (Excel, 1997) was not included. The agency does not have access to this study to perform an independent assessment of its conduct and; however, Greim et al. (2015) stated that the study “is considered unreliable for carcinogenicity evaluation” and there were “several deviations from the OECD Test Guideline 453”.

4.3 Assessment of Animal Carcinogenicity Studies

The agency considers many factors when interpreting the results of carcinogenicity studies. The 2005 EPA Guidelines for Carcinogen Risk Assessment are intended as a guidance only and does not provide a checklist for determining whether tumor findings are related to treatment. These guidelines emphasize the importance of weighing multiple lines of evidence in reaching conclusions regarding human carcinogenic potential of chemicals. Evaluation of observed tumor findings takes into consideration both biological and statistical significance. There are several factors in the 2005 EPA Guidelines for Carcinogen Risk Assessment used in the weight-of-evidence evaluation of individual studies. For this evaluation, the interpretation of the evidence related to tumor findings is described below. The agency is soliciting comment from the SAP regarding several of these factors as they relate to the interpretation of studies as part of Charge Question #3.

Dose Selection

Doses should be selected based on relevant toxicological information. Caution is taken in administering an excessively high dose that would confound the interpretation of the results to humans. As mentioned above, the 2005 EPA Guidelines for Carcinogen Risk Assessment recommends that the highest dose level selected should elicit signs of toxicity without substantially altering the normal life span due to effects other than tumors; or without inducing inappropriate toxicokinetics (e.g., overwhelming absorption or detoxification mechanisms); however, the high dose is not recommended to exceed 1,000 mg/kg/day (OCSPP 870.4200; OCSPP 870.4300). Doses should provide relevant dose-response data for evaluating human hazard for human health risk assessment. In the case of glyphosate, the low (oral) systemic toxicity and limited pharmacokinetic (PK) data for this chemical make it difficult to define a maximum tolerated dose (MTD) for the cancer bioassays. A large number of the carcinogenicity studies conducted with glyphosate approach or exceed the limit dose. The 2005 EPA Guidelines for Carcinogen Risk Assessment state that “weighing of the evidence includes addressing not only the likelihood of human carcinogenic effects of the agent but also the conditions under which such effects may be expressed”. As such, the agency puts less weight on observations of tumors that occur near or above the limit dose.

Statistical analyses to evaluate dose response and tumor incidences

The main aim of statistical evaluation is to determine whether exposure to the test agent is associated with an increase in tumor development, rather than due to chance alone. Statistical analyses should be performed on each tumor type separately. The incidence of benign and malignant lesions of the same cell type, usually within a single tissue or organ, are considered

separately, but may be combined when scientifically defensible (McConnell *et al.*, 1986). Trend tests and pairwise comparison tests are the recommended tests for determining whether chance, rather than a treatment-related effect, is a plausible explanation for an apparent increase in tumor incidence. The 2005 Guidelines for Carcinogen Risk Assessment states that

“A trend test such as the Cochran-Armitage test (Snedecor and Cochran, 1967) asks whether the results in all dose groups together increase as dose increases. A pairwise comparison test such as the Fisher exact test (Fisher, 1950) asks whether an incidence in one dose group is increased over that of the control group. By convention, for both tests a statically significant comparison one for which p is less than 0.05 that the increased incidence is due to chance. Significance in either kind of test is sufficient to reject the hypothesis that chance accounts for the result.”

In the current evaluation, the Cochran-Armitage Test for Trend (Snedecor and Cochran, 1967; one-sided) was used. For pairwise comparisons, the Fisher Exact Test (Fisher, 1950; one-sided) was used in the current evaluation to determine if incidences observed in treated groups were different from concurrent controls. Furthermore, the 2005 EPA Guidelines for Carcinogen Risk Assessment state that “considerations of multiple comparisons should also be taken into account”. Multiple comparison methods control the familywise error rate, such that the probability of Type I error (incorrect rejection of the null hypothesis or “false positive”) for the pairwise comparisons in the family does not exceed the alpha level. In the current evaluation, a Sidak correction method was used to adjust for multiple comparisons.

For the current evaluation, statistical significance observed in either test is judged in the context of all of the available evidence. Statistically significant responses may or may not be biologically significant and vice versa (Hsu and Stedeford, 2010; EPA, 2005). If a trend was found to be statistically significant, a closer examination of the tumor incidence was taken to determine whether the data demonstrate a monotonic dose-response where an increase in tumor incidence is expected with corresponding increase in dose. Therefore, statistically significant results with fluctuating tumor incidence across doses are not weighed as heavily as those displaying a monotonic dose-response. If a pair-wise comparison was found to be statistically significant, a closer examination of the tumor incidence and other lines of evidence was taken to determine whether the response was biologically significant. Factors considered in determining the biological relevance of a response are discussed below.

Given that statistical evaluations were performed at different times for each study, all statistical analyses were reanalyzed for the purposes of this evaluation to ensure consistent methods were applied (TXR# 0057494).

Historical Control Data

As indicated in the 2005 EPA Guidelines for Carcinogen Risk Assessment (Section 2.2.2.1.3), the standard for determining statistical significance of tumor incidence comes from a comparison of tumors in dosed animals with those in concurrent control animals. Additional insight into the statistical and/or biological significance of a response can come from the consideration of

historical control data (Tarone, 1982; Haseman, 1995; EPA, 2005). Historical control data can add to the analysis, particularly by enabling identification of uncommon tumor types or high spontaneous incidence of a tumor in a given animal strain. Generally speaking, statistically significant increases in tumors should not be discounted simply because incidence rates in the treated groups are within the range of historical controls or because incidence rates in the concurrent controls are somewhat lower than average.

Historical control data are also useful to determine if concurrent control tumor incidences are consistent with previously reported tumor rates (Haseman, 1995). Given the large number of age-related tumor outcomes in long-term rodent bioassays, and thus the large number of potential statistical tests run, caution is taken when interpreting results that have marginal statistical significance or in which incidence rates in concurrent controls are unusually low in comparison with historical controls since there may be an artificial inflation of the differences between concurrent controls and treated groups. Consequently, in the current evaluation, unusually low incidence in concurrent controls was noted when applicable and considered as part of the weight-of-evidence for the tumor findings. Identification of common or uncommon situations prompts further thought about the meaning of the response in the current study in context with other observations in animal studies and with other evidence about the carcinogenic potential of the agent.

Evidence of supporting preneoplastic lesions or related non-neoplastic lesions

Carcinogenicity rodent studies are designed to examine the production of tumors as well as preneoplastic lesions and other indications of chronic toxicity that may provide evidence of treatment-related effects and insights into the way the test agent produces tumors (EPA, 2005). As such, the presence or lack of supporting preneoplastic or other related non-neoplastic changes were noted in the current evaluation of each study and considered in the weight-of-evidence.

Additional Considerations

Other observations can strengthen or lessen the significance of tumor findings in carcinogenicity studies. Such factors include: uncommon tumor types; tumors at multiple sites; tumors in multiple species, strains, or both sexes; progression of lesions from preneoplastic to benign to malignant; reduced latency of neoplastic lesions (i.e., time to tumor); presence of metastases; unusual magnitude of tumor response; and proportion of malignant tumors (EPA, 2005). The agency considers all of the above factors when determining the significance of tumor findings in animal carcinogenicity studies.

4.4 Summary of Animal Carcinogenicity Studies

A total of 9 chronic toxicity/carcinogenicity studies in the rat and 6 carcinogenicity studies in the mouse were considered acceptable and evaluated in the weight-of-evidence analysis for glyphosate. This includes all of the studies that were part of the 2015 CARC evaluation plus an additional 5 studies identified from the systematic review. In the 2015 CARC evaluation, for some of the studies considered, the CARC relied on summary data that was provided in the supplement to the Greim et al. (2015) review article. Due to the ongoing data collection effort and the acquiring of studies not previously submitted, the agency no longer needs to rely on the

Greim et al. (2015) review article for the study data generated in relevant studies, allowing for a more complete and independent analysis. It should be noted that studies have been cited differently in this evaluation as compared to Greim et al. (2015) so these alternative citations have been noted for applicable studies.

The carcinogenicity studies conducted in the rat and mouse that were considered for the analysis are discussed in Sections 4.5 and 4.6, respectively. In these sections, short study summaries are presented which include information on the study design (including test material, strain of animal used, and doses and route of administration) as well as study findings including effects on survival, general toxicity observed, relevant non-neoplastic lesions, and the incidence and characterization of any tumor findings. The characterization of the tumor response(s) is based on the considerations previously discussed in Section 4.3 for interpreting the significance of tumor findings in animal carcinogenicity studies. The rat and mouse carcinogenicity studies are all summarized in Table 4.11 and Table 4.18, respectively.

4.5 Rat Carcinogenicity Studies with Glyphosate

4.5.1 Burnett et al., 1979 (MRID 00105164)

In a two-year chronic/carcinogenicity oral study, glyphosate (as an aqueous monosodium salt solution) was administered to groups of 90 albino rats/sex/dose at doses of 0, 3, 10, or 30 mg/kg/day (M/F) for 24 months through oral intubation (gavage).

A higher mortality rate was noted in the control group in comparison to the treated groups after 12 and 24 months of testing. No histopathological alterations were observed. There were no treatment-related increases in tumor incidences in the study; however, the highest dose tested in this study was 30 mg/kg/day, which was not considered a maximum tolerable dose to assess the carcinogenic potential of glyphosate.

4.5.2 Lankas, 1981 (MRID 00093879)¹⁶

In a chronic toxicity/carcinogenicity study, groups of Sprague-Dawley rats (50/sex/dose) were fed diets containing glyphosate (98.7%, pure) at dietary doses of 0, 3/3, 10/11, and 31/34 mg/kg/day (M/F).

There were no treatment-related effects on survival at any dose level. As in Burnett (1979), the highest dose tested of approximately 32 mg/kg/day was not considered a maximum tolerable dose to assess the carcinogenic potential of glyphosate. Consequently, a second study (Stout and Ruecker, 1990) was conducted at higher doses, which is summarized in the Section 4.5.3.

**Table 4.1. Testicular Interstitial Cell Tumors in Male Sprague-Dawley Rats (Lankas, 1981)
Cochran-Armitage Trend Test & Fisher's Exact Test Results**

	0 mg/kg/day	3.05 mg/kg/day	10.3 mg/kg/day	31.49 mg/kg/day
--	-------------	----------------	----------------	-----------------

¹⁶ In Greim et al. (2015), the same study is cited as Monsanto (1981).

**Table 4.1. Testicular Interstitial Cell Tumors in Male Sprague-Dawley Rats (Lankas, 1981)
Cochran-Armitage Trend Test & Fisher's Exact Test Results**

Incidence (%)	0/50 (0)	3/47 (6)	1/49 (2)	6/44 (12)
Raw p-value =	0.009**	0.121	0.500	0.013*
Sidak p-value =	--	0.321	0.875	0.039*

Note: Trend test results denoted at control; * denotes significance at p=0.05; ** denotes significance at p=0.001.

A statistically significant trend was reported for the testicular interstitial tumors; however, closer examination of the tumor incidence indicates that the data do not demonstrate a monotonic dose response with greater incidence observed at the low-dose as compared at the mid-dose. The incidence at the high dose was found to be statistically significant as compared to the concurrent controls. The observed incidence of interstitial cell tumors in concurrent controls (0%) appears to be unusually low for this tumor type as compared to historical controls provided in the study report for this tumor type (mean = 4.5%; range = 3.4%-6.7%) resulting in an artificial difference at the high dose. Furthermore, the observed incidence of interstitial cell tumors in the glyphosate-treated groups were within the normal biological variation for this tumor type in this strain of rat. There was an absence of pre-neoplastic or related non-neoplastic lesions (e.g., interstitial cell hyperplasia). As a result, the statistically significant results do not appear to be biologically significant and are not supported by any histopathological observations. Based on the weight-of-evidence for this study, the agency does not consider the increases in interstitial cell tumors in the testes to be treatment-related.

4.5.3 Stout and Ruecker, 1990 (MRID 41643801)¹⁷

In a chronic toxicity/carcinogenicity study, groups of Sprague-Dawley rats (60/sex/dose) were fed diets containing glyphosate (96.5%, pure) at dietary doses of 0, 89/113, 362/457 or 940/1183 mg/kg/day M/F for 24 months. The highest dose tested in this study approaches or exceeds the highest dose recommended in the test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200 and OCSPP 870.4300). Tumor findings at these high doses are given less weight.

There was no significant increase in mortality. The most frequently seen tumors were pancreatic cell adenomas, hepatocellular adenomas, and thyroid C-cell adenomas in males. A discussion of each tumor type by organ is presented below:

1. Pancreas: Tumor incidences of pancreatic islet cell tumors in male rats and corresponding historical control values are presented in Tables 4.2 and 4.3, respectively. The incidence of pancreatic islet cell tumors lacked monotonic dose-responses and trend analyses were not statistically significant. Statistical significance was observed with raw (unadjusted) p-values for the incidence of adenomas at the low-dose (89 mg/kg/day) and high-dose (940 mg/kg/day) when comparing to concurrent controls; however, none of the incidences were statistically significant with an adjustment for multiple comparisons (p=0.052 at the low-dose and p=0.120 at the high-dose). The statistical significance of

¹⁷ In Greim et al. (2015), the same study is cited as Monsanto (1990).

the pairwise comparisons with the concurrent control group may have been due to the unusually low incidences in the controls and not to an actual treatment-related response. The mean incidence of pancreatic islet cell adenomas in historical control data provided for laboratory (Monsanto Environmental Health Laboratory; MRID No. 41728701) was 5.3% and ranged from 1.8% to 8.3% indicating the concurrent control incidence for this tumor type was at the lower bound of the range. Carcinomas were only observed in the control group and the combined analyses did not yield any statistically significant pairwise comparisons. There were no supporting preneoplastic or other related non-neoplastic changes observed and no evidence of progression from adenomas to carcinomas. Based on a weight-of-evidence for this study, the agency does not consider these increases in pancreatic islet cell tumors to be treatment-related.

Table 4.2. Pancreatic Islet Cell Tumors in Male Sprague-Dawley Rats (Stout and Ruecker, 1990) Cochran-Armitage Trend Test & Fisher's Exact Test Results.				
Tumor Type	0 mg/kg/day	89 mg/kg/day	362 mg/kg/day	940 mg/kg/day
Adenoma Incidence (%)	1/43 ^a (2)	8/45 (18)	5/49 (10)	7/48 ^b (15)
Raw p-value =	0.176	0.018*	0.135	0.042*
Sidak p-value =	--	0.052	0.352	0.120
Carcinoma Incidence (%)	1/43 ^c (2)	0/45 (0)	0/49 (0)	0/48 (0)
Raw p-value =	- ^d	1.000	1.000	1.000
Sidak p-value =	--	1.000	1.000	1.000
Combined Incidence (%)	2/43 (2)	8/45 (18)	5/49 (10)	7/48 (15)
Raw p-value =	0.242	0.052	0.275	0.108
Sidak p-value =	--	0.149	0.619	0.289

Note: Trend test results denoted at control; * denotes significance at p=0.05.

- a. Number of tumor-bearing animals/Number of animals examined, excluding those that died or were sacrificed prior to study week 55.
- b. First adenoma in the study was observed at week 81 in the 940 mg/kg/day group.
- c. First carcinoma in the study was observed at week 105 in the controls.
- d. Trend p-value not reported since tumor incidence decreased with increasing dose.

Historical control data on the incidence of pancreatic islet cell adenomas in male Sprague-Dawley rats in 2-year studies (1983–1989) conducted at the testing facility (Monsanto Environmental Health Laboratory; MRID No. 41728701) are presented below in Table 4.3.

Table 4.3. Historical Control Data — Pancreatic Islet Cell Adenomas in Male Sprague- Dawley Rats (MRID No. 41728701).								
Study No.	1	2	3	4	5	6	7	Mean
Study Year	07/83	02/85	10/85	6/85	9/88	1/89	3/89	-
Tumor Incidence	2/68	5/59	4/69	1/57	5/60	3/60	3/59	-
Percentage (%)	2.9%	8.5%	5.8%	1.8%	8.3%	5.0%	5.1%	5.3%

2. Liver: Tumor incidences of liver tumors in male rats are presented in Tables 4.4. There was a statistically significant dose trend for liver adenomas only. Closer examination of the incidence indicates a relatively flat response at the low- and mid-dose with only an increase observed at the high-dose (940 mg/kg/day); however, the incidence of liver adenomas at the high-dose was not statistically significant when compared to the concurrent controls. Carcinomas and combined adenomas/carcinomas lacked statistical significance in trend and pairwise comparisons (Table 4.4). Except for a single animal at the mid-dose late in the study (89 weeks), no hyperplasia, preneoplastic foci or other non-neoplastic lesions were observed. Furthermore, there was no evidence of progression from adenomas to carcinomas. Given the lack of both statistical significance and corroborative lesions to support the tumor finding, the agency does not consider these increases in liver tumors to be treatment-related.

Table 4.4. Hepatocellular Tumors in Male Sprague-Dawley Rats (Stout and Ruecker, 1990) Cochran-Armitage Trend Test & Fisher's Exact Test Results				
Tumor Type	0 mg/kg/day	89 mg/kg/day	362 mg/kg/day	940 mg/kg/day
Adenoma Incidence (%) Raw p-value = Sidak p-value =	2/44 ^a (5) 0.022* --	2/45 (4) 0.700 0.973	3/49 (6) 0.551 0.910	7/48 ^b (15) 0.101 0.274
Carcinoma Incidence (%) Raw p-value = Sidak p-value =	3/44 (7) - ^d -	2/45 (4) 0.827 0.995	1/49 (2) 0.954 1.000	2/48 ^c (4) 0.845 0.996
Combined Incidence (%) Raw p-value = Sidak p-value =	5/44 (11) 0.078 --	4/45 (9) 0.769 0.988	4/49 (8) 0.808 0.993	9/48 (19) 0.245 0.569

Note: Trend test results denoted at control; * denotes significance at p=0.05.

a. Number of tumor-bearing animals/Number of animals examined, excluding those that died or were sacrificed prior to study week 55.

b. First adenoma in the study was observed at week 88 in the 940 mg/kg/day group.

c. First carcinoma in the study was observed at week 85 in the 940 mg/kg/day group.

d. Trend p-value not reported since tumor incidence decreased with increasing dose.

3. Thyroid: Tumor incidences of thyroid tumors in male and female rats are presented in Tables 4.6 and 4.7, respectively. For males, no statistically significant trends were observed for adenomas, carcinomas, or combined adenomas/carcinomas. For females, a statistically significant trend was observed for adenomas and combined adenomas/carcinomas with no statistically significance in pairwise analyses. Therefore, although there may be an indication of a dose-response in females, the increases observed in the glyphosate treated groups were not considered to be different than those observed in the concurrent controls. Non-neoplastic lesions (thyroid C-cell hyperplasia) were

observed; however, there was a lack of a monotonic dose-response for these histopathological findings and no dose-related increase in severity to support tumor findings (Table 4.8). There was also no evidence of progression from adenomas to carcinomas. Based on a weight-of-evidence for this study, the agency does not consider these increases in thyroid tumors to be treatment-related.

Table 4.6. Thyroid C-Cell Tumors in Male Sprague-Dawley Rats (Stout and Ruecker, 1990) Cochran-Armitage Trend Test & Fisher's Exact Test Results				
Tumor Type	0 mg/kg/day	89 mg/kg/day	362 mg/kg/day	940 mg/kg/day
Adenoma Incidence (%) Raw p-value = Sidak p-value =	2/54 ^{a, b} (4) 0.079 --	4/55 (7) 0.348 0.723	8/58 (14) 0.060 0.168	7/58 (12) 0.099 0.269
Carcinoma Incidence (%) Raw p-value = Sidak p-value =	0/54 (0) 0.457 --	2/55 ^c (4) 0.252 0.441	0/58 (0) 1.000 1.000	1/58 (4) 0.518 0.768
Combined Incidence (%) Raw p-value = Sidak p-value =	2/54 (4) 0.087 --	6/55 (11) 0.141 0.367	8/58 (14) 0.060 0.168	8/58 (14) 0.060 0.168

Note: Trend test results denoted at control.

- a. Number of tumor-bearing animals/Number of animals examined, excluding those that died or were sacrificed prior to study week 55.
- b. First adenoma in the study was observed at week 54 in the controls.
- c. First carcinoma in the study was observed at week 93 in the 89 mg/kg/day group.

Table 4.7. Thyroid C-Cell Tumors in Female Sprague Dawley Rats Cochran-Armitage Trend Test & Fisher's Exact Test Results (Stout and Ruecker, 1990).				
Tumor Type	0 mg/kg/day	113 mg/kg/day	457 mg/kg/day	1183 mg/kg/day
Adenoma Incidence (%) Raw p-value = Sidak p-value =	2/57 ^a (4) 0.040* --	2/60 (7) 0.710 0.976	6/59 ^b (10) 0.147 0.380	6/55 (11) 0.124 0.328
Carcinoma Incidence (%) Raw p-value = Sidak p-value =	0/57 (0) 0.494 --	0/60 (0) 1.000 1.000	1/59 ^c (2) 0.509 0.509	0/55 (0) 1.000 1.000
Adenoma/Carcinoma Incidence (%) Raw p-value = Sidak p-value =	2/57 (4) 0.042* --	2/60 (3) 0.710 0.976	7/59 (12) 0.090 0.246	6/55 (11) 0.124 0.328

Note: Trend test results denoted at control; * denotes significant at p=0.05.

- a. Number of tumor-bearing animals/Number of animals examined, excluding those that died or were sacrificed prior to study week 55.

- b. First adenoma in the study was observed at week 72 in the controls.
c. First carcinoma in the study was observed at week 93 in the 457 mg/kg/day group.

Table 4.8. Thyroid Non-Neoplastic Lesions (Stout and Ruecker, 1990)				
Males				
Dose	0 mg/kg/day	89 mg/kg/day	362 mg/kg/day	940 mg/kg/day
Total Incidences of thyroid C-cell hyperplasia and severity scores	5/60 (8%) Diffuse (moderate) – 1 Multi-focal (minimal) – 3 Focal (mild) – 1	1/60 (2%) Focal (mild) – 1	6/60 (10%) Focal (minimal) – 4 Multi-focal (minimal) – 1 Multi-Focal (mild) – 1	5/60 (8%) Focal (minimal) – 2 Focal (mild) – 1 Multi-focal (mild) – 1 Multi-focal (moderate) – 1
Females				
	0 mg/kg/day	113 mg/kg/day	457 mg/kg/day	1183 mg/kg/day
Thyroid C-cell hyperplasia and severity scores	10/60 (17%) Diffuse (moderate) – 1 Focal (mild) – 1 Focal (minimal) – 1 Focal (mild) – 1 Focal (moderate) – 1 Multi-focal (minimal) – 3 Multi-focal (moderate) – 1 Diffuse (moderate) – 1	5/60 (8%) Focal (mild) – 3 Focal (minimal) – 1 Multi-focal (minimal) – 1	9/60 (15%) Focal (minimal) – 4 Multi-focal (minimal) – 2 Multi-focal (mild) – 3	5/60 (8%) Focal (mild) – 1 Focal (minimal) – 1 Multi-focal (mild) – 2 Diffuse (moderate) – 1

*Data taken from pages 1071-2114 of the study report.

4.5.4 Atkinson et al., 1993a (MRID 496317023)¹⁸

In a combined chronic toxicity/carcinogenicity study, glyphosate (98.9% pure) was administered to 50 Sprague-Dawley rats/sex/dose in the diet at doses of 0, 11/12, 112/109, 320/347, and 1147/1134 mg/kg/day for 104 weeks (M/F) for 104 weeks. An additional 35 rats/sex/dose were included for 1-year interim sacrifice.

No adverse effects on survival were seen in either sex across the doses tested. There were no changes in histopathological findings observed. There were no treatment-related increases in tumor incidences in the study.

4.5.5 Brammer, 2001 (MRID 49704601)¹⁹

In a combined chronic toxicity/carcinogenicity study, glyphosate acid (97.6% pure) was administered to groups of Wistar rats in the diet. Groups of 52 rats/sex received diets containing doses of 0, 121/145, 361/437 or 1214/1498 mg/kg/day for 24 months, in males/females, respectively. The highest dose tested in this study exceeds the highest dose recommended in the

¹⁸ Note: In Greim et al. (2015), the same study is cited as Cheminova (1993a).

¹⁹ Note: In Greim et al. (2015), the same study is cited as Syngenta (2001).

test guidelines on how to conduct carcinogenicity studies (OCSP 870.4200 and OCSP 870.4300).

A statistically significant higher survival ($p=0.02$) was observed in males at the highest dose tested at the end of 104 weeks relative to concurrent controls, and a statistically significant trend for improved survival was observed in treated males ($p=0.03$). The inter-current (early) deaths were 37/52, 36/52, 35/52, and 26/52 for the control, low-, mid-, and high-dose groups, respectively. The terminal deaths were 16/52, 17/52, 18/52, and 26/52 for the control, low-, mid- and high-dose groups, respectively. There were no treatment-related non-neoplastic lesions in any organs of either sex at any dose level tested. As shown in Table 4.9, a statistically significant trend in the incidences of liver adenomas was observed in male rats; however, a monotonic dose-response was not observed upon closer examination of the incidence data. Tumor incidences appear to fluctuate with increases observed at the low- and high-dose and no tumors observed in the control and mid-dose. Statistical significance with raw (unadjusted) p -values was observed for the tumor incidence at the high-dose (1214 mg/kg/day) when compared to concurrent controls; however, it was not statistically significant with an adjustment for multiple comparisons ($p=0.056$). Tumor findings at these high doses are given less weight. The improved survival in the high-dose group may help explain a modestly higher incidence of an age-related background tumor like liver adenomas and this corresponds with the lack of associated lesions. Given that the tumor findings did not reflect a monotonic dose response and the high dose tumors were not statistically significant with an adjustment for multiple comparisons, the agency does not consider these increases in liver adenomas to be treatment-related.

Table 4.9. Liver Adenomas in Male Wistar Rats (Brammer, 2001) Cochran-Armitage Trend Test and Fisher's Exact Test Results.				
	0 mg/kg/day	121 mg/kg/day	361 mg/kg/day	1214 mg/kg/day
Adenoma Incidence (%)	0/52 ^a (0)	2/52 (4)	0/52 (0)	5/52 (10)
Raw p -value =	0.008**	0.248	1.000	0.028*
Sidak p -value =	--	0.434	1.000	0.056

Note: Trend test results denoted at control; * denotes significance at $p=0.05$; ** denotes significance at $p=0.01$.
a. Number of tumor-bearing animals/Number of animals examined.

4.5.6 Pavkov and Wyand 1987 (MRIDs 40214007, 41209905, 41209907)

Glyphosate trimesium salt (sulfosate, 56.2% pure) was tested in a 2-year chronic feeding/carcinogenicity study in male and female Sprague-Dawley (CrI:CD[SD]BR) rats. Sixty animals/sex were tested in control group 1 (basal diet, no vehicle), 80/sex were tested in control group 2 (basal diet plus propylene glycol at 1% w/w vehicle) and in the low and mid-dose groups, and 90/sex were tested in the high dose group. The following dose levels were tested: 0, 4.2/5.4, 21.2/27 or 41.8/55.7 mg/kg/day in males and females respectively.

Treatment had no effect on survival. There were no changes in histopathological findings observed. There were no treatment-related increases in tumor incidences in the study.

4.5.7 Suresh, 1996 (MRID 49987401)²⁰

In a combined chronic toxicity/carcinogenicity study, glyphosate (96.0-96.8% pure) was administered to groups of Wistar rats in the diet. Groups of 50 rats/sex/group received diets containing 0, 6.3/8.6, 59.4/88.5, and 595.2/886 mg/kg/day glyphosate for 24 months in males and females respectively. The highest dose tested in females in this study approaches the highest dose recommended in the test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200 and OCSPP 870.4300).

No adverse effects on survival were observed in either sex across the doses tested. There were no changes in histopathological findings observed. There were no treatment-related increases in tumor incidence observed in the study.

4.5.8 Enemoto, 1997 (MRID 50017103-50017105)²¹

In a combined chronic toxicity and carcinogenicity study, groups of 50 Sprague-Dawley rats/sex/group received daily dietary doses of 0, 104/115, 354/393 and 1127/1247 mg/kg bw/day glyphosate for males and females, respectively. In addition, 10 rats/sex/group were included for interim sacrifices at 26, 52, and 78 weeks. The highest dose tested in this study exceeds the highest dose recommended in the test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200 and OCSPP 870.4300).

There were no changes in mortality at any of the doses tested. There were no changes in histopathological findings observed. There were no treatment-related increases in tumor incidence observed in the study.

4.5.9 Wood et al., 2009a (MRID 49957404)²²

In a combined chronic toxicity/carcinogenicity study, glyphosate (95.7% pure) was administered to groups of Wistar rats in the diet. Groups of 51 rats/sex/group received diets containing 0, 95.0, 316.9, and 1229.7 mg/kg/day glyphosate for males and female, respectively. The highest dose tested in this study exceeds the highest dose recommended in the test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200 and OCSPP 870.4300).

No adverse effects on survival were seen in either sex across the doses tested. There were no treatment-related preneoplastic or related non-neoplastic lesions in either sex at any dose level.

In female rats, mammary gland tumors were noted. Tumor incidences for mammary gland adenomas, adenocarcinomas, and combined adenomas/adenocarcinomas in female mice are presented in Table 4.10. Statistically significant trends were observed for the adenocarcinoma and combined analyses. Tumor incidence for adenocarcinomas was not statistically significant

²⁰ Note: In Greim et al. (2015), the same study is cited as Feinchemie Schwebda (1996).

²¹ Note: In Greim et al. (2015), the same study is cited as Arysta Life Sciences (1997b).

²² Note: In Greim et al. (2015), the same study is cited as NuFarm (2009b).

in pairwise comparisons as compared to concurrent controls. Marginal statistical significance was observed with the raw (unadjusted) p-value for combined mammary gland tumors at the high-dose (1229.7 mg/kg/day) when comparing to concurrent controls; however, with an adjustment for multiple comparisons, the increased incidence at the high-dose was not statistically significant (p=0.132). There was also no evidence of progression from adenomas to carcinomas. Based on a weight-of-evidence for this study, the agency does not consider these increases in mammary gland tumors in female rats to be treatment-related.

Table 4.10. Mammary Gland Tumor Incidences in Female Rats (Wood et al., 2009a) Fisher's Exact Test and Cochran-Armitage Trend Test Results				
Tumor Type	0 mg/kg/day	95.0 mg/kg/day	316.9 mg/kg/day	1229.7 mg/kg/day
Adenoma Incidence (%) Raw p-value = Sidak p-value =	0/51 (0) 0.062 --	0/51 (0) 1.000 1.000	0/51 (0) 1.000 1.000	2/51 (4) 0.248 0.248
Adenocarcinoma Incidence (%) Raw p-value = Sidak p-value =	2/51 (4) 0.042* --	3/51 (6) 0.500 0.875	1/51 (2) 0.879 0.998	6/51 (12) 0.135 0.352
Combined Incidence (%) Raw p-value = Sidak p-value =	2/51 (4) 0.007** --	3/51 (6) 0.500 0.875	1/51 (2) 0.879 0.998	8/51 (16) 0.046* 0.132

Note: Trend test results denoted at control; * denotes significance at p=0.05; ** denotes significant at p=0.01.

4.5.10 Summary of Rat Data

In 5 of the 9 rat studies conducted with glyphosate, no tumors were identified for detailed evaluation. Of the remaining 4 rat studies, a statistically significant trend was observed for tumor incidences in the testes, pancreas, liver, thyroid, or mammary gland; however, the agency determined that these tumor findings are not considered to be related to treatment. Although a statistically significant trend was obtained, closer examination of the incidence data across doses did not demonstrate a monotonic dose response in several instances. Some of the tumor incidences at the highest dose tested (approaching or exceeding 1,000 mg/kg/day for almost all studies) were statistically significant from concurrent controls using raw (unadjusted) p-values; however, none of the pairwise comparisons were found to be statistically significant following adjustment for multiple comparisons, except the testicular tumors seen in a single study. Furthermore, these high-dose tumors were given less weight. There was no evidence of corroborating pre-neoplastic or related non-neoplastic lesions or evidence of tumor progression (progression from pre-neoplastic to malignancy) to support biological significance of tumor findings. In a limited number of cases, the agency considered historical control data to inform the relevance of a tumor increase when incidence rates in the concurrent controls were unusually low.

Table 4.11. Summary of Rat Carcinogenicity Studies			
Study	Dose Range	Pre-Neoplastic or Related Non-Neoplastic Lesions	Tumors Incidences, Statistical Significance, and Related Comments
Burnett et al. (1979) Albino rats	0, 3, 10 or 30 mg/kg/day for 24 months [M/F]	None observed	There were no treatment-related increases in tumor incidences.
Lankas (1981) Sprague-Dawley rats	98.7% Technical in diet 0, 3/3, 10/11, and 31/34 mg/kg/day [M/F]	None observed	Statistically significant trend observed for testicular interstitial cell tumors; however, did not observe monotonic dose-response with higher incidence at low-dose than mid-dose. Incidences were 0/50 in controls, 3/47 at low-dose, 1/49 at mid-dose, and 6/44 at high-dose. Increased incidence at high-dose statistically significant, but unusually low control incidence (based on historical control data in study report) inflated increase at high-dose.
Stout and Ruecker (1990) Sprague-Dawley rats	96.5% Technical in diet 0, 89/113, 362/457 and 940/1183 mg/kg/day [M/F] for 24 months	None observed	Pancreatic tumors lacked statistically significant trend. Tumor incidence for pancreatic adenomas in males were 1/43 in controls, 8/45 at the low-dose, 5/49 at the mid-dose, and 7/48 at the high-dose. Concurrent control incidence for this tumor type was at the lower bound of the historical control range. No statistically significant pairwise comparisons, including the highest dose tested which is approaching/exceeding 1,000 mg/kg/day. Statistically significant trend for liver adenomas in males with only an increase at high-dose. Incidences were 2/44 in controls, 2/45 at the low-dose, 3/49 at the mid-dose, and 7/48 at the high-dose. No statistically significant pairwise comparisons, including the highest dose tested which is approaching/exceeding 1,000 mg/kg/day. No statistically significant trend for thyroid C-cell tumors in males. For females, statistically significant trend for adenomas and combined adenomas/carcinomas. Incidences for adenomas were 2/57 in controls, 2/60 at the low-dose, 6/59 at the mid-dose, and 6/55 at the high-dose. Similar incidences were seen for combined except the mid-dose was 7/59. No statistically significant pairwise comparisons, including the highest dose tested which is approaching/exceeding 1,000 mg/kg/day.
Atkinson et al. (1993a) Sprague-Dawley rats	98.9% Technical in diet 0, 11/12, 112/109, 320/347, and 1147/1134 mg/kg/day for 104 weeks (M/F)	None observed	There were no treatment-related increases in tumor incidences, including the highest dose tested which exceeded 1,000 mg/kg/day.

Table 4.11. Summary of Rat Carcinogenicity Studies			
Study	Dose Range	Pre-Neoplastic or Related Non-Neoplastic Lesions	Tumors Incidences, Statistical Significance, and Related Comments
Brammer. (2001) Wistar rats	97.6% Technical in diet 0, 121/145, 361/437 and 1214/1498 mg/kg/day [M/F]	None observed	Statistically significant trend in liver adenomas in males. Incidences were 0/52 in controls, 2/52 at the low-dose, 0/52 at the mid-dose, and 5/52 at the high-dose. No statistically significant pairwise comparisons when adjusting for multiple comparisons, including the highest dose tested which exceeded 1,000 mg/kg/day.
Pavkov and Wyand (1987) Sprague-Dawley rats	56.2% Technical (Trimesium salt; Sulfosate) 0, 4.2/5.4, 21.2/27 and 41.8/55.7 mg/kg/day [M/F]	None observed	There were no treatment-related increases in tumor incidences.
Suresh (1996) Wistar rats	96.0-96.8% Technical in diet 0, 6.3/8.6, 59.4/88.5, and 595.2/886 mg/kg/day [M/F]	None observed	There were no treatment-related increases in tumor incidences, including the highest dose tested which exceeded 1,000 mg/kg/day.
Enemoto (1997) Sprague-Dawley rats	94.61-97.56% Technical in diet 0, 104/115, 354/393 and 1127/1247 mg/kg/day [M/F]	None observed	There were no treatment-related increases in tumor incidences, including the highest dose tested which exceeded 1,000 mg/kg/day.
Wood et al. (2009a) Wistar rats	95.7% Technical in diet 0, 86/105, 285/349 or 1077/1382 mg/kg/day [M/F]	None observed	Statistically significant trends were observed for the mammary gland adenocarcinoma and combined adenoma/adenocarcinoma analyses. Incidences for adenocarcinomas were 2/51 in controls, 3/51 at the low-dose, 1/51 at the mid-dose, and 6/51 at the high-dose. Similar incidences observed for combined adenoma/adenocarcinomas except incidence at high-dose was 8/51. No statistically significant pairwise comparisons when adjusting for multiple comparisons, including the highest dose tested which exceed 1,000 mg/kg/day.

4.6 Mouse Carcinogenicity Studies with Glyphosate

4.6.1 Reyna and Gordon, 1973 (MRID 00061113)

In an 18-month carcinogenicity study, groups of 50 Swiss white mice/sex/dose were fed glyphosate at dietary levels of approximately 17 mg/kg/day and 50 mg/kg/day. There was no effect on survival at any of the doses tested. There were no changes in histopathological findings observed. There were no treatment-related increases in tumor incidence observed in the study. Although only ten mice/sex/dose were examined for histopathological changes, there were no statistically significant increases in tumors observed in the study; therefore, this deficiency would not impact the overall conclusion regarding tumor findings.

4.6.2 Knezevich and Hogan, 1983 (MRID 00130406)²³

Groups of 50 male and female CD-1 mice received glyphosate (99.78%, pure) at dietary doses of 0, 161/195, 835/968, 4945/6069 mg/kg/day for males and females, respectively for 24 months. The highest dose tested in this study far exceeds the highest dose recommended in the test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200 and OCSPP 870.4300). Furthermore, the mid-dose tested in this study was approaching 1,000 mg/kg/day. Tumor findings at these high doses are given less weight.

No effect on survival was observed. There were no corroborating lesions to support any tumor findings in this study.

A low incidence of renal tubule adenomas, which are considered rare, were noted in males. The incidences of renal tubule adenomas following initial evaluation of the study were reported as follows: 0/49 in the controls; 0/49 at the low-dose; 1/50 at the mid-dose; and 3/50 at the high dose (TXR No. 0004370). In 1985, the registrant directed a re-evaluation of the original renal sections by a consulting pathologist. This re-evaluation identified a small renal tubule adenoma in one control male mouse, which was not diagnosed as such in the original pathology report. In 1986, at the request of the agency, additional renal sections (3 sections/kidney/mouse spaced at 150 micron intervals) were evaluated in all control and all glyphosate-treated male mice in order to determine if additional tumors were present. The additional pathological and statistical evaluations concluded that the renal tumors in male mice were not compound-related.

Subsequently, the agency requested a Pathology Work Group (PWG) evaluate the kidney sections. The PWG examined all sections of the kidney, including the additional renal sections, and were blinded to treatment group. The renal tubular-cell lesions diagnosed by the PWG are presented below in Table 4.12 with results from statistical analyses. The PWG noted that because differentiation between tubular-cell adenoma and tubular-cell carcinoma is not always clearly apparent and because both lesions are derived from the same cell type, it is appropriate to combine the incidences from these two tumor types for purposes of evaluation and statistical

²³ Note: In Greim et al. (2015), the same study is cited as Monsanto (1983).

analysis. The PWG unanimously concluded that these lesions are not compound-related based on the following considerations: 1) renal tubular cell tumors are spontaneous lesions for which there is a paucity of historical control data for this mouse stock; 2) there was no statistical significance in a pairwise comparison of treated groups with the concurrent controls and there was no evidence of a statistically significant linear trend; 3) multiple renal tumors were not found in any animal; and 4) compound-related nephrotoxic lesions, including pre-neoplastic changes, were not present in male mice in this study (TXR No. 0005590).

Table 4.12. Kidney Tubular Cell Tumors in Male CD-1 Mice (Knezevich and Hogan, 1983) Cochran-Armitage Trend Test & Fisher's Exact Test Results.				
Tumor Type	0 mg/kg/day	161 mg/kg/day	835 mg/kg/day	4945 mg/kg/day
Adenoma Incidence (%) Raw p-value = Sidak p-value =	1/49 (2) 0.4422 --	0/49 (0) 1.000 1.000	0/50 (0) 1.000 1.000	1/50 (2) 0.758 0.986
Carcinoma Incidence (%) Raw p-value = Sidak p-value =	0/49 (0) 0.063 --	0/49 (0) 1.000 1.000	1/50 (2) 0.505 0.755	2/50 (4) 0.253 0.441
Combined Incidence (%) Raw p-value = Sidak p-value =	1/49 (2) 0.065 --	0/49 (0) 1.000 1.000	1/50 (2) 0.758 0.986	3/50 (6) 0.316 0.680

Note: Trend test results denoted at control.

Histopathological examinations noted chronic interstitial nephritis and tubular epithelial changes (basophilia and hypertrophy) in the kidneys of male rats in the study (Table 4.13). The increased incidence of chronic interstitial nephritis in males lacked a dose-response. The incidence in controls of bilateral interstitial nephritis was higher than low-dose group and approximately the same as the mid-dose group. Unilateral chronic interstitial nephritis was only seen in 1 animal in the low- and high-dose groups. Furthermore, chronic interstitial nephritis is not considered to be a precursor lesion for tubular neoplasms. A monotonic dose-response was not observed for the epithelial basophilia and hypertrophy, such that the incidence fluctuated with dose and the lowest incidence was observed at the highest dose tested. There was no increase in supporting preneoplastic or related non-neoplastic renal tubular lesions (e.g., tubular epithelial necrosis/regeneration, hyperplasia) observed in male mice.

Table 4.13. Kidney Histopathological Alterations in Male CD-1 Mice (Knezevich and Hogan, 1983)				
Males				
Dose	0 mg/kg/day	161 mg/kg/day	835 mg/kg/day	4945 mg/kg/day
Bilateral Chronic Interstitial Nephritis	5/49 (10%)	1/49 (2%)	7/50 (14%)	11/50 (22%)

Unilateral Chronic Interstitial Nephritis	0/49 (0%)	1/49 (2%)	0/49 (0%)	1/50 (2%)
Proximal Tubule Epithelial Basophilia and Hypertrophy	15/49 (31%)	10/49 (20%)	15/50 (30%)	7/50 (14%)

*Data taken from page 305 and 306, and the study pathology report; incidences were moderate diffuse

Based on the weight-of-evidence for this study, the agency concurs with the PWG conclusion, following a thorough examination of all kidney sections, that the renal tubular neoplasms are not treatment-related with a lack of statistical significance in the trend and pairwise tests. Although there was an increase in chronic interstitial nephritis at the highest dose tested, this finding is not considered relevant to the tubular neoplasms.

4.6.3 Atkinson, 1993b (MRID 49631702)²⁴

In a carcinogenicity study, glyphosate (>97% pure) was administered to groups of 50 CD-1 mice/sex/dose in the diet for 104 weeks at doses of 0, 98/102, 297/298, 988/1000 mg/kg/day for males and females, respectively. No interim sacrifices were performed.

There was no effect on survival in the study. There were no preneoplastic lesions or related non-neoplastic lesions observed. As shown in Table 4.14, hemangiosarcomas were found in 4/45 (9%) of high-dose male mice (1000 mg/kg/day) compared to none in the concurrent controls or other treated groups. Hemangiosarcomas are commonly observed in mice (generally more common in males for CD-1 strain) as both spontaneous and treatment-related tumors arising from endothelial cells. As vascular tumors, they can occur at different sites, with liver and spleen tending to be the most common sites in mice. In the high-dose mice with hemangiosarcomas, one had the tumors present in the liver and spleen, one had the tumor present in the liver only, one had the tumors present in the liver, spleen, and prostate, and one had the tumor present in the spleen only. A statistically significant trend was observed ($p=0.00296$). Closer examination of the incidence indicates a relatively flat response at the low- and mid-dose with only an increase observed at the high-dose; however, the incidence of hemangiosarcomas at the high-dose was not statistically significant when compared to the concurrent controls. Based on a weight-of-evidence for this study, the agency does not consider these increases in hemangiosarcomas in male mice to be treatment-related.

Table 4.14. Hemangiosarcomas in Male CD-1 Mice (Atkinson, 1993b)				
Cochran-Armitage Trend Test and Fisher's Exact Test Results.				
Dose (mg/kg/day)	0	100	300	1000
Hemangiosarcoma Incidence (%)	0/47 ^a (0)	0/46 (0)	0/50 (0)	4/45 (9)
Raw p-value =	0.003**	1.000	1.000	0.053
Sidak p-value =	--	1.000	1.000	0.053

Note: Trend test results denoted at control; * denotes significance at $p=0.05$; ** denotes significance at $p=0.01$.
a= Number of tumor bearing animals/Number of animals examined, excluding those that died before week 52.

²⁴ Note: In Greim et al. (2015), the same study is cited as Cheminova (1993b).

4.6.4 Wood et al., 2009b (MRID 49957402)²⁵

In a feeding study conducted in 2009, CD-1 mice (50/sex/dose) received glyphosate (95.7%) for 80 weeks at dietary dose levels of 0, 71.4/97.9, 234.2/299.5, or 810/1081.2 mg/kg/day for males and females, respectively. The highest dose tested in this study approaches or exceeds the highest dose recommended in the test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200 and OCSPP 870.4300).

There was no effect on survival in the study. In male mice at the high dose, there were increases in the incidences of lung adenocarcinomas and malignant lymphomas. A discussion of each tumor type is presented below:

1. Lung: Tumor incidence for lung adenomas, adenocarcinomas, and combined adenomas/adenocarcinomas are presented in Table 4.15. A statistically significant trend was only noted for the adenocarcinomas. Closer examination of the tumor incidence indicates the dose-response was relatively flat at the low- and mid-dose with only an increase observed at the high-dose; however, the incidence of lung adenocarcinomas at the high-dose (810 mg/kg/day) was not statistically significant when compared to the concurrent controls. There were no treatment-related preneoplastic or related non-neoplastic lesions observed. There was also no evidence of progression from adenomas to carcinomas. Based on a weight-of-evidence for this study, the agency does not consider these increases in lung tumors to be treatment-related.
2. Malignant lymphoma: Tumor incidence for malignant lymphoma are also presented in Table 4.16. A statistically significant trend was observed and the incidence at the high-dose (810 mg/kg/day) was statistically significantly elevated as compared to concurrent controls with the raw (unadjusted) p-value; however, with an adjustment for multiple comparisons, the increased incidence at the high-dose was not statistically significant ($p=0.082$). Historical control data were also considered to better understand the significance of the reported increased incidence of lymphoma. Historical control data from the same laboratory and same supplier are preferred; however, this data were not available for consideration with the study report. The 2005 EPA Guidelines for Carcinogen Risk Assessment does not prohibit the use of historical control data from other sources; however, it does state it should be used with caution. For this strain of mouse, the mean incidence for untreated animals is approximately 4.5% (range: 1.5%-21.7%) based on historical control data from Charles River (59 studies performed from 1987-2000; Giknis and Clifford, 2005) and Huntingdon Laboratories (20 studies from 1990-2002; Son and Gopinath, 2004). Although the data are not from the performing laboratory, it does indicate that the incidence in concurrent controls in this study was low, which can contribute to the pairwise significance observed at the highest dose tested with the raw (unadjusted) p-value. Based on a weight-of-evidence for this study, the agency does not consider the increase in malignant lymphoma to be treatment-related.

²⁵ Note: In Greim et al. (2015), the same study is cited as NuFarm (2009a).

Table 4.15. Lung Tumors in Male CD-1 Mice (Wood et al., 2009b) Fisher's Exact Test and Cochran-Armitage Trend Test Results.				
Dose (mg/kg/day)	0	71.4	234.2	810
Lung Adenoma Incidence (%)	9/51 (18)	7/51 (14)	9/51 (18)	4/51 (8)
Raw p-value =	_ ^b	0.793	0.602	0.964
Sidak p-value =	-	0.991	0.937	1.000
Lung Adenocarcinoma (%)	5/51 ^a (10)	5/51 (10)	7/51 (14)	11/51 (22)
Raw p-value =	0.028*	0.630	0.380	0.086
Sidak p-value =	--	0.949	0.762	0.237
Lung Combined Incidence (%)	14/51 (27)	12/51 (24)	16/51 (31)	15/51 (29)
Raw p-value =	0.336	0.752	0.414	0.500
Sidak p-value =	--	0.985	0.799	0.875

Note: Trend test results denoted at control; * denotes significance at p=0.05;** denotes significance at p=0.01
a= Number of tumor bearing animals/Number of animals examined.

b = Trend p-value not reported since tumor incidence decreased with increasing dose.

Table 4.16. Malignant Lymphomas in Male CD-1 Mice (Wood et al., 2009b) Fisher's Exact Test and Cochran-Armitage Trend Test Results.				
Dose (mg/kg/day)	0	71.4	234.2	810
Malignant Lymphoma Incidence (%)	0/51 (0)	1/51 (2)	2/51 (4)	5/51 (10)
Raw p-value =	0.007**	0.500	0.248	0.028*
Sidak p-value =	--	0.875	0.574	0.082

Note: Trend test results denoted at control; * denotes significance at p=0.05;** denotes significance at p=0.01
a= Number of tumor bearing animals/Number of animals examined.

4.6.5 Sugimoto, 1997 (MRID 50017108 - 50017109)²⁶

In a carcinogenicity study, glyphosate (purity 97.56 and 94.61%; two lots) was administered to groups of 50 male and 50 female Specific-Pathogen-Free (SPF) ICR (Crj: CD-1) mice/dose in the diet at dose levels of 0, 165/153.2, 838.1/786.8, or 4348/4116 mg/kg/day for males and females, respectively, for 18 months. The highest dose tested in this study far exceeds the highest dose recommended in the test guidelines on how to conduct carcinogenicity studies (OCSPP 870.4200 and OCSPP 870.4300). Furthermore, the mid-dose tested in this study was approaching 1,000 mg/kg/day. Tumor findings at these high doses are given less weight.

There were no treatment-related effects on mortality or survival. There were no changes in histopathological findings observed.

²⁶Note: In Greim et al. (2015), the same study is cited as Arysta Life Sciences (1997b)

Hemangiomas in female mice were found to occur at different sites. The tumor incidences are presented in Table 4.17. A statistically significant trend was observed. Tumor incidence at the high-dose, which was approximately 4 times the recommended high-dose in test guidelines (4116 mg/kg/day), was statistically significant with the raw (unadjusted) p-value as compared to concurrent controls; however, with an adjustment for multiple comparisons, the high dose tumors were not statistically significant (p=0.055). Based on a weight-of-evidence for this study, the agency does not consider these increases in hemangiomas in female rats to be treatment-related.

Table 4.17. Hemangioma Incidences (Sugimoto, 1997) Fisher's Exact Test and Cochran-Armitage Trend Test Results				
Tumor Type	0 mg/kg/day	153.2 mg/kg/day	786.8 mg/kg/day	4116 mg/kg/day
Hemangioma Incidence (%)	0/50 (0)	0/50 (0)	2/50 (4)	5/50 (10)
Raw p-value =	0.002**	1.000	0.247	0.028*
Sidak p-value =	--	1.000	0.434	0.055

Note: Trend test results denoted at control; * denotes significance at p=0.05; ** denotes significance at p=0.01.

4.6.6 Pavkov and Turnier, 1987 (MRIDs 40214006, 41209907)

Glyphosate trimesium salt (sulfosate, 56.2% pure) was tested in a 2-year chronic feeding/carcinogenicity study in male and female CD-1 mice. Sixty animals/sex were tested in control group 1 (basal diet, no vehicle), 80/sex were tested in control group 2 (basal diet plus propylene glycol at 1% w/w vehicle) and in the low- and mid-dose groups, and 90/sex were tested in the high-dose group. The following dose levels were tested: 0, 11.7/16, 118/159, and 991/1341 mg/kg/day for males and females, respectively.

No adverse effects on survival were seen in either sex across the doses tested. There were no changes in histopathological findings observed. There were no treatment-related increases in tumor incidence observed in the study.

4.6.7 Summary of Mouse Data

No tumors were identified for detailed evaluation in 2 of the 6 mouse carcinogenicity studies. In the remaining 4 mouse studies, 3 observed a statistically significant trend in tumor incidences in the hemangiosarcomas, lung adenomas, malignant lymphomas or hemangiomas; however, the agency determined that none of the tumors observed in the mouse are treatment related. Although a statistically significant trend was obtained, closer examination of the incidence data across doses did not demonstrate a monotonic dose response in several instances. Some of the tumor incidences at the highest dose tested (approaching or exceeding 1,000 mg/kg/day for almost all studies) were statistically significant from concurrent controls using raw (unadjusted) p-values; however, none of the pairwise comparisons were found to be statistically significant following adjustment for multiple comparisons. Furthermore, these high-dose tumors were given less weight. There was no evidence of corroborating pre-neoplastic or related non-neoplastic lesions or evidence of tumor progression (progression from pre-neoplastic to malignancy) to support

biological significance of tumor findings. In a limited number of cases, the agency considered historical control data to inform the relevance of a tumor increase when incidence rates in the concurrent controls were unusually low.

Table 4.18. Summary of Mouse Carcinogenicity Studies			
Study	Dose Range	Pre-Neoplastic or Related Non-Neoplastic Lesions	Tumors Incidences, Statistical Significance, and Related Comments
Reyna and Gordon (1973) Swiss white mice	0, 17 or 50 mg/kg/day for 18 months	None observed	There were no treatment-related increases in tumor incidence.
Knezevich and Hogan (1983) CD-1 mice	99.78% Technical in diet 0, 161/195, 835/968, 4945/6069 mg/kg/day for [M/F] for 24 months.	Chronic interstitial nephritis lacked dose-response and not considered relevant to renal tumors. Tubular epithelial changes in kidney were approximately the same in controls, low- and mid-doses and then decreased at high-dose.	The incidences of renal tubule adenomas were: 1/49 (2%) in the controls; 0/49 at the low-dose; 1/50 at the mid-dose; and 3/50 (6%) at the high dose. No statistical significance in trend or pairwise comparisons, including the mid- and high-doses which approached or exceeded 1,000 mg/kg/day.
Atkinson et al. (1993b). CD-1 mice	97.5 - 100.2% Technical in diet 0, 98/102, 297/298, 988/1000 mg/kg/day for 104 weeks (M/F)	None observed	Statistically significant trend for hemangiosarcomas that were only observed in 4/45 (9%) high-dose male mice. Increased incidence was not statistically significant from the concurrent controls at all doses, including the highest dose tested which is approximately 1,000 mg/kg/day.
Wood et al. (2009b) CD-1 mice	95.7% Technical in diet 0, 71.4/97.9, 234.2/299.5, or 810/1081.2 mg/kg/day [M/F] for 80 weeks	None observed	Statistically significant trend for lung adenocarcinomas with incidences of 5/51 in controls, 5/51 at the low-dose, 7/51 at the mid-dose, and 11/51 at the high-dose. No statistical significance in pairwise comparisons. Statistically significant trend for malignant lymphoma with incidences of 0/51 in controls, 1/51 at the low-dose, 2/51 at the mid-dose, and 5/51 at the high-dose. Incidence in concurrent controls for this tumor type was low. No statistically significant pairwise results with multiple comparison adjustment, including the highest dose tested which was approaching 1,000 mg/kg/day.
Sugimoto (1997) CD-1 mice	94.61 – 97.56% Technical in diet 0, 165/153.2, 838.1/786.8, or 4348/4116 mg/kg/day [M/F] for 18 months	None observed	Statistically significant trend for hemangiomas female mice with incidences of 0/50 in controls, 0/50 at the low-dose, 2/50 at the mid-dose, and 5/50 at the high-dose. No statistically significant pairwise results with multiple comparison adjustment, including the mid- and high-doses which approached or exceeded 1,000 mg/kg/day.

Table 4.18. Summary of Mouse Carcinogenicity Studies			
Study	Dose Range	Pre-Neoplastic or Related Non-Neoplastic Lesions	Tumors Incidences, Statistical Significance, and Related Comments
Pavkov and Turnier (1987) CD-1 mice	56.2% Technical (Trimesium salt; Sulfosate) 0, 11.7/16, 118/159, and 991/1341 mg/kg/day [M/F] for 24 months.	None observed	There were no treatment-related increases in tumor incidence, including the highest dose tested which approached/exceeded 1,000 mg/kg/day.

4.7 Absorption, Distribution, Metabolism, Excretion (ADME)

The 2005 EPA Guidelines for Carcinogen Risk Assessment also permit analysis of other key data that may provide valuable insights into the likelihood of human cancer risk from exposure to a chemical, such as information regarding the absorption, distribution, metabolism, and excretion (ADME) of a test chemical. EPA's Harmonized Test Guidelines for pesticides include a series of studies for characterizing a chemical's metabolism and pharmacokinetics. As described in the test guideline (OCSPP 870.7485), testing of the disposition of a test substance is designed to obtain adequate information on its: absorption, distribution, biotransformation (metabolism), and excretion, which can all collectively aid in understanding the chemical's mechanism of toxicity. Basic pharmacokinetic/toxicokinetic parameters determined from these studies can also provide information on the potential for accumulation of the test substance in tissues and/or organs and the potential for induction of biotransformation as a result of exposure to the test substance. These data can be used to assess the adequacy and relevance of the extrapolation of animal toxicity data (particularly chronic toxicity and/or carcinogenicity data) to estimate human risk.

Oral exposure is considered the primary route of concern for glyphosate. The maximum absorption from the GI tract for glyphosate was estimated to be ~30% with one study showing up to 40% based upon radiolabel detected in the urine. In general, the amounts of glyphosate detected in tissues were negligible indicating low tissue retention following dosing. Parent glyphosate is the principal form excreted in urine and feces. The primary route of excretion following oral administration of glyphosate is the feces, as verified by the intravenous dosing and bile cannulation experiments. Within the dose ranges tested, elimination was essentially complete by 24 hours indicating that glyphosate does not bioaccumulate.

Multiple studies examined the pharmacokinetics of a single dose of radiolabeled glyphosate ranging from 5.6 – 400 mg/kg. Across these studies, time to reach peak plasma concentrations (T_{max}) appeared to increase with increasing dose; however, the reported range of T_{max} (1-5.5 hours) suggests only a slight shift in absorption kinetics occurs despite large increases in dose. In the one study that tested two doses (NTP, 1992), data graphically show that peak blood levels were only roughly 3-fold with a 10-fold increase between the two doses. Reported area under the curve (AUC) values indicated conflicting results regarding whether linear or non-linear absorption kinetics was occurring at higher doses.

In general, EPA and OECD guideline ADME studies are designed for a different purpose and do not provide the information needed to adequately determine whether linear kinetics is still occurring at high doses of glyphosate. These studies are often limited to one or two doses and do not include time course data. A well-conducted pharmacokinetic study testing multiple doses is needed to conclusively make this determination.

4.8 Discussion

Glyphosate has been extensively tested in rodents to evaluate its carcinogenic potential. A total of 15 rodent carcinogenicity studies were considered to be adequate for this analysis. Nine studies were conducted in the rat and 6 studies were conducted in the mouse. When a potential tumor signal was identified in a study, the agency considered several factors. Consistent with the EPA's 2005 Guidelines for Carcinogen Risk Assessment, the agency evaluated the tumor responses for both statistical and biological significance by considering factors such as historical control data; rarity of tumor types; tumors at multiple sites; tumors in multiple species, strains, or both sexes; progression of lesions from preneoplastic to benign to malignant; reduced latency of neoplastic lesions (i.e., time to tumor); presence of metastases; unusual magnitude of tumor response; proportion of malignant tumors; and dose-related increases. When these factors were considered together, the agency made a determination of whether or not the observed tumor was related to treatment with glyphosate. A weight of the evidence approach was used to determine the carcinogenic potential of glyphosate in rodents.

In 5 of the 9 rat studies conducted with glyphosate, no tumors were identified for detailed evaluation. Of the remaining 4 rat studies, a statistically significant trend was observed for tumor incidences in the testes, pancreas, liver, thyroid, or mammary gland; however, the agency determined that these tumor findings are not considered to be related to treatment, as described in Section 4.5, due to lack of pairwise statistical significance, lack of a monotonic dose response, absence of preneoplastic or non-neoplastic lesions, no evidence of tumor progression, and/or historical control information (in limited instances). Lastly, tumors seen in individual rat studies were not reproduced in other studies, including those conducted in the same animal species and strain at similar or higher doses.

In 2 of the 6 mouse studies, no tumors were identified for detailed evaluation. In the remaining 4 mouse studies, 3 observed a statistically significant trend in tumor incidences in the hemangiosarcomas, lung adenomas, malignant lymphomas or hemangiomas; however, the agency determined that none of the tumors observed in the mouse are treatment related, as described in Section 4.6, due to lack of pairwise statistical significance, lack of a monotonic dose response, absence of preneoplastic or non-neoplastic lesions, no evidence of tumor progression, and/or historical control information (in limited instances). Lastly, tumors seen in individual mouse studies were not reproduced in other studies, including those conducted in the same animal species and strain at similar or higher doses.

In addition to the lines of evidence considered when determining if a tumor was treatment-related within a study, the agency also looked across all of the relevant studies to determine if the tumor findings were reproducible in other studies conducted in the same species and strain. Increased incidence of testicular, pancreatic, thyroid and mammary gland tumors were seen in only one study and were not reproduced in the other four studies for that strain at similar or higher doses. An increased incidence of hepatocellular adenomas were seen in one study with Sprague-Dawley rats and one study with Wistar rats, but this tumor type was not significantly

increased in the other six studies tested in these rat strains at similar or higher doses. In the mice, an increase in the incidence of renal tumors, hemangiosarcomas, lung adenomas, malignant lymphoma and hemangiomas were reported only in a single study and findings were not seen in the four other studies conducted in CD-1 mice at similar or higher doses.

When looking across the studies at doses where potential tumor signals were identified, doses below 500 mg/kg/day consistently showed no increased incidence of tumors with the single exception of the testicular tumors in SD rats (Lankas, 1981), where an increase in incidence was seen at approximately 31.5 mg/kg/day. However, as discussed in Section 4.5.2, the testicular tumor data do not show a monotonic dose response, the concurrent controls appear to be unusually low for this tumor, there were no pre-neoplastic or related non-neoplastic lesions, and this tumor type was not seen in other studies at doses up to 35-fold higher in the same strain of rat. As a result, the increased incidence in testicular tumors was not considered treatment-related based on the weight-of-evidence for the study. Even if the tumor findings observed above 500 mg/kg/day were considered indicative of treatment-related effects, the 2005 EPA Guidelines for Carcinogen Risk Assessment state that the “weighing of the evidence includes addressing not only the likelihood of human carcinogenic effects of the agent but also the conditions under which such effects may be expressed”. As such, the high doses (~1,000 mg/kg/day or greater) where these tumor findings were observed were considered in the context of potential exposure to glyphosate in residential and occupational settings. As previously discussed in Section 1.4, oral exposure is the primary route of concern for glyphosate. In residential/non-occupational settings, children 1-2 years old are considered the most highly exposed subpopulation with an estimate of potential combined exposure of 0.47 mg/kg/day. The estimated maximum potential exposure for occupational workers is 7 mg/kg/day. The estimate of exposure children and occupational workers is at least 2,000-fold and 140-fold lower, respectively, than the doses (~1000 mg/kg/day) where increases in tumor incidences were typically observed in the rodent studies. Based on these exposure estimates, the high dose tumor findings are not considered relevant for human health risk assessment.

Based on the weight-of-evidence, the agency has determined that any tumor findings observed in the rat and mouse carcinogenicity studies for glyphosate are not considered treatment-related. Tumor findings observed at the highest doses tested were also not reproduced in studies in the same animal strain at similar or higher doses. Furthermore, even if the high-dose tumors were considered treatment-related, these findings are not considered relevant for human health risk assessment based on the use pattern and potential exposures for glyphosate.

5.0 Data Evaluation of Genetic Toxicity

5.1 Introduction

Genotoxicity is a broad term for any damage to the genetic material, whether the damage is transient or permanent. Transient damage refers to unintended modifications to the structure of DNA, which may or may not undergo successful repair. Permanent damage refers to heritable changes in the DNA sequence, known as mutations. Types of mutations include: 1) changes in single base pairs, partial, single or multiple genes, or chromosomes, 2) breaks in chromosomes that result in transmissible deletion, duplication or rearrangement of chromosome segments, and 3) mitotic recombination (OECD, 2015). In somatic cells, DNA-reactive chemicals can cause cancer if the mutations occur within regulatory genes that control cell growth, cell division and differentiation, such as proto-oncogenes, tumor suppressor genes and/or DNA damage response genes (OECD, 2015). Additionally, DNA damage may signal the cell to undergo apoptosis (cell death) rather than cell division and, therefore, the damage is not “fixed” as a mutation and is not passed along to daughter cells.

Evaluation of genotoxicity data entails a weight-of-evidence approach that includes consideration of the various types of genetic damage that can occur. Since no single genotoxicity assay evaluates the many types of genetic alterations that can be induced by a chemical, one must employ a battery of genotoxicity tests to adequately cover all the genetic endpoints important for regulatory decisions. EPA, like other regulatory agencies, considers genotoxicity information as part of the weight of evidence when assessing the potential of a chemical to induce cancer in humans. Under FIFRA, OPP requires genotoxicity tests of the technical grade active ingredient for the registration of both food and non-food use pesticides. The current genotoxicity test battery (40 CFR Part 158.500) for pesticide registration consists of:

- 1) Bacterial reverse mutation test (typically conducted in bacteria strains *Salmonella typhimurium* and *Escherichia coli*),
- 2) *in vitro* mammalian (forward) gene mutation and *in vitro* mammalian chromosomal aberration test, and
- 3) *in vivo* test for micronucleus induction (mammalian erythrocyte micronucleus test) or *in vivo* chromosomal aberration test (mammalian bone marrow chromosomal aberration test).

In cases where equivocal or inconsistent results are obtained for the same endpoint in different test systems, additional testing may be required. Test Guidelines on how to conduct the genotoxicity tests have been published by the agency and have been harmonized with the Organization for Economic Cooperation and Development (OECD, 2015; Cimino 2006). These guidelines identify specific test species, genetic endpoints, test conditions, exposure durations as well information on how to report data and interpret the results. The test guidelines provide a level of consistency and predictability for regulatory compliance and regulatory decision making.

5.2 Scope of the Assessment Considerations for Study Quality Evaluation

Previous genotoxicity assessments conducted as part of the CARC reviews for glyphosate in 1991 and 2015, considered only studies conducted with glyphosate technical and included only studies that provided adequate characterization of the test material (*i.e.* purity information provided). In the current analysis, a fit-for-purpose systematic review process was conducted to identify relevant genotoxicity data from regulatory studies and published literature from open sources (published and unpublished) for both glyphosate technical and glyphosate-based formulations. Studies conducted with glyphosate formulations that were identified and considered relevant for genotoxicity evaluation are summarized in table form in Appendix F. As described in Section 7.0 of this document, glyphosate formulations are hypothesized to be more toxic than glyphosate alone. The agency is collaborating with NTP to systematically investigate the mechanism(s) of toxicity for glyphosate and glyphosate formulations. However the focus of this section is the genotoxic potential of glyphosate technical.

As described previously in Section 2.1.3, the list of studies identified in this process were also cross-referenced with genotoxicity review articles for glyphosate from the open literature [Kier and Kirkland (2013), and Williams *et al.* (2000)], as well as recent international evaluations of glyphosate (IARC 2015, EFSA 2015, JMPR 2016). The current analysis also includes studies conducted by other registrants that were not previously available to the agency. Sixteen studies for glyphosate technical that were included in Kier and Kirkland (2013) were not available to the agency; therefore, data and study summaries provided in the review articles were relied upon in the current review and are identified in the data tables with a footnote. The Kier and Kirkland (2013) article serves as the original publication for these studies and provided relevant information on study design and conditions as well as summary data. The data set includes *in vitro* and *in vivo* studies conducted in mammalian systems, with the exception of standard bacterial test strains, which have a long history of detecting chemicals that are mutagenic in humans. Studies conducted in non-mammalian species (e.g. worms, fish, reptiles, plants), were excluded because they were considered to be not relevant for informing genotoxic risk in humans.

When evaluating the quality of the published and unpublished data for inclusion in the analysis, the agency considered the reporting quality (how well a study was reported), the study design and how well the study was conducted. Critical elements in study design and interpretation for genotoxicity tests are described in the various EPA and OECD test guidelines. Elements such as test conditions (e.g. solubility, pH, osmolarity, and cytotoxicity) and study design (e.g. number of test organisms, doses selected, use of positive and negative controls; blinded evaluation) were used to evaluate the quality of published and non-published studies. In cases where inappropriate testing conditions or study design clearly had an impact on the outcome the study, the study was excluded from the analysis. For example, early studies by Majeska (1982) were excluded from the analysis since it was clearly demonstrated that altered pH by the test chemical can result in false positive responses in several of *in vitro* genotoxicity tests (Majeska, 1985d,e,f). In other cases, particularly with the published literature studies, where test conditions and/or study design differed from what is generally considered as acceptable following in the EPA or OECD guidelines, the differences are noted, but the studies were not excluded from analysis unless the condition made the study unreliable. Summaries of relevant genotoxicity studies can be found in TXR# 0057499. Studies that were excluded from the analysis are listed in Appendix G.

The studies evaluating the genetic toxicity of the active ingredient glyphosate are presented in the following sections according to the type of genetic endpoints evaluated: mutations, chromosomal aberrations and other assays evaluating DNA damage. *In vitro* and *in vivo* assays are discussed separately according to the genetic endpoint. For the purpose of this analysis, glyphosate and its salts are considered together when evaluating the genotoxic potential of the active ingredient glyphosate.

5.3 Tests for Gene Mutations for Glyphosate Technical

5.3.1 Bacterial Mutagenicity Assays

Bacteria have traditionally been employed as a primary test organism for the detection of chemical mutagens. The bacterial reverse mutation assay is routinely performed in the test strains of *Salmonella typhimurium* and *Escherichia coli*. These test strains are mutant strains that are deficient for the synthesis of an essential amino acid. The assay detects mutations that revert the test strains back to wild type for amino acid synthesis and the revertants are identified by their ability to grow in culture medium deficient of the specific amino acid(s). This mutagenicity test identifies point mutations, which includes base substitutions and deletions and insertions of up to a few base pairs (OECD 471). The tests are typically conducted in the presence and absence of an exogenous source of metabolic activation (e.g., S9 microsomal fraction of activated liver homogenates) to identify potential mutagenic metabolites.

Glyphosate has been extensively evaluated for its potential to induce mutations in bacteria. Most of the studies considered consist of the full battery of bacterial strains (*i.e.* the recommend strains in EPA and OECD Test Guidelines) and were evaluated at appropriate test concentrations (up to cytotoxic or assay limit concentrations).

EPA identified 27 studies that tested glyphosate technical in bacterial mutagenicity assays by means of the standard plate incorporation method or the pre-incubation modification of the standard assay. Glyphosate was negative in the presence and absence of metabolic activation in all the studies. The results of the bacterial reversion mutation assays evaluating glyphosate technical are presented in Table 5.1

Table 5.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Technical.						
Test/Endpoint	Test System	Concentrations	Purity	Results	Reference	Comments
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA1537, TA98 and TA100 and WP <i>uvrA</i> ± S9	156-5000 µg/plate	95.68%	Negative ± S9	Akanuma (1995) [MRID 50017102]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA535, TA1537, TA98 and TA100 and <i>E. coli</i> WP2P and WP2P <i>uvrA</i> ± S9	100-5000 µg/plate in DMSO	95.6% glyphosate acid	Negative ± S9	Callander (1996) [MRID 44320617]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA 1535, TA1537, TA98 and TA100 and <i>E. coli</i> WP2P and WP2P <i>uvrA</i> ± S9	100-5000 µg/plate in water	60% potassium glyphosate salt	Negative ± S9	Callander (1999) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA97a, TA98, TA100 and TA102, ± S9	25-2000 µg in aqueous solution	Not provided	Negative ± S9	Chruscielska <i>et al.</i> (2000)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 ± S9	10-1000 µg/plate	98.4%	Negative ± S9	Flowers and Kier (1978) [MRID 00078620]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, TA1537 ± S9	31.6-3160 µg/plate	98.8%	Negative ± S9	Flügge (2009a) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, TA1537 ± S9	31.6-3160 µg/plate	96.4% technical	Negative ± S9	Flügge (2010b) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA1537, TA98 and TA100	310-5000 µg/plate (+S9); 160-2500 µg/plate (-S9)	98.6%	Negative ± S9	Jensen (1991a) [MRID 49961502]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, TA1537 ± S9	1-1000 µg/plate	98.05%	Negative ± S9	Miyaji (2008) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537, TA1538 ± S9	5000 µg/plate	Not reported	Negative ± S9	Moriya <i>et al.</i> (1983)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA97, TA98 and TA100 ± S9	33-10,000 µg/plate	99%	Negative ± S9	NTP (1992)	Hamster and rat S9

Table 5.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Technical.						
Test/Endpoint	Test System	Concentrations	Purity	Results	Reference	Comments
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535 and TA97a ± S9	1-5000 µg/plate	61.27 % Glyphosate isopropyl-amine salt	Negative ± S9	Ranzani (2000) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, TA1537 ± S9	648-5000 µg/plate	98.01%	Negative ± S9	Ribeiro do Val (2007) [MRID 50000903]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. Coli</i> WP2 <i>uvrA</i> ± S9	31.6-5000 µg/plate	96.0% technical	Negative ± S9	Schreib (2010) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA1537, TA1538, TA98, TA100 and <i>E. coli</i> WP2 <i>hcr</i> ± S9	10-5000 µg/plate	98.4%	Negative ± S9	Shirasu <i>et al.</i> (1978) [MRID 00078619]	Published in Li & Long, 1988
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP <i>uvrA</i> ± S9	3-5000 µg/plate (plate-incorporation), 33-5000 µg/plate (pre-incubation test)	95.1%	Negative ± S9	Sokolowski (2007a) [MRID 49957406]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP <i>uvrA</i> ± S9	3-5000 µg/plate (plate-incorporation) 33 – 5000 µg/plate (pre-incubation test)	97.7%	Negative ± S9	Sokolowski (2007b) [MRID 49957407]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP <i>uvrA</i> ± S9	3-5000 µg/plate (plate-incorporation) 33-5000 µg/plate (pre-incubation test)	95.0%	Negative ± S9	Sokolowski (2007c) [MRID 49957408]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP <i>uvrA</i> ± S9	3-5000 µg/plate	96.66% technical	Negative ± S9	Sokolowski (2009a) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> pKM 101 and WP2 pKM 101 ± S9	3-5000 µg/plate	96.3% glyphosate acid	Negative ± S9	Sokolowski (2009b) [MRID 49961801]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP <i>uvrA</i> ± S9	3-5000 µg/plate	97.16 %	Negative ± S9	Sokolowski (2010) [MRID 50000902]	

Table 5.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Technical.						
Test/Endpoint	Test System	Concentrations	Purity	Results	Reference	Comments
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537, TA1538 ± S9	1-1000 µg/plate	96.0%	Negative ± S9	Suresh (1993a) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP <i>uvrA</i> ± S9	0-5000 µg/plate	95.3%	Negative ± S9	Thompson (1996) [MRID 49957409]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, TA1537 ± S9	31.6-5000 µg/plate	98.2%	Negative ± S9	Wallner (2010) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98 and TA100 ± S9	25 µg/plate	Not reported	Negative ± S9	Wilderman and Nazar (1982)	Rat S9 and plant cell-free homogenates were used for metabolic activation
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA1537, TA1538, TA98 and TA100 ± S9	0.12-10 mg/plate –S9 0.56-15 mg/plate +S9	90% glyphosate trimesium salt	Negative ± S9	Majeska et al. (1982a) [MRID 00126612]	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA1537, TA98 and TA100 ± S9	0.005-50 µL/mL	55.6% glyphosate trimesium salt	Negative ± S9	Majeska (1985a) [MRID 00155527]	

¹ Study was cited in Kier and Kirkland (2013). Supplementary information about the study was provided online including test guideline, test material purity, control chemicals and summary data tables.

5.3.2 *In vitro* Tests for Gene Mutations in Mammalian Cells

In vitro gene mutation studies in mammalian cells are conducted in cell lines with reporter genes for forward mutations. The most common reporter genes are the endogenous thymidine kinase (TK) gene, endogenous hypoxanthine-guanine phosphoribosyl transferase (HPRT) gene and the xanthine-guanine phosphoribosyl transferase transgene (XPRT). Mutations that occur within these reporter genes result in mutant cells that are resistant to the cytotoxic effect of the pyrimidine analogue trifluorothymidine (for TK) or the purine analogue 6-thioguanine (for HPRT and XPRT) (OPPTS 870.5330). Suitable cell lines for this assay include L5178Y mouse lymphoma cells, Chinese hamster ovary (CHO) cells, hamster AS52 and V79 lung fibroblasts and human TK6 lymphoblastoid cells. Similar to other *in vitro* assays, chemicals are tested both in the presence and absence of S9 metabolic activation.

A total of four studies were conducted for (forward) mutations in mammalian cells (Table 5.3). Three studies were conducted with a high purity concentration of glyphosate technical ($\geq 95.6\%$) and the remaining study was performed with glyphosate trimesium salt. In four of the assays, mouse lymphoma L5178Y TK^{+/−} cells were the target organism and one was conducted in CHO cells with the HPRT endpoint. Glyphosate technical and the glyphosate trimesium salt were negative in the mouse lymphoma cell assays (Jensen, 1991b; Clay, 1996; Majesak, 1985b) when tested up to the current guideline limit concentration and glyphosate was negative in CHO/HPRT cells when tested up to cytotoxic concentrations (Li, 1983a).

Table 5.2. <i>In vitro</i> Mammalian Gene Mutation Assays: Glyphosate Technical.						
Test/Endpoint	Test System	Concentrations/ Conditions	Test Material Purity	Results	Reference	Comments
Gene Mutations in Mammalian Cells	Mouse lymphoma L5178Y TK ^{+/−} cells ± S9	296–1000 µg/mL	95.6%	Negative	Clay (1996) ¹	Relative survival was 90% (-S9) and 57% (+S9) at top concentration
Gene Mutations in Mammalian Cells	Mouse lymphoma L5178Y TK ^{+/−} cells ± S9	520–4200 µg/mL (+S9); 610–5000 µg/mL (-S9)	98.6%	Negative	Jensen (1991b) [MRID 49961504]	Reported no significant reduction in cloning efficiency at any concentration.
Gene Mutations in Mammalian Cells	Chinese hamster ovary (CHO) cells, HPRT locus ± S9	500–25000 µg/mL (+S9); 500–22500 µg/mL (-S9)	98.7%	Negative	Li (1983a); [MRID 00132681]	Tested S9 from 1–10% Cytotoxic at 22.5 mg/mL (-S9, and with 1,2 and 10% S9) and at 17.5 mg/ml (10% S9)
Gene Mutations in Mammalian Cells	Mouse lymphoma L5178Y TK ^{+/−} cells ± S9	1–5 µl/mL	55.6% <i>Glyphosate trimesium salt</i>	Negative	Majeska (1985b) [MRID 00155530]	Negative with pH adjusted

¹ Study was cited in Kier and Kirkland (2013). Supplementary information about the study was provided online including test guideline, test material purity, control chemicals and summary data tables.

5.4 *In vitro* Tests for Chromosomal Abnormalities

Cytogenetic assays are tests that can detect chemicals that cause structural chromosomal damage (clastogenicity) or affect the segregation of chromosomes during cell division and alter chromosome number (aneuploidy). Generally, there are two types of *in vitro* cytogenetic assays that identify chemicals inducing chromosomal abnormalities: chromosomal aberration assays and micronucleus assays. Although chromosomal damage observed in these assays are not considered heritable mutations, chemicals that can induce these types of chromosomal damage can also induce transmissible mutations to daughter cells indicating their role in cancer (Yauk *et al.*, 2015; OECD 2015). In addition, assays such as (fluorescence *in situ* hybridization (FISH)) can provide additional mechanistic information on the formation of chromosomal abnormalities. It is important to note that factors such as cytotoxicity, solubility of the test substance, changes in pH or osmolality play a significant role in the outcome of the assay. Like other *in vitro* assays, compounds are generally tested in the presence or absence of S9 metabolic activation to determine if metabolism affects the genotoxic activity of the parent compound and to determine if potential genotoxic metabolites are formed.

5.4.1 *In vitro* Mammalian Chromosomal Aberration Test

Chromosomal aberration assays detect both structural chromosomal and numerical aberrations. Structural chromosomal aberrations are of two types: chromatid and chromosome and include breaks, deletions and rearrangements (OPPTS 870.5375, OECD 2015). Numerical chromosomal aberrations generally results from the loss of an entire chromosome mostly due to damage in the spindle fiber resulting in aneuploidy. The types of cells that are most commonly used in chromosomal aberration assays include established cell lines such as Chinese hamster lung (CHL) and CHO cells or primary cell cultures such as human or other mammalian peripheral blood lymphocytes. In this assay, cells are typically sampled at a time equivalent to the length of approximately 1.5 cell cycles from the start of treatment. Prior to harvesting, cells are treated with Colcemid® or colchicine to arrest cells at the first metaphase stage of the cell cycle following the beginning of exposure to the test article. Once harvested, the cells are stained and metaphase cells are evaluated microscopically for various types of chromosome aberrations. (OECD TG 473). Data should be presented in a way that indicates the percentage of affected cells in the population of cells scored (e.g., % cells with aberrations or # aberrant cells/100 cells). Gaps should not be included in the analysis; they are scored but gaps alone in the absence of any additional chromosomal aberrations (e.g., a fragment or a ring chromosome) are not sufficient to define a cell as aberrant.

Glyphosate technical was evaluated in eight chromosomal aberrations tests to determine its potential to induce clastogenic effects *in vitro*. The findings are presented in Table 5.3. Six of the eight studies were negative. The two positive studies were both from the same laboratory where, Lioi *et al.* reported an increase in chromosomal aberrations at glyphosate concentrations of 8.5µM and above in bovine lymphocytes (Lioi *et al.*, 1998b) and at all concentrations of glyphosate tested (7-170 µM) in human lymphocytes (Lioi *et al.*, 1998a) following a 72-hour exposure period. No chromosomal aberrations were observed as a result of exposure to glyphosate in one study using CHO cells (Majeska, 1985c) and in two studies with CHL cells

(Matsumoto, 1995; and Wright, 1996). Sivikova and Dianovsky (2006) reported no statistically significant increases in chromosomal aberrations in bovine lymphocytes treated with glyphosate (62% pure) at concentrations up to 1120 μ M following 24-hour exposure. (Sivikova and Dianovsky, 2006). In studies conducted with human lymphocytes treated with glyphosate ($\geq 95\%$) for 24-96 hours at concentrations, no increase in chromosomal aberrations were seen at concentrations as high as 6000 μ M (Fox, 1998; and Manas *et al.*, 2009).

5.4.2 *In vitro* Mammalian Micronucleus Test

The *in vitro* micronucleus test can detect the induction of micronuclei in the cytoplasm of cells in the interphase stage of the cell cycle. Micronuclei form from acentric chromosome fragments (i.e., chromosome fragments lacking a centromere) or when whole chromosomes are unable to migrate to the cellular poles during anaphase prior to cell division. (OECD 487). Thus, the micronucleus assay can detect both structural and numerical chromosomal changes. It should be noted, however, that additional work is required to distinguish whether induced micronuclei have arisen from a clastogenic versus an aneugenic mechanism, *e.g.*, staining micronuclei to detect the presence of kinetochore proteins. The assay is typically performed with cell lines or primary cell cultures of human or rodent origin. The assay can be conducted with the addition of cytochalasin B which inhibits cytokinesis resulting in the formation of binucleated cells. The presence of binucleated cells, indicates that cells have undergone one round of mitosis, a necessary prerequisite for micronucleus formation.

Six studies evaluated glyphosate technical for its potential to induce micronuclei *in vitro* (Table 5.4). Four of the six studies were positive and the remaining two studies were equivocal. In a study by Koller *et al.* (2012), TR146 cells (derived from a human neck metastasis of buccal epithelial origin) were treated for 20 minutes with up to 20 mg/L (~ 0.12 mM) glyphosate (95%), the authors reported a statistically significant increase in binucleated cells with micronuclei at 15 (~ 0.09 mM) and 20 (~ 0.12 mM) mg/L, and also indicated significant apoptosis and necrosis at 20 mg/L. The short exposure period in this study was unusually short (20 minutes) and was conducted in a tumor cell line that had not been well characterized in regards to its degree of chromosomal instability and DNA damage and repair capacity. In another study, Roustan *et al.* (2014) reported positive findings +S9 only in CHO cells treated with glyphosate (unknown purity) at 10- 100 μ g/mL with little evidence of a dose response over that concentration range.

Two other studies evaluated glyphosate technical in human lymphocytes (Mladinic *et al.*, 2009a, 2009b). These studies used an exposure protocol that is different from the OECD recommendations for the *in vitro* micronucleus assay. OECD recommends that whole blood or isolated lymphocytes are cultured in the presence of a mitogen (*e.g.* phytohemagglutinin; PHA) prior to exposure of a test chemical in order to detect micronuclei formed via an aneugenic mechanism. However, in these two studies, blood cells were exposed to glyphosate for 4 hours, washed, and then treated with PHA to stimulate cell division. Both studies reported a statistically significant increase in micronucleated cells at 580 μ g/mL (~ 3.4 mM), but not at lower concentrations, following 4-hour exposures in the presence of S9. The frequency of micronucleated cells (+S9) ranged from 11.3 to 28.7 in one study (Mladinic *et al.*, 2009a) and 33.3 to 65.2 in the other study (Mladinic *et al.*, 2009b) over the 1000-fold concentration range. No statistically significant increases in micronucleated cells were seen in either study in the absence of S9 activation. When cells were evaluated with vital stains, cells treated with 580

µg/mL showed a significant ($p < 0.05$) increase in the percentage of cells undergoing apoptosis and necrosis compared to the negative controls.

Piesova et al. (2004, 2005) conducted two *in vitro* micronucleus studies using glyphosate technical (62%) up to 560 µM in bovine lymphocytes. In the 2004 study, bovine lymphocytes from two donors were treated for 24 or 48 hours without S9 metabolic activation, and for 2 hours (with and without S9 activation) or 48 hours (-S9) in the 2005 study. Both studies yielded similar results following 48-hour exposure to glyphosate. In both cases, the authors reported a weak induction of micronuclei in one donor at 280 µM and at 560 µM in the second donor. The induction was approximately 2-fold ($p < 0.05$), but with no clear dose response. No effects on micronuclei induction were seen at the 2- or 24-hour time points; however, with these early time points it is unlikely that one cell division has occurred during or after treatment. .

Table 5.3. <i>In vitro</i> Tests for Chromosome Aberrations in Mammalian Cells- Glyphosate Technical						
Test/Endpoint	Test System	Concentrations/ Conditions	Test Material Purity	Results	Reference	Comments
<i>In vitro</i> Chromosomal Aberration	Chinese hamster ovary (CHO) cells	4-10 µl/mL, ± S9	55.6% <i>Glyphosate trimesium salt</i>	Negative	Majeska (1985c) [MRID 00155530]	pH adjusted (7.4-7.6)
<i>In vitro</i> Chromosomal Aberration	Chinese Hamster lung (CHL) cells	±S9: 0, 250, 500, 1000 and 2000 µg/mL; 24 and 48 h treatment - S9; 6 h treatment ±S9 harvest 24 h	95.68%	Negative	Matsumoto (1995) [MRID 50017106]	Decline in pH noted at 500 and 1000 µg/mL.
<i>In vitro</i> Chromosomal Aberration	Chinese hamster lung (CHL) cells	-S9: 24 & 48-hr exposure: 0-1250 µg/mL; +S9: 0-1250 µg/mL	95.3%	Negative	Wright (1996) [MRID 49957410]	Excessive decrease in pH >1250 µg/mL
<i>In vitro</i> Chromosomal Aberration	Bovine lymphocytes	-S9 only: 0, 7, 85 and 170 µM; 72 h exposure	≥98%	Positive (all concs.)	Lioi <i>et al.</i> (1998b)	
<i>In vitro</i> Chromosomal Aberration	Bovine lymphocytes	±S9: 0, 28, 56, 140, 280, 560 and 1120 µM; 24 h exposure	62.0%	Negative	Sivikova and Dianovsky (2006)	Decreased MI and PI at ≥ 560 µM
<i>In vitro</i> Chromosomal Aberration	Human lymphocytes	±S9: 100-1250 µg/mL cultures analyzed; 68 & 92 h	95.6%	Negative	Fox (1998) [MRID 49961803]	Excessive decrease in pH >1250 µg/mL
<i>In vitro</i> Chromosomal Aberration	Human lymphocytes	-S9 only: 0, 5.0, 8.5, 17.0 and 51.1 µM; 72 h exposure	≥98%	Positive ≥ 8.5 µM	Lioi <i>et al.</i> (1998a)	No significant ↓ in MI observed.
<i>In vitro</i> Chromosomal Aberration	Human lymphocytes	-S9: 0, 200, 1200 and 6000 µM; 48 h exposure	96.0%	Negative	Manas <i>et al.</i> (2009)	No toxicity observed up to 6000 µM

¹ Study was cited in Kier and Kirkland (2013). Supplementary information about the study was provided online including test guideline, test material purity, control chemicals and summary data tables.

CA= chromosomal aberrations, MI= mitotic index, PI= proliferation index.

Table 5.4. *In vitro* Tests for Micronuclei Induction in Mammalian Cells- Glyphosate Technical

Test/Endpoint	Test System	Concentrations/ Conditions	Test Material Purity	Results	Reference	Comments
<i>In vitro</i> Cytokinesis Block Micronucleus Assay (with FISH analysis)	TR146 cells (human- derived buccal carcinoma cell line)	10, 15 and 20 mg/L; 20 minute exposure.	95%	Positive Statistically significant (p<0.05) increase in MN at 15 and 20 mg/L.	Koller <i>et al.</i> (2012)	Apoptosis and necrosis reported at 20 mg/L Also reported ↑ in NB and NPB
<i>In vitro</i> Cytokinesis Block Micronucleus Test	CHO-K1 cells	5 - 100 µg/mL, ±S9	Not stated	Negative –S9 Positive +S9 at 10- 100 µg/mL	Roustan et al., (2014)	No clear dose response
<i>In vitro</i> Cytokinesis Block Micronucleus Test	Bovine lymphocytes (2 donors)	0, 28, 56, 140, 280 and 560 µM 24 & 48 h exposure	62%	24 h: Negative 48 h: Equivocal ↑ MN at 280 µM only (donor A) ↑ MN at 560 µM only (donor B)	Piesova, 2004	No dose-response No significant decrease in CBPI observed.
<i>In vitro</i> Cytokinesis Block Micronucleus Test	Bovine lymphocytes (2 donors)	0, 28, 56, 140, 280 and 560 µM; 2 h (±S9) and 48 h (-S9) exposure	62%	2 h: Negative 48 h: Equivocal ↑ MN at 280 µM only (donor A) and at 560 µM only (donor B)	Piesova, 2005	No dose-response; No significant decrease in CBPI observed. Metabolic activation had no effect on MN formation after 2 h exposure.

Table 5.4. <i>In vitro</i> Tests for Micronuclei Induction in Mammalian Cells- Glyphosate Technical						
Test/Endpoint	Test System	Concentrations/ Conditions	Test Material Purity	Results	Reference	Comments
<i>In vitro</i> Cytokinesis Block Micronucleus Assay (with FISH analysis)	Human lymphocytes (treated with cytochalasin B)	4h treatment $\pm$ S9; 0.5, 2.91, 3.50, 92.8 and 580 μ g/mL; harvested 72 h	98.0%	Negative –S9 Positive +S9, $\uparrow$ MN at 580 μ g/mL, but not at 0.5-92.8 μ g/mL Also observed $\uparrow$ in NB at 580 μ g/mL ($\pm$ S9); $\uparrow$ NPB at 580 μ g/mL (+S9)	Mladinic <i>et al.</i> (2009a)	Cells were exposed to glyphosate and washed prior to treatment with PHA. Authors did not report being blind to treatment.
<i>In vitro</i> Cytokinesis Block Micronucleus Assay (with FISH analysis)	Human lymphocytes (treated with cytochalasin B)	4h treatment $\pm$ S9; 0.5, 2.91, 3.50, 92.8 and 580 μ g/mL	98%	Negative –S9 Positive +S9 $\uparrow$ MN at 580 μ g/mL, but not at 0.5 -92.8 μ g/mL $\uparrow$ apoptosis and necrosis at 580 μ g/mL (-S9); $\uparrow$ apoptosis at $\geq$ 2.91 μ g/mL and necrosis at 580 μ g/mL (+S9) $\uparrow$ in NB at 580 μ g/mL ($\pm$ S9) and NPB at 580 μ g/mL (+S9)	Mladinic <i>et al.</i> (2009b)	Cells were exposed to glyphosate and washed prior to treatment with PHA. Authors did not report being blind to treatment. .

CBPI= cytokinesis block proliferation index, FISH= fluorescent in situ hybridization; MN= micronuclei; NB= nuclear buds; NPB= nucleoplasmic bridges.

5.5 *In Vivo* Genetic Toxicology Tests

5.5.1 *In Vivo* Assays for Chromosomal Abnormalities

5.5.1.1 Mammalian Bone Marrow Chromosomal Aberration Assays

The *in vivo* mammalian bone marrow chromosomal assay detects the ability of a chemical to cause structural chromosomal damage in cells in the bone marrow. The assay is typically conducted in rodents (mouse or rat) and detects both chromosome-type and chromatid-type aberrations. Chromatid-type aberrations are expressed when a single chromatid break occurs and/or a reunion between chromatids, and chromosome-type aberrations result from damage expressed in both sister chromatids (OPPTS 870.5385). In this test, animals are exposed (typically via oral route or intraperitoneal injection) and sacrificed at sequential intervals. Prior to sacrifice, animals are treated with a spindle inhibitor such as colchicine or Colcemid® to arrest cells at metaphase. Chromosome preparations from the bone marrow are stained and scored for chromosomal aberrations. (OPPTS 870.5385). Generally, the optimal time to detect chromosomal aberrations in the bone marrow is 24 hours after treatment.

Three *in vivo* mammalian bone marrow chromosomal assays were conducted with glyphosate technical for regulatory purposes and all were negative (Table 5.8). In the first study, Sprague Dawley rats were administered glyphosate (98%) at 0 or 1000 mg/kg and the bone marrow was sampled at 6, 12 or 24 hours after dosing. No significant increase in bone marrow chromosomal aberrations were observed (Li, 1983b). In the second study, Swiss albino mice were treated twice by oral gavage (24 hours apart) with 0 or 5000 mg/kg glyphosate technical (96.8%) resulting in no significant increase in bone marrow chromosomal aberrations (Suresh, 1994). In a third study conducted with glyphosate trimesium salt, no increase in chromosomal aberrations were seen in the bone marrow of rats treated by oral gavage with up to 188 mg/kg (Majeska, 1982c).

5.5.1.2 Rodent Dominant Lethal Test

Dominant lethal mutations cause embryonic or fetal death. The induction of a dominant lethal mutation after exposure to a chemical indicates that the test chemical has affected the germinal tissue (sperm at some point in development, from stem cell to spermatocyte). Dominant lethal effects are considered to result from chromosomal damage (structural or numerical), but may also reflect gene mutations or systemic toxicity (OPPTS 870.5450, OECD 2016). In this test, male rodents are treated with the test material and mated with (untreated) virgin females. The females animals are sacrificed at an appropriate time and the uteri are examined to determine the number of implants, and live and dead embryos. Two dominant lethal studies were identified. One study was conducted in the rat (Suresh, 1992) where male rats were dosed by oral gavage with glyphosate up to 5000 mg/kg. The other study (Rodney, 1980) was conducted in male mice treated with up to 2000 mg/kg glyphosate (98.7%) by oral gavage. No significant increase in dominant lethal mutations were observed in either study (Table 5.5).

5.5.1.3 *In Vivo* Mammalian Erythrocyte Micronucleus Assays

The mammalian micronucleus test is the most commonly conducted *in vivo* test to detect clastogenic or aneugenic chemicals. The test identifies chemicals that induce micronuclei in proerythrocytes (progenitor cells) by assessing micronucleus frequency in immature erythrocytes (polychromatic erythrocytes, PCEs) sampled from the bone marrow or from the peripheral blood (reticulocytes). This test is typically conducted in mice or rats. When bone marrow erythroblasts develop into erythrocytes, the main nucleus is extruded following the final cell division (erythrocytes are the only mammalian cell that does not contain a nucleus). Any micronuclei formed after the final cell division may remain in the cytoplasm following extrusion of the main nucleus. The visualization of micronuclei is facilitated by the lack of a nucleus in these cells (OPPTS 870.5395, OECD 474). Micronuclei can originate from acentric chromosomes, lagging chromosome fragments, or whole chromosomes; thus, micronuclei are biomarkers of both altered chromosome structure or chromosome number. The assay is based on an increase in the frequency of micronucleated erythrocytes in treated animals, in either peripheral blood samples or bone marrow samples (OPPTS 870.5395). Additional mechanistic information on the formation of chromosomal abnormalities can be obtained from the incorporation of centromeric and telomeric fluorescent probes (FISH) assay. . According to EPA test guidelines, a single dose of the test substance may be used in this test if the dose is the maximum tolerated dose (MTD), a dose that produces some indication of bone marrow cytotoxicity (e.g., a reduction in the proportion of immature erythrocytes (PCEs) to total erythrocytes by >50%) or a maximum limit dose of 5000 mg/kg. The routes of administration for this test are typically oral or intraperitoneal injection and generally involve a single administration.

Glyphosate technical has been extensively evaluated for micronuclei induction in *in vivo* studies. Fourteen studies were conducted for regulatory purposes, four were identified from the open literature, and one study was conducted by the U.S. National Toxicology Program (NTP). This included nine studies with administration of glyphosate by the intraperitoneal (i.p.) route and 10 studies by the oral route. The findings are presented in Table 5.10. Of the nine i.p. studies, seven (Costa, 2008; Chruscielska *et al.*, 2000; Durward, 2006; Gava, 2000; Marques, 1999; Rank *et al.*, 1993 and Zaccaria, 1996) were negative. These studies tested doses as high as 2016 mg/kg (single and double administration) with sampling times at 24 and 48 hours post-dose. Two positive findings were reported when glyphosate technical was administered by i.p. Bolognesi *et al.* (1997) reported a significant increase in micronuclei in the bone marrow of male Swiss CD mice 24 hours after i.p. treatment with 300 mg/kg glyphosate technical (99.9%). The dose in this study was administered as ½ dose (150 mg/kg) injections 24 hours apart to 3 male mice. Manas *et al.* (2009) evaluated glyphosate technical (96%) in BALB/c male and female mice (5/sex/dose) administered 50, 100 or 200 mg/kg by two i.p. injections, 24 hours apart. The results showed a significant increase in micronucleated erythrocytes at 200 mg/kg, but not at 50 or 100 mg/kg. It should be noted that doses that resulted in the positive responses in these two studies were above the reported i.p. LD50 value (130 mg/kg) for glyphosate in mice (NTP 1992).

Glyphosate technical was also evaluated in nine micronucleus assays with administration by the oral route in mice and one in the rat. Eight of the nine oral studies in the mouse were negative for micronuclei induction. The single positive response was seen in female mice treated with two 5000 mg/kg (limit dose) doses, 24 hours apart with bone marrow sampling at 24 hours post-dose (Suresh, 1993b). No increase was observed at lower doses (50 and 500 mg/kg) in females or at any dose in males. The eight negative oral studies in mice included single dose administrations of 5000 mg/kg and bone marrow analysis at 24, 48, and/or 72 hours (Jensen, 1991c; Fox and Mackay, 1996) and one or two administrations of glyphosate technical with top doses between 30 and 2000 mg/kg (Honarvar, 2005; Honarvar, 2008; Jones, 1999; and Zoriki-Hosmi, 2007). It should be noted that evaluations at 48 and 72 hours post dose may be too late to detect chemically-induced micronucleated PCEs in the bone marrow as these cells may have already migrated into the peripheral blood. No significant increase in micronucleated erythrocytes were seen in male or female mice following 13-weeks of dietary (feed) administration of glyphosate technical at doses up to 3393 mg/kg/day (NTP, 1992). In the single study that evaluated micronuclei induction in rats, glyphosate technical did not induce significant induce micronuclei in CD1 rats treated by oral gavage at doses up to 2000 mg/kg (Flügge, 2009b). When glyphosate trimesium salt was evaluated, no increase in micronuclei induction was seen in mice treated orally up to 1100 and 800 mg/kg in males and females, respectively (Majeska, 1987).

Table 5.5. <i>In Vivo</i> Tests for Chromosomal Aberrations in Mammals- Glyphosate Technical.							
Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Chromosomal Aberration Test	Sprague Dawley rats (males and females)	Intraperitoneal injection; sampled at 6, 12 and 24 h after treatment	0, 1000 mg/kg (6/sex/dose/sampling time)	98%	Negative	Li (1983b) [MRID 00132683]	No toxicity observed. A separate study using ¹⁴ C-glyphosate showed that glyphosate reaches BM 0.5 h after dosing with ½ life elimination at 7.6 h. Peak BM value was 400 ppm, corresponding to 2000 ppm plasma value.
Bone Marrow Chromosomal Aberration Test	Sprague Dawley rats (males and females) Vehicle: distilled water	Oral gavage, sampling after 6, 12, 24, 48 h and 5 d	0, 21, 63 and 188 mg/kg	58.5% <i>Glyphosate trimesium salt</i>	Negative	Majeska (1982c) [MRID 00132176]	
Bone Marrow Chromosomal Aberration Test	Swiss Albino mice (males and females) Vehicle: peanut oil	Oral gavage (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 5000 mg/kg (5/sex/dose)	96.8%	Negative	Suresh (1994) [MRID 49987408]	Significant (p<0.05) decrease in bw of females at high dose.
Rodent Dominant Lethal Test	CD-1 mice Each dosed male mated with 2 females/week for 8 weeks	Oral gavage	0, 200, 800, and 2000 mg/kg	98.7%	Negative	Rodwell (1980) [MRID 00046364]	
Rodent Dominant Lethal Test	Wistar rat Each dosed male mated with 1 female/week for 10 weeks	Oral gavage	0, 200, 100 and 5000 mg/kg	96.8%	Negative	Suresh (1992) [MRID 49987404]	

Table 5.6. In Vivo Tests for Micronuclei Induction in Mammals- Glyphosate Technical.

Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Micronucleus Test	Swiss CD1 mice (males only)	Intraperitoneal injection; 2 injections of half the dosage of 300 mg/kg 24 h apart; sampling at 6 and 24 h	0, 300 mg/kg (3/dose)	99.9%	Positive Stat significant increase in MN at 24 h	Bolognesi <i>et al.</i> (1997)	Material and methods indicate 3 animals/dose; however, Table 1 of article indicates 4 animals were evaluated.
Bone Marrow Micronucleus Test	Balb C mice (males and females) Vehicle: Saline	Intraperitoneal Injection (two injections, 24 h apart); sampling after 24 h (last treatment)	0, 50, 100, and 200 mg/kg (5/sex/dose)	96%	Positive ↑MN at 200 mg/kg, but not at 50 or 100 mg/kg	Manas <i>et al.</i> (2009)	No significant signs of toxicity observed.
Bone Marrow Micronucleus Test	C3H mice (males only) Vehicle: water	Intraperitoneal Injection (single treatment); sampling after 24, 48 and 72 h	0, 300 mg/kg	Not reported	Negative	Chruscielska <i>et al.</i> (2000)	
Bone Marrow Micronucleus Test	Swiss Albino mice (males and females) Vehicle: corn oil	Intraperitoneal Injection (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 15.62, 31.25, and 62.5 mg/kg (5/sex/dose)	980 g/kg Glyphosate technical	Negative#	Costa (2008) ¹	OECD guideline 474 #Was not tested up to limit dose and did not demonstrate that compound was tested up to toxic dose. No mention of BM toxicity or clinical signs.

Table 5.6. In Vivo Tests for Micronuclei Induction in Mammals- Glyphosate Technical.

Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Micronucleus Test	CrI:CD-1TM(ICR) BR mice (males only ¹) Vehicle: PBS	Intraperitoneal Injection (single treatment); sampling after 24 and 48 h (high dose only)	0, 150, 300 and 600 mg/kg (7/dose)	95.7%	Negative	Durward (2006) [MRID 49957411]	Clinical signs reported at ≥ 150 mg/kg. Significant $\downarrow$ in %PCEs reported at 24 h in 600 mg/kg group. $\uparrow$ in MN PCEs observed at 600 mg/kg (1.9 ± 0.7 vs. 1.0 ± 1.2 control; $p < 0.05$), at 24 h, but not 48 h, within historical control range.
Bone Marrow Micronucleus Test	Swiss Albino mice (males and females) Vehicle: water	Intraperitoneal Injection (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 1008, 2016, and 3024 mg/kg 5/sex/dose	612.7 g/kg (glyphosate technical Nufarm)	Negative	Gava (2000) ¹	LD50 was 4032 mg/kg Mortality observed in 1 animal at high dose (only 4 m/f scored for MPCEs). No effect on PCE/NCE.
Bone Marrow Micronucleus Test	Swiss Albino mice (males and females) Vehicle: water	Intraperitoneal Injection (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 187.5, 375 and 562.5 mg/kg 5/sex/dose	954.9 g/kg (glyphosate technical Nufarm)	Negative	Marques (1999) [MRID 49957412]	LD50 was 750 mg/kg No significant signs of toxicity observed in main study
Bone Marrow Micronucleus Test	NMRI-Bom mice	Intraperitoneal Injection (single treatment); sampling after 24 h (all doses) and 48 h (150 and 200 mg/kg)	0, 150, and 200 mg/kg (5/sex/dose)	glyphosate isopropyla mine (purity not specified)	Negative	Rank <i>et al.</i> (1993)	

Table 5.6. In Vivo Tests for Micronuclei Induction in Mammals- Glyphosate Technical.

Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Micronucleus Test	Swiss albino mice (males and females)	Intraperitoneal Injection (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 68, 137, and 206 mg/kg (	360 g/L	Negative	Zaccaria (1996) [MRID 49961501]	Doses selected were reported as corresponding to 25, 50 and 75% LD ₅₀
Bone Marrow Micronucleus Test	CD-1 mice (males and females) Vehicle: saline	Oral gavage (single treatment); sampling after 24 and 48 h	0, 5000 mg/kg 5/sex/dose	95.6%	Negative	Fox and Mackay (1996) [MRID 44320619]	No significant signs of toxicity observed
Bone Marrow Micronucleus Test	NMRI mice (males and females) Vehicle: PEG 400	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 mg/kg 5 sex/dose	97.73%	Negative	Honarvar (2005) ¹	OECD guideline 474 No significant signs of toxicity observed
Bone Marrow Micronucleus Test	NMRI mice (males only) Vehicle: 0.5% carboxymethylcellulose	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 mg/kg (5/dose)	99.1%	Negative	Honarvar (2008) [MRID 49961802]	No significant signs of toxicity observed
Bone Marrow Micronucleus Test	NMRI mice (males and females) Vehicle: 0.5% carboxymethylcellulose	Oral gavage (single treatment); sampling after 24, 48 and 72h	0, 5000 mg/kg; 5/sex/dose	98.6%	Negative	Jensen (1991c) [MRID 49961503]	No significant signs of toxicity observed
Bone Marrow Micronucleus Test	CD-1 mice (males only) ¹ Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h	0, 2000 mg/kg 5/dose	59.3% potassium glyphosate salt	Negative	Jones (1999) ¹	OECD guideline 474 No significant signs of toxicity observed
Bone Marrow Micronucleus Test	Swiss albino mice; (males and females)	Oral gavage (2 treatments, 24 h apart); sampling	0, 50, 500, 5000 mg/kg 5/sex/dose	96.8% glyphosate acid	Positive in females at 5000	Suresh (1993b) [MRID 49987407]	No significant signs of toxicity observed

Table 5.6. In Vivo Tests for Micronuclei Induction in Mammals- Glyphosate Technical.

Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
	Vehicle: peanut oil	after 24 h (last treatment)			mg/kg only. Negative in males at all doses		
Bone Marrow Micronucleus Test	Swiss mice (males only) Vehicle: corn oil	Oral gavage (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 8, 15 and 30 mg/kg (6/dose)	980.1 g/kg	Negative	Zoriki Hosomi (2007) [MRID 50000901]	OECD guideline 474 No significant signs of toxicity observed
Bone Marrow Micronucleus Test	CD-1 mice (males and females) Vehicle: distilled water	Oral gavage , Sampling 24, 48 and 72 h after treatment	Males: 0, 700, 900 and 1100 mg/kg Females: 0, 400, 600 and 800 mg/kg	55.3% <i>Glyphosate trimesium salt</i>	Negative	Majeska (1987) [MRID 40214004]	
Bone Marrow Micronucleus Test	B6CF3 Mice (males and females)	Oral (dietary). MN assay conducted following 13 week feed study.	0, 205/213, 410/421, 811/844, 1678/1690 and 3393/3393 mg/kg (m/f) (10/sex/dose)	99%	Negative	NTP (1992)	
Bone Marrow Micronucleus Test	CD Rats (males and females) Vehicle: 0.8% hydroxypropylmethyl cellulose	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 mg/kg (5/sex/dose)	98.8%	Negative	Flügge (2009b) ¹	OECD guideline 474 No significant signs of toxicity observed

¹ Study was cited in Kier and Kirkland (2013). Supplementary information about the study was provided online including test guideline followed, test material purity, control chemicals and summary data tables.

²Only males tested; report indicated that there were no difference between sexes seen in range finding study.

CA= chromosomal aberrations, MPCE= micronucleated polychromatic erythrocytes, NCE= normochromatic erythrocytes, PCE=polychromatic erythrocytes.

5.6 Additional Genotoxicity Assays Evaluating Primary DNA Damage

There are a number of genotoxicity assays that evaluate primary DNA damage, but do not measure the consequence of the genetic damage (*i.e.*, mutation or chromosomal damage). As discussed in the Guidance Document on Revisions to OECD Genetic Toxicology Test Guidelines (OECD 2015), the endpoints measured in primary DNA damage tests such as DNA adducts, comet assay, or unscheduled DNA synthesis may lead to cell death or may initiate DNA repair, rather than a mutation. These types of assays can, however, provide mechanistic information when interpreting positive findings in other genotoxicity tests or when determining whether a chemical is acting through a mutagenic mode of action. Additionally, indirect mechanisms of DNA damage such as oxidative DNA damage can be detected by these test systems. Oxidative damage results from oxidative stress, which occurs when there is a disturbance in the balance between the production of reactive oxygen species (ROS) and antioxidant defense systems. Normal cellular metabolism is a source of endogenous reactive oxygen species that accounts for background levels of oxidative damage in normal cells. Some types of oxidative damage are repairable while others lead to serious consequences in the cell. (Cooke et al, 2003). The various assays evaluating primary DNA damage in glyphosate technical are presented in Table 5.7 Details of the findings are discussed below.

Glyphosate technical is not electrophilic and is not considered to be DNA-reactive. In a study to evaluate the potential for glyphosate to directly interact with DNA, Peluso et al. (1998) reported that glyphosate technical did not form DNA adducts in mice when tested up to 270 mg/kg via i.p. Bolognesi et al. (1997) reported an increase in the oxidative damage biomarker 8-hydroxydeoxyguanosine (8-OHdG) in the liver 24 h after i.p. injection of 300 mg/kg in mice. No increase in 8-OHdG was seen in the kidney with glyphosate technical. The dose in this study was high (300 mg/kg) for an i.p. injection and within the i.p. LD₅₀ range (134- 545 mg/kg) that has been reported elsewhere (WHO, 1994).

The comet assay, also known as single cell gel electrophoresis (SCGE), is a sensitive and rapid method to detect DNA strand breaks in individual cells. In this assay, individual cells are embedded in agarose. The cells are then lysed (which digests the cellular and nuclear membranes) and the DNA is allowed to unwind under alkaline or neutral conditions. During electrophoresis, chromatin (which is in a supercoiled state) that has undergone steric relaxation due to DNA damage migrates away from the nucleoid (nucleus) toward the anode, yielding images that resemble a comet. The intensity of the comet tail relative to the comet head reflects the amount of DNA breakage (Tice et al., 2000; Collins et al., 2008). The comet assay can detect single and double strand breaks resulting from direct interactions with DNA, alkali labile sites, or transient DNA breaks resulting during DNA excision repair. These types of strand breaks may be, (a) repaired with no persistent effect, (b) be lethal to the cell or (c) be fixed as a mutation (OECD TG 489). DNA strand breaks in the comet assay can be measured by endpoints such as percent tail DNA (also referred to as % tail intensity), tail length, and tail moment. However, % tail DNA is the recommended metric for evaluating and interpreting results using this assay (OECD TG 489).

The five studies that evaluated glyphosate technical using the comet assay are summarized in Table 5.12. Two of the studies were conducted using tumor cell lines. Koller et al. (2012) reported positive comet effects (increased tail intensity) in a human buccal carcinoma cell line (TR146) following a 20-minute treatment with ≥ 20 mg/L (~ 0.118 mM) glyphosate. Although no evidence of cytotoxicity was reported in this study, the authors did report an increase in apoptosis and necrosis at the same concentrations (≥ 20 mg/L) when the same cell line was tested for *in vitro* micronuclei induction (discussed previously). In a study using Hep-2 cells (presumably a HeLa cell derivative), Manas et al. (2009) reported a statistically significant increase in mean tail length, and tail intensity at all concentrations (3.0-7.5 mM) tested. In a comet study conducted on human lymphocytes, Alvarez-Moya et al. (2014) reported significant increases in tail length only (but not % tail DNA) following treatment with glyphosate concentrations of 0.7-700 μ M. Mladinic et al. (2009a) evaluated DNA damage in non-dividing human lymphocytes ($\pm$ S9) following treatment from 0.5 to 580 μ g/mL using the standard alkaline comet method and a modified comet method that detects DNA damage due to oxidative damage (human 8-hydroxyguanine DNA-glycosylase, hOGG1 comet method). In this study, the authors reported statistically significant increases in tail intensity at 3.5 μ g/mL and higher in the absence of S9, with significance only at 580 μ g/mL (~ 3.4 mM) in the presence of S9 using the alkaline method. This concentration also resulted in increased apoptosis and necrosis as well as an increase in plasma total antioxidant capacity (TAC) and changes in plasma lipid peroxidation (thiobarbituric reactive substances, TBARs); however, only a dose-related increase in tail length (not % tail DNA) was observed at 580 μ g/mL (+S9) using the hOGG1 method. When the Manas et al. (2013) evaluated blood and liver cells following a 14 day drinking water study in mice treated with 40 and 400 mg/kg/day glyphosate, significant increases in tail intensity, tail length and tail moment were reported were observed at both doses in both tissues (except for DNA tail intensity in liver at 40 mg/kg); however, there were no substantial effects on oxidative stress measurements suggesting that DNA damage reported may not be due to oxidative damage.

The Unscheduled DNA Synthesis (UDS) test with mammalian liver cells *in vitro* identifies substances that induce DNA repair after excision and removal of a segment of damaged DNA. The test is typically conducted in liver cells, which have relatively few cells in the S-phase of the cell cycle. The assay measures the incorporation of radiolabeled nucleotide [3 H]-thymidine into DNA during the repair process in non-S phase cells. (OPPTS 870.5555). Substances that produce either a statistically significant dose-related increase or statistically significant and reproducible increase in 3 H-TdR incorporation in at least one test point are considered to be positive in this test. A UDS study that evaluated glyphosate technical in rat primary hepatocytes was negative (Williams, 1978). Glyphosate technical was also negative in a DNA repair test conducted in bacteria (Rec-A test) (Shirasu, 1978).

In an alkaline elution assay, which detects single strand DNA breaks, Bolognesi et al. (1997) reported an increase in single strand breaks (i.e. increased DNA elution rate) in the liver and kidney 4 hours after a single i.p. injection of 300 mg/kg. The elution rate returned to control

levels at 24 hours. Glyphosate technical was also negative in a DNA repair test conducted in *Bacillus subtilis* H17 (rec⁺) and M45 (rec⁻) bacterial Rec-A test (Shirasu, 1978).

Finally, the sister chromatid exchange (SCE) test is an assay that can measure the consequence of primary DNA damage. The mechanism(s) of action for chemical induction of SCE is unclear. The SCE assay detects the exchange of DNA between two sister chromatid arms within a single chromosome. The assay can be performed *in vitro* or *in vivo*. Following exposure, cells/animals are treated with bromodeoxyuridine (BrdU) to allow for the differentiation of the two sister chromatids (harlequin staining) and prior to harvest are treated with a spindle inhibitor to accumulate cells in metaphase. The chromosome preparations are then stained and analyzed for SCEs (OPPTS 870.5900, 870.5915). The SCE studies that evaluated glyphosate technical are also presented in Table 12. Positive SCE findings were reported in all four studies; two evaluating bovine lymphocytes (Lioi, 1988b, Sivikova and Dianovksy, 2006) and two studies evaluating human lymphocytes (Lioi, 1988a; Bolognesi et al., 1997). In all four studies the induction did not demonstrate a clear dose response.

Additionally, although it is recognized that mechanisms other than genotoxicity may be involved in cell transformation, glyphosate trimesium salt was evaluated in the Balb/3T cell transformation assay (an *in vitro* tumor formation assay) and was negative up to 5.0 mg/ml (Majeska, 1982b).

Table 5.7 Assays for Detecting Primary DNA Damage- Glyphosate Technical.							
Test/Endpoint	Test System	Route of Administration	Doses/ Concentrations	Test Material Purity	Results	Reference	Comments
DNA Adducts ³² P-postlabeling	Swiss CD1 mice (males and females) Liver and kidney evaluated	Intraperitoneal injection; 24 h exposure	0, 130 and 270 mg/kg	Not reported	Negative	Peluso <i>et al.</i> (1998)	
DNA oxidative damage: 8-OHdG formation	Swiss CD-1 mice (males) liver and kidney evaluated	Intraperitoneal injection (single dose); sampling 4 and 24 h after injection	0, 300 mg/kg (3/dose)	99.9%	Kidney: negative Liver: positive (24 h)	Bolognesi <i>et al.</i> (1997)	
Single-cell gel electrophoresis (SCGE) assays- Comet assay	TR146 cells (human-derived buccal epithelial cell line).	NA (<i>in vitro</i>)	-S9: 10-2000 mg/L; 20 minute exposure.	95%	Positive Increased DNA migration at >20 mg/L	Koller <i>et al.</i> (2012)	Also measured multiple cellular integrity parameters to assess cytotoxicity. No clear evidence of cytotoxicity seen except for increase in enzyme activity (indicative of membrane damage) in LDHe (extracellular lactate dehydrogenase) assay at >80 mg/L. No mention of monitoring pH
Single-cell gel electrophoresis (SCGE) assays- Comet assay	Hep-2 cells	NA (<i>in vitro</i>)	0, 3, 4.5, 6, 7.5, 9, 12 and 15 mM	96%	Positive Stat. significant increase in mean tail length, and tail intensity at all concs.	Manas <i>et al.</i> (2009)	The authors did not report a source for the Hep-2 cells. The agency presumes that this is a HeLa derived cervical carcinoma cell line.

Table 5.7 Assays for Detecting Primary DNA Damage- Glyphosate Technical.							
Test/Endpoint	Test System	Route of Administration	Doses/ Concentrations	Test Material Purity	Results	Reference	Comments
Single-cell gel electrophoresis (SCGE) assays- Comet assay	Human lymphocytes	NA (<i>in vitro</i>)	0, 0.7, 7, 70, 700 µM	96%	Positive at all doses (increase in tail length only)	Alvarez-Moya <i>et al.</i> , (2014)	Issues were identified with this study resulting in a low quality ranking. These include: 1) blood was washed with PBS and then held at 4° C for an indeterminate amount of time before exposure to glyphosate. (2) Cells were treated for 20 hours at room temperature. (3) The same amount of damage was reported across 2 orders of magnitude concentration.
Single-cell gel electrophoresis (SCGE) assays- Comet assay	Human lymphocytes; ±S9 Alkaline and hOOG1 Comet assays performed	NA (<i>in vitro</i>)	0, 0.5, 2.91, 3.5, 92.8 and 580 µg/mL	98%	Positive ±S9	Mladinic <i>et al.</i> (2009a)	<u>The alkaline comet assay</u> -S9: ↑ in mean tail length at 580 µg/mL and ↑ in tail intensity at ≥ 3.5 µg/mL). +S9: ↑ DNA tail length at ≥3.5 µg/mL. Tail intensity ↑ only at 580 µg/mL <u>hOOG1 comet assay:</u> -S9 no effect on tail length, ↑tail intensity only at 3.50 µg/mL +S9: ↑ tail length at 580 µg/mL, no effect on tail intensity compared to controls at any conc.

Table 5.7 Assays for Detecting Primary DNA Damage- Glyphosate Technical.							
Test/Endpoint	Test System	Route of Administration	Doses/ Concentrations	Test Material Purity	Results	Reference	Comments
Single-cell gel electrophoresis (SCGE) assays- Comet assay with oxidative stress measures	Balb/C mice; evaluated blood and liver	Drinking water (14 days)	0, 40, and 400 mg/kg	96%	Positive Blood and liver at both doses	Manas et al. (2013)	Only minor effects seen on oxidative stress measurements (TBARs, SOD, CAT)
Sister Chromatid Exchange (SCE)	Bovine lymphocytes (3 donors)	NA (<i>in vitro</i>)	-S9: 0, 17, 85 and 170 µM; 72 h exposure	≥98%	Positive Significant (p>0.05) increase in SC/cell at all concentrations	Lioi (1998b)	1.8-, 2.1-, 1.6-fold increases, respectively
Sister Chromatid Exchange (SCE)	Human lymphocytes	NA (<i>in vitro</i>)	-S9: 0, 5, 8.5, 17 and 51 µM; 72 h exposure	≥98%	Positive Significant (p>0.05) increase in SCE/cell at ≥ 8.5 µM	Lioi (1998a)	1.9-, 2.8-, and 2.6-fold increase at 8.5, 17 and 51 µM, respectively
Sister Chromatid Exchange (SCE)	Human lymphocytes	NA (<i>in vitro</i>)	-S9: 0, 0.33, 1,3 and 6 mg/mL; 72 h exposure	99.9%	Positive	Bolognesi <i>et al.</i> (1997)	Very limited information was provided on the methods used in this paper. Authors report a dose –dependent increase in SCE frequency; however, no statistical analysis for dose response was reported. Data presented graphically with no error bars.
Sister Chromatid Exchange (SCE)	Human lymphocytes	NA (<i>in vitro</i>)	28, 56, 140, 280, 560 and 1120 µM; 24 h exposure ±S9	62%	Positive	Sivikova and Dianovsky (2006)	The increases in SCEs observed did not show a clear concentration related increase across a 40-fold increase in the concentrations tested

Table 5.7 Assays for Detecting Primary DNA Damage- Glyphosate Technical.

Test/Endpoint	Test System	Route of Administration	Doses/ Concentrations	Test Material Purity	Results	Reference	Comments
Alkaline elution assay- DNA single strand breaks	Swiss CD-1 mice (males) liver and kidney evaluated	Intraperitoneal injection (single dose); sampling 8 and 24 h after injection	0, 300 mg/kg (3/dose)	99.9%	Positive (Increased elution rate) at 4 hours in liver and kidney At 24 h, elution rate returned to control levels	Bolognesi <i>et al.</i> (1997)	Return to control values may indicate DNA repair or reflect rapid elimination of compound
DNA Repair Test (Rec-A test)	<i>B. subtilis</i> H17 (rec+) and M45 (rec-)	NA (<i>in vitro</i>)	20-2000 µg/disk	98.4%	Negative	Shirasu (1978) [MRID 00078619]	
Unscheduled DNA synthesis (DNA repair)	F-344 rat primary hepatocytes	NA (<i>in vitro</i>)	0, 0.0125, 0.0625, 0.125, 0.6.5, 1.25, 12.5, 125 µg/mL	98%	Negative	Li and Long (1988)	
Cell Transformation Assay	BALB/3T cells	NA (<i>in vitro</i>)	0.313-5.0 mg/mL	90% Glyphosate trimesium salt	Negative	Majeska (1982b) [MRID 00126616]	

h- hour; CAT= catalase, G6PD= glucose 6-phosphate dehydrogenase, NA= not applicable, hOOG1 =, TBARs= thiobarbituric acid reactive substances, SOD= superoxide dismutase

5.7 Summary and Discussion

The genotoxic potential of glyphosate has been extensively investigated using a variety of test systems and genetic endpoints. This assessment focuses only on test systems that the agency considered relevant for assessing genotoxic risks in humans. The totality of the genetic toxicology information was evaluated using a weight of evidence approach to determine the genotoxic potential of glyphosate. This involves the integration of *in vitro* and *in vivo* results as well as an overall evaluation of the quality, consistency, reproducibility, magnitude of response, dose-response relationship and relevance of the findings. In the weight of evidence analysis, studies evaluating endpoints that measured gene mutations and chromosomal aberrations (i.e. permanent DNA damage) were given more weight than endpoints reflecting DNA events that may be transient or reversible such as primary DNA damage (e.g., comet assays). *In vivo* studies in mammals were given the greatest weight and more weight was given to doses and routes of administration that were considered relevant for evaluating genotoxic risk based on human exposure to glyphosate. Also, since the molecular mechanisms underlying the observation of SCEs are unclear and thus, the consequences of increased frequencies of SCEs are unclear, the data from this test were given low weight in the overall analysis. A summary of the various lines of evidence of considered in the weight of evidence evaluation for the genotoxic potential of the active ingredient glyphosate is presented below.

Evidence of primary DNA damage

Glyphosate technical is not considered to be electrophilic and did not induce DNA adducts in the liver or kidney at an i.p. dose of 270 mg/kg. However, evidence of DNA strand breaks were reported in a number mammalian cell studies using the comet assay. Additionally, transient increases in alkali labile sites in the liver and kidney of mice and an induction of 8-OHdG in DNA were seen in the livers of mice following i.p. injections with 300 mg/kg glyphosate. These effects were seen at high doses for the i.p. route in mice (LD₅₀ for mouse =130 mg/kg; NTP, 1992). However, due to technical limitations identified in a number of these studies (e.g. use of cancer cell lines that have not been well-characterized, atypical exposure protocols and no indication of blind to treatment), caution should be exercised in interpreting the results.

In vitro mutations

Glyphosate technical was negative in all 39 studies for mutagenicity in bacteria. In the four studies that tested for gene mutations in mammalian cells *in vitro*, no increase in mutations were observed.

In vitro chromosomal alterations

Mixed results were observed in studies evaluating *in vitro* chromosomal alterations with glyphosate treatment. Three SCE studies reported positive findings (Lioi, 1998a, b; Bolognesi *et al.*, 1997) bovine and human lymphocytes. As stated previously, low weight is given to SCE results in the overall analysis given the uncertainty regarding the consequence of increases in the frequencies of SCEs. The SCE responses were weak and not concentration dependent. Eight of the 10 studies measuring *in vitro* chromosomal aberrations were negative. The two positive

findings were reported by Lioi et al., one study was conducted with bovine lymphocytes and the other with human lymphocytes. The authors reported positive findings in these studies at concentrations much lower than four other studies that reported negative results using the same cell types. Additionally, in both studies, Lioi et al. used an atypical exposure protocol of 72 hours which is very long for analyzing one round of mitosis. Furthermore, in both studies, nearly the same level effect for aberration frequency and percent of cells with aberrations were observed for the same concentrations of glyphosate and the two other chemicals tested in those experiments.

Four of the six studies evaluating micronuclei induction *in vitro* were positive and two showed equivocal results. Three of the positive responses required S9 activation, two conducted with human lymphocytes and one conducted with CHO cells. The remaining positive micronucleus study was conducted using a TR146 cells which is a tumor cell line derived from human buccal mucosa. The authors state that this cell line had not been previously used for genotoxicity testing. It is difficult to interpret any genotoxicity findings conducted in a tumor cell line that has not been well-characterized regarding its DNA damage response and repair capacity, and its degree of chromosomal instability.

Glyphosate was negative in all three L5178Y mouse lymphoma cell studies which may detect chromosomal damage in addition to mutations.

Mammalian *in vivo* chromosomal alterations

All three *in vivo* mammalian studies evaluating chromosomal aberrations with glyphosate technical were negative. Two studies were conducted in rats (i.p. and oral) and one was conducted in mice (oral). In addition glyphosate was also negative in a rodent dominant lethal test. Glyphosate was negative in 15 of the 19 bone marrow micronucleus studies evaluated. In two of the positive studies, glyphosate technical was administered by i.p. injection. In these studies, the authors reported positive findings at doses of 200-300 mg/kg. Based on the available ADME data for glyphosate, assuming 30% oral absorption, an oral dose of ~700-1000 mg/kg would be needed to achieve a dose of 200-300 mg/kg in the blood. Seven other i.p. studies in mice reported no increase in micronuclei induction at doses up to 3000 mg/kg. The remaining positive finding was reported in an oral gavage study in mice where an approximately 2-fold increase in micronuclei were reported in females only at a dose of 5000 mg/kg, which is considerably higher than the current guideline recommended limit dose of 2000 mg/kg. The effect was not seen in the 7 other oral gavage studies in mice when glyphosate was tested at similar doses. In addition, glyphosate was negative for micronuclei induction following a 13 week dietary study with a dose up to approximately 3000 mg/kg/day. A negative finding was also reported in the only study that evaluated *in vivo* micronuclei induction in the rat using doses up to 2000 mg/kg.

In a meta-analytic review of micronuclei frequency across mammalian and non-mammalian species (primarily fish, amphibians, reptiles and plants), Ghisi et al. (2016), not surprisingly, reported that different responses were observed when comparing mammalian results to phylogenetically distant non-mammalian species for micronuclei induction. Their analyses included most, but not all, of the mammalian studies that the agency evaluated and determined to

be negative for micronuclei induction. The authors reported a statistically significant increase in micronuclei by the i.p. route across the studies in the data set they considered; however, when glyphosate was administered by the oral route (which is the most physiologically relevant route for human exposure to glyphosate), no significant difference was observed.

Conclusion for Glyphosate

The overall weight of evidence indicates that there is no convincing evidence that glyphosate induces mutations *in vivo* via the oral route. When administered by i.p. injection, the micronucleus studies were predominantly negative. In the two cases where an increase in micronuclei were reported via this route, the effects occurred above the reported i.p. LD50 for mice and were not observed in other i.p. injection studies at similar or higher doses. While there is limited evidence genotoxic for effects in some *in vitro* experiments, *in vivo* effects were given more weight than *in vitro* effects particularly when the same genetic endpoint was measured, which is consistent with current OECD guidance. The only positive findings reported *in vivo* were seen at relatively high doses that are not relevant for human health risk assessment.

6.0 Data Integration & Weight-of-Evidence Analysis Across Multiple Lines of Evidence

6.1 Background

In 2010, OPP developed a draft “Framework for Incorporating Human Epidemiologic & Incident Data in Health Risk Assessment” which provides the foundation for evaluating multiple lines of scientific evidence (U.S. EPA, 2010). OPP’s draft framework is consistent with updates to the World Health Organization/International Programme on Chemical Safety MOA/human relevance framework, which highlights the importance of problem formulation and the need to integrate information at different levels of biological organization (Meek et al, 2014).

One of the key components of the agency’s draft framework is the use of modified Bradford Hill Criteria (Hill, 1965) like those described in the 2005 Guidelines for Carcinogen Risk Assessment. These criteria are used to evaluate the experimental support considers such concepts as strength, consistency, dose response, temporal concordance and biological plausibility in a weight-of-evidence analysis.

6.2 Dose-Response and Temporal Concordance

Given the lack of consistent positive findings particularly at doses < 1000 mg/kg/day across the lines of evidence, lack of mechanistic understanding, and lack of biological activity in mammalian systems to the parent compound glyphosate, there are few data to assess key events in the biological pathway and any associated temporal or dose concordance. Temporal concordance can be assessed using the experimental animal studies and epidemiological studies that evaluated exposure prior to outcomes. Similarly, dose concordance can be assessed using findings of apical outcomes in experimental animal studies, as well as epidemiological studies that utilize exposure metrics that are stratified by the number of exposure days.

A prospective cohort study is designed to collect exposure information prior to the development of cancer. As such, exposure is known to occur before the outcome. In De Roos *et al.* (2005), a prospective cohort study, no association was observed between glyphosate exposure and numerous cancer subtypes in the AHS cohort. Although the median follow-up time following recruitment into the cohort was approximately 7 years, it does not represent the amount of time subjects were exposed. Study participants provided pesticide exposure information prior to enrollment in the study and this information was used to evaluate has cumulative lifetime days of exposure and intensity-weighted cumulative days of exposure. An updated analysis of the AHS cohort is anticipated with a longer follow-up period, which includes the time period after the introduction of glyphosate-tolerant crops and the subsequent substantial increase in glyphosate use. The updated AHS cohort analysis will further elucidate the impact of increased glyphosate use due to glyphosate-tolerant crops. In De Roos *et al.* (2005), effect estimates did not increase across categories of increasing exposure for almost all cancer types, including NHL, in the prospective cohort study.

Two case-control studies evaluating the risk of NHL (Eriksson *et al.*, 2008 and McDuffie *et al.*, 2001) observed increased effect estimates in the highest exposure categories analyzed. Eriksson *et al.* (2008) found a greater effect estimate for subjects with >10 days (based on the median days of exposure among controls) and >10 years of exposure (for latency analysis) when compared to subjects with ≤10 days and 1-10 years of exposure, respectively; however, this analysis did not appear to adjust for co-exposures to other pesticides. By dividing the total number of exposed cases and controls using these exposure metrics, wider confidence intervals were observed indicating reduced power from smaller sample sizes. This may indicate that a longer follow-up time is needed to detect the risk for NHL; however, given the latency analysis of NHL was limited to Eriksson *et al.* (2008) and lack of NHL latency understanding in general, further studies are needed to determine the true latency of NHL. McDuffie *et al.* (2001), stratifying based on the average number of days per year of exposure, observed similar effect estimates in the lower exposure category (>0 and ≤2 days/year) while a greater effect estimate was observed in the highest exposure category (>2 days/year). The results from these two case-control studies conflict with the results observed in the cohort study (De Roos *et al.*, 2005), where no dose-response was seen across three exposure categories (stratified by tertiles; however, the case-control studies did not adjust for co-exposure to other pesticides. It is also difficult to make conclusions regarding dose-response with only two exposure categories used for the analyses by Eriksson *et al.* (2008) and McDuffie *et al.* (2001). It should also be noted that these analyses combine all NHL subtypes, which may have etiological differences (Morton *et al.*, 2014). Although some studies did provide effect estimates for subtypes, as stated in Section 3.5.2, these were not considered in the current evaluation due to the limited sample sizes. At this time, there are no data available to evaluate dose-response for NHL subtypes.

Furthermore, as discussed in Section 3.6, a dose-response relationship was not observed following the dramatic increase in glyphosate use due to the introduction of genetically engineered glyphosate-tolerant crops in 1996. Due to the change in use pattern, if a true association exists between glyphosate exposure and NHL, this large increase in use would be expected to result in a corresponding increase in risk of NHL associated with glyphosate exposure; therefore, higher effect estimates would be expected in more recent years. This trend was not observed though. For example, some of the highest adjusted risk measures for NHL

were reported for study years prior to 1996. Furthermore, it would also be expected that higher effect estimates would be reported in countries with higher use of glyphosate and/or that use glyphosate-tolerant crops, such as the United States and Canada, as compared to countries that exhibit less use. Once again, this trend was not observed with NHL studies, such that effect estimates for studies conducted in Sweden (Eriksson *et al.*, 2008; Hardell *et al.*, 2002) were similar or higher than those reported in the United States (De Roos *et al.*, 2003; De Roos *et al.*, 2005) and Canada (McDuffie *et al.*, 2001).

With respect to animal carcinogenicity studies, key events in a MOA/AOP are evaluated to confirm that they precede tumor appearance. This temporal concordance evaluation cannot be conducted for glyphosate since a MOA/AOP has not been established. In general, the tumor incidences lacked a monotonic dose-response. It should be noted, however, that no preneoplastic or related non-neoplastic lesions were reported in any of the animal carcinogenicity studies to support any observed tumors. Furthermore, genotoxicity assays did not support a direct mutagenic MOA. While there is limited evidence of genotoxic in some *in vitro* endpoints, multiple *in vivo* do not support a genotoxic risk at relevant human exposure levels.

6.3 Strength, Consistency, and Specificity

A large database is available for evaluating the carcinogenicity potential of glyphosate. Across animal carcinogenicity and genotoxicity studies, results were consistent. For epidemiological studies, only one or two studies were available for almost all cancers investigated. The largest number of studies was available investigating NHL; however, there were conflicting results across studies.

In epidemiological studies, there was no evidence of an association between glyphosate exposure and solid tumors, leukemia, and HL. This conclusion is consistent with those recently conducted by IARC, EFSA, and JMPR. The available data for multiple myeloma are not considered adequate to assess carcinogenic potential at this time.

At this time, a conclusion regarding the association between glyphosate exposure and risk of NHL cannot be supported based on the available data due to conflicting results. Chance and/or bias cannot be excluded as an explanation for observed associations. The magnitude of adjusted risk estimates for never/ever use were relatively small ranging from 1.0 (no association) to 1.85 in adjusted analyses, with the widest confidence interval observed for the highest effect estimate indicating the estimate is less reliable. All of the estimates were not statistically significant with half of the effect estimates approximately equal to 1, while the other half of the effect estimates ranged from 1.5-1.85. As a result, studies of at least equal quality provided conflicting results. There were various limitations identified in Section 3.6 for these studies that could impact calculated effect estimates and explain the weak responses observed in these studies. Meta-risk ratios using these studies were also of small magnitude ranging from 1.3-1.5. As discussed in Section 3.6, meta-analyses should be interpreted with caution and are susceptible to the same limitations identified for individual studies.

Although none of the effect estimates were below 1 using the never/ever exposure metric, risk estimates were all below 1 (0.6-0.9) when using cumulative lifetime and intensity-weighted

cumulative exposure metrics in the prospective cohort study (De Roos et al., 2005). As discussed in Section 6.2, two case-control studies that investigated an exposure-response relationship conflicted with the extensive analyses conducted by De Roos et al. (2005). This may be due to differences in confounding control, differences associated with study design, limitations from small sample sizes, and/or, as some may suggest, a potentially short follow-up time in the cohort. It should also be noted that publication bias may play a role in this evaluation given there is a tendency to only publish positive results and potential concerns regarding glyphosate have only been raised in recent years.

A total of 15 (9 rat and 6 mouse) animal carcinogenicity studies with glyphosate, glyphosate acid, or glyphosate salts were analyzed for the current evaluation. Although increases in tumor incidences were observed in some studies, none were considered treatment-related based on weight-of-evidence evaluations. In 7 of these studies, no tumors were identified for detailed evaluation. In the remaining studies, tumor incidences were not increased at doses <500 mg/kg/day, except for the testicular tumors observed in one study. The high dose tumors, as well as the testicular tumors, were not reproduced in other studies, including those testing the same animal strain with similar or higher dosing. Additionally, the tumors typically lacked a monotonic dose response, pairwise significance, and/or corroborating preneoplastic lesions.

Over 80 genotoxicity studies with the active ingredient glyphosate were analyzed for the current evaluation. The overall weight-of-evidence indicates that there is no convincing evidence that glyphosate is genotoxic *in vivo* via the oral route. When administered via i.p. injection the studies were predominantly negative. In the two cases where an increase in micronuclei were reported via this route, the effects were not observed in other i.p. injection studies at similar or higher doses. Technical glyphosate was negative in all gene mutation studies. There was limited evidence of positive findings in studies evaluating primary DNA damage; however, as discussed in Section 5.6, the endpoints measured in these assays are less specific in regards to detecting permanent DNA changes (mutations) and can be attributed to other factors, such as cytotoxicity or cell culture conditions. Although some positive findings were reported for chromosomal alterations *in vitro*, these findings were limited to a few studies and are not supported by the *in vivo* studies that are the most relevant for human risk assessment.

Overall, there is remarkable consistency in the database for glyphosate across multiple lines of evidence. For NHL, observed associations in epidemiological studies were non-statistically significant and were of relatively small magnitude. Chance and/or bias cannot be excluded as an explanation for the observed associations. For all other cancer types, there were no associations found; however, only one or two studies were available for evaluation of most cancer types. Across species, strain, and laboratory, tumor incidence was not increased at doses <500 mg/kg/day, except the testicular tumors which were only seen in one study. Observed tumors were not reproduced in other studies, including those conducted using the same strain at similar or higher doses. The genotoxicity studies demonstrate that glyphosate is not directly mutagenic or genotoxic *in vivo*.

6.4 Biological Plausibility and Coherence

The Guidelines for Carcinogen Risk Assessment (U.S. EPA, 2005) include the following guidance regarding the criteria of biological plausibility and coherence:

“evaluation of the biological plausibility of the associations observed in epidemiologic studies reflects consideration of both exposure-related factors and toxicological evidence relevant to identification of potential modes of action (MOAs). Similarly, consideration of the coherence of health effects associations reported in the epidemiologic literature reflects broad consideration of information pertaining to the nature of the biological markers evaluated in toxicologic and epidemiologic studies. [p.39].”

The genotoxicity studies demonstrate that glyphosate is not directly mutagenic or genotoxic *in vivo*. The available data regarding non-cancer endpoints also do not provide any support for a carcinogenic process for glyphosate, and have shown glyphosate has relatively low toxicity. Laboratory animals generally display non-specific effects (e.g., clinical signs, reduced body weight) following glyphosate exposure at relatively high-doses, and no preneoplastic or related non-neoplastic lesions were observed to corroborate any of the observed tumors in the carcinogenicity studies. As discussed in Section 4.2, metabolism studies demonstrate low oral absorption and rapid excretion of glyphosate. The data are not sufficient to determine whether linear kinetics is occurring at high doses where tumors were observed. In the carcinogenicity test guideline (OCSPP 870.4200) and the 2005 Guidelines for Carcinogen Risk Assessment, inappropriate toxicokinetics (e.g., overwhelming absorption or detoxification mechanisms) should be avoided. A study evaluating the toxicokinetic profile of glyphosate using multiple doses is needed to further investigate the pharmacokinetic properties between low- and high-dose levels.

Although many of the studies included in this document focus on the potential for glyphosate to cause a cancer outcome, the agency is also aware of a limited number of studies in the open literature that have shown glyphosate and its metabolite, AMPA, can inhibit proliferation and promote apoptosis in cancer cells indicating the compounds have potential to be developed into therapeutic drugs for cancer treatment (Li *et al*, 2013; Parajuli *et al.*, 2015; Parajuli *et al.*, 2016). It is unknown if this is due to lack of additional studies that have investigated these compounds for cancer treatment or if this may be due to publication bias. The bias towards only publishing positive and/or novel results can hamper the ability to evaluate whether there are plausible biological mechanisms for observed outcomes and/or sufficient information to support a cause-and-effect interpretation of an association. Overall, this further supports the need for mechanistic data to elucidate the true mammalian MOA/AOP for glyphosate. There is a distinct lack of mechanistic understanding for the toxicity of glyphosate in mammals and the plant MOA is not relevant for mammalian systems.

As noted previously, tumor incidence in animal carcinogenicity studies was typically only increased at the highest doses tested (≥ 1000 mg/kg/day). It is very unlikely for people to be exposed to such large doses of glyphosate via the oral route. Glyphosate is registered for pre- and post-emergence application to a variety of fruit, vegetable, and field crops, as well as desiccant applications to several commodities. The highest dietary exposure value for any population subgroup in an unrefined chronic dietary analysis would be 0.23 mg/kg/day for children (1-2 years old). Since glyphosate also has residential uses, including application to turf,

there is also the potential for children at this age to be exposed via incidental oral exposures (e.g., hand to mouth, object to mouth and soil ingestion) from playing on treated lawns. The highest exposure for the incidental oral and dermal exposures would be 0.16 mg/kg/day (from hand-to-mouth behaviors for children) and 0.08 mg/kg/day, respectively. Combining exposures from the dietary and residential exposures for children would, therefore, result in an aggregate exposure of 0.47 mg/kg/day. These calculations use a number of assumptions that have been extensively peer-reviewed²⁷ and yet the potential oral exposure of glyphosate for the most highly exposed residential population subgroup is more than 2,000 times lower than the highest doses tested in the animal carcinogenicity studies (see Appendix E for more details regarding these calculations). The maximum potential exposure calculated for occupational handlers would be 7 mg/kg/day, which is still significantly lower than the highest doses tested in the animal carcinogenicity studies. As a result, even though tumors were observed in animal carcinogenicity studies, the possibility of being exposed to these excessive dietary doses over time is considered implausible based on the currently registered use pattern and not relevant to human health risk assessment.

6.5 Uncertainty

When evaluating a database, it is also important to assess the uncertainties associated with the available data. When uncertainty is high there is less confidence in the exposure and effect estimates and, therefore, informs the reliability of results. Understanding the sources of uncertainty within a database can help characterize observed results and aid in developing new research with reduced uncertainty.

In some instances, the agency did not have access to all of the data underlying the studies analyzed for the current evaluation. This includes all of the epidemiological studies, 17 genotoxicity studies, and 1 animal carcinogenicity study. For these studies, the agency had to rely upon information the study authors reported. Without the raw data, statistical analyses could not be replicated or recalculated. On the other hand, studies that include full reports with detailed methodology, analytically measured doses, and individual animal data may provide a higher level of confidence. It also allows the agency to perform its own evaluation of the data using current practices and policies.

Several uncertainties have already been identified throughout this document. There are numerous metabolism studies available for glyphosate; however, the data are not sufficient to determine whether linear kinetics is occurring at high doses where tumors were observed in animal carcinogenicity studies. In the carcinogenicity test guideline (OCSPP 870.4200) and the 2005 Guidelines for Carcinogen Risk Assessment, inappropriate toxicokinetics (e.g., overwhelming absorption or detoxification mechanisms) should be avoided. A study evaluating the toxicokinetic profile of glyphosate using multiple doses is needed to further investigate the pharmacokinetic properties between low- and high-dose levels.

²⁷ Using the 2012 Standard Operating Procedures for Residential Exposure Assessment. Available: <http://www2.epa.gov/pesticide-science-and-assessing-pesticide-risks/standard-operating-procedures-residential-pesticide>

With respect to the epidemiological data, the database is limited for each investigated cancer with only one or two studies available. Although six studies were used in the evaluation of NHL, the results were constrained by the limitations of the individual studies, such as small sample size/limited power, missing data, and control selection issues. The quality of the exposure assessment is a major concern since the validity of the overall study results depend in large part on the ability of the study to correctly quantify and classify a subject's exposure. There was no direct information on pesticide exposure or absorbed dose because the exposures were self-reported. All of the studies conducted exposure assessments through questionnaires and interviews that are susceptible to recall bias, which can result in exposure misclassification. The study with the highest ranking (De Roos *et al.*, 2005) did not find an association between glyphosate exposure and NHL; however, it has been noted that the median follow-up time for this study was ~7 years. A longer follow-up from the AHS cohort would be beneficial to better understand whether there is an association between glyphosate exposure and NHL. An update from the AHS cohort would also provide a more recent evaluation of glyphosate exposure and cancer outcomes. Many of the studies were conducted prior to the introduction of glyphosate-tolerant crops in 1996, which resulted in a dramatic increase of glyphosate use in subsequent use. More recent studies will help further elucidate the association between glyphosate exposure and cancer outcomes during this period of time.

Another consideration is that farmers and other applicators apply formulations, not the active ingredient alone. It is possible that different formulations were used across and/or within the different epidemiological studies. Formulations are end-use products that are sold as a mixture of registered pesticidal active ingredients, such as glyphosate, and additional substances that increase the effectiveness of a pesticidal product, which are often referred to as "inert ingredients." For example, inert ingredients may act as a solvent to allow a pesticide active ingredient to penetrate a plant's outer surface, may facilitate and accentuate the dispersion of the product, or may extend the pesticide product's shelf-life²⁸. Inert ingredients and the proportion of these inert ingredients vary across formulations. It has been hypothesized that glyphosate formulations may be more toxic than glyphosate alone. Glyphosate has been studied in a multitude of studies and there are studies that have been conducted on numerous formulations that contain glyphosate; however, there are relatively few research projects that have attempted to systematically compare glyphosate and the formulations in the same experimental design. Furthermore, there are even less instances of studies comparing toxicity across formulations. This is one aspect of the uncertainty in the database that the agency has been working to address by developing a strategic research plan in collaboration with NTP (see Section 7.0).

It is recognized that these uncertainties exist for the current database; however, the available data are adequate for evaluating the carcinogenic potential of glyphosate and determine the cancer classification using the 2005 Guidelines for Carcinogen Risk Assessment. As discussed in Section 6.3, there are a large number of studies available and the database is remarkably consistent across these studies.

²⁸ <https://www.epa.gov/pesticide-registration/inert-ingredients-overview-and-guidance>

6.6 Evaluation of Cancer Classification per the 2005 EPA Guidelines for Carcinogen Risk Assessment

6.6.1 Introduction

In the 2005 Guidelines for Carcinogen Risk Assessment, five classification descriptors are provided:

- Carcinogenic to Humans
- Likely to be Carcinogenic to Humans
- Suggestive Evidence of Carcinogenic Potential
- Inadequate Information to Assess Carcinogenic Potential
- Not Likely to be Carcinogenic to Humans

Descriptors are assigned using all available data from the multiple lines of evidence. The following text has been excerpted/summarized from the guidelines regarding these descriptors:

Choosing a descriptor is a matter of judgment and cannot be reduced to a formula. Each descriptor may be applicable to a wide variety of potential data sets and weights of evidence. The weight-of-evidence, including the selected descriptor, is presented as a narrative laying out the complexity of information that is essential to understanding the hazard and its dependence on the quality, quantity, and type(s) of data available. The descriptors and narratives are intended to permit sufficient flexibility to accommodate new scientific understanding and new testing methods. The descriptors represent points along a continuum of evidence; consequently, there are gradations and borderline cases that are clarified by the full weight-of-evidence narrative. Rather than focusing simply on the descriptor, the entire range of information included in the weight-of-evidence narrative should be considered.

The weight-of-evidence presented in Sections 6.2-6.5 and based on the available epidemiological, animal carcinogenicity, and genotoxicity data for glyphosate was considered for each classification descriptor. For each descriptor, the guidelines provide examples and/or conditions for when the descriptor may be appropriate and the weight-of-evidence for glyphosate is assessed to determine which descriptor is supported by the available data. As stated in the 2005 EPA Guidelines for Carcinogen Risk Assessment, “the entire range of information included in the weight-of-evidence should be considered”. Based on all of the available data, the weight-of-evidence clearly do not support the descriptors “carcinogenic to humans” and “likely to be carcinogenic to humans” at this time. According to the 2005 Cancer Guidelines, “carcinogenic to humans” is appropriate “when there is convincing epidemiologic evidence of a causal association between human exposure and cancer.” Similarly, “likely to be carcinogenic to humans” descriptor is appropriate “when the weight of the evidence is adequate to demonstrate carcinogenic potential to humans but does not reach the weight of evidence for the descriptor.”

In epidemiological studies, there was no evidence of an association between glyphosate exposure and solid tumors, leukemia, or HL. The available data for multiple myeloma are not considered adequate to assess carcinogenic potential and a conclusion regarding the association between

glyphosate exposure and risk of NHL cannot be determined based on the available data due to conflicting results and various limitations identified in studies investigating NHL. In 7 of the 15 animal carcinogenicity studies, no tumors were identified for detailed evaluation. In the remaining 8 studies, tumor incidences were not increased at doses <500 mg/kg/day, except for testicular tumors. The tumors observed at doses at or exceeding 1,000 mg/kg/day are not considered relevant to human health risk assessment. Tumor findings were not reproduced in studies in the same animal strain at similar or higher doses. Furthermore, the tumors often lacked a monotonic dose response, pairwise significance, and/or corroborating preneoplastic lesions. The mammalian MOA/AOP is unknown for glyphosate and precursor events are unknown; however, the genotoxicity data were highly reproducible and consistent with a clear demonstration that glyphosate does not have a mutagenic MOA.

The descriptor “inadequate information to assess carcinogenic potential” is used when available data are judged inadequate for applying one of the other descriptors. Given the extensive size of the glyphosate database, which includes a multitude of well-designed and well-conducted studies, this classification descriptor is not supported. The epidemiological data at this time are limited and study results appear to be inconsistent for some cancer types. However, it is important to note that epidemiological studies are not available for most pesticides. Similarly, for most pesticides, generally, only two animal bioassays are available. EPA routinely evaluates human cancer potential using the small, more typical datasets. As such, for glyphosate, given the significant amount of information across multiple lines of evidence, the agency believes the database is sufficient to designate a cancer classification descriptor for glyphosate and that “inadequate information to assess carcinogenic potential” is not appropriate.

The remaining two cancer classification descriptors (“*Suggestive Evidence of Carcinogenic Potential*” and “*Not Likely to Be Carcinogenic to Humans*”) from the 2005 EPA Guidelines for Carcinogen Risk Assessment are described in detail below. Subsequently, these descriptors are discussed in the context of whether the available evidence do or do not support them.

“Suggestive Evidence of Carcinogenic Potential”

This descriptor is appropriate when a concern for potential carcinogenic effects in humans is raised, but the data are judged not sufficient for a stronger conclusion. It covers a spectrum of evidence associated with varying levels of concern for carcinogenicity. Depending on the extent of the database, additional studies may or may not provide further insights.

Some examples of when this descriptor may be appropriate include the following:

- If a small, and possibly not statistically significant, increase in tumor incidence observed in a single animal or human study that does not reach the weight-of-evidence for the descriptor of “likely to be carcinogenic to humans.” The study generally would not be contradicted by other studies of equal quality in the same population group or experimental system;
- If there is evidence of a positive response in a study whose power, design, or conduct limits the ability to draw a confident conclusion (but does not make the study fatally flawed), but where the carcinogenic potential is strengthened by other lines of evidence;

- If there is a small increase in a tumor with a high background rate in that sex and strain, when there is some but insufficient evidence that the observed tumors may be due to intrinsic factors that cause background tumors and not due to the agent being assessed (when there is a high background rate of a specific tumor in animals of a particular sex and strain, then there may be biological factors operating independently of the agent being assessed that could be responsible for the development of the tumors). In this case, the reasons for determining that the tumors are not due to the agent are explained; or
- If there is a statistically significant increase at one dose only, but no significant response at the other doses and no overall trend.

“Not Likely to Be Carcinogenic to Humans”

This descriptor is appropriate when the available data are considered robust for deciding that there is no basis for human hazard concern. In some instances, there can be positive results in experimental animals when there is strong, consistent evidence that each MOA in experimental animals does not operate in humans. In other cases, there can be convincing evidence in both humans and animals that the agent is not carcinogenic.

This descriptor would be appropriate if any of the following was observed:

- Animal evidence demonstrates lack of carcinogenic effects in both sexes in well-designed and well-conducted studies in at least two appropriate animal species in the absence of other animal or human data suggesting a potential for cancer effects, or
- Convincing and extensive experimental evidence showing that the only carcinogenic effects observed in animals are not relevant to humans, or
- Convincing evidence that carcinogenic effects are not likely by a particular exposure route, or
- Convincing evidence that carcinogenic effects are not likely below a defined dose range.

6.6.2 Discussion of Evidence to Support Cancer Classification Descriptors

As stated above, the available data and weight-of-evidence clearly do not support the descriptors “carcinogenic to humans”, “likely to be carcinogenic to humans”, or “inadequate information to assess carcinogenic potential”. The following discusses the remaining cancer classification descriptors and how the evidence does or does not support the descriptors.

It could be argued that the “suggestive evidence of carcinogenic potential” descriptor would be appropriate. The evidence to support this includes:

- Non-statistically significant effect estimates greater than the null were reported for NHL across studies and meta-analyses based on ever/never use ranged from 1.3-1.5.
- There was limited evidence of a possible exposure-response relationship between glyphosate exposure and NHL.

- In several animal carcinogenicity studies, a statistically significant trend was observed. In some instances, tumor incidences at the highest dose tested were statistically significant as compared to concurrent controls using raw (unadjusted) p-values.
- Positive responses were observed in a limited number of genotoxicity assays evaluating chromosomal and primary DNA damage.

However, according to the 2005 EPA Guidelines for Carcinogen Risk Assessment, in order for the above evidence to support the “suggestive evidence of carcinogenic potential” descriptor, “the study generally would not be contradicted by other studies of equal quality in the same population group or experimental system”. Furthermore, the guidelines state that “rather than focusing simply on the descriptor, the entire range of information included in the weight-of-evidence narrative should be considered”. For the epidemiological studies evaluating NHL, half of the studies reported effect estimates for ever/never use ranging from 1.5-1.85, with the widest confidence interval observed for the highest effect estimate indicating the effect estimate is less reliable. In the other half of the studies, which were of equal or higher quality, the reported effect estimates were approximately equal to the null. All of the effect estimates were non-statistically significant. There were conflicting results in exposure-response assessments investigating glyphosate exposure and the risk of NHL. Although two-case control studies (McDuffie et al., 2001; Eriksson et al., 2008) reported elevated effect estimates when analyzing for exposure-response relationships across two exposure categories, extensive analyses in a study of equal or higher quality (De Roos et al., 2005) for cumulative lifetime exposure and intensity-weighted cumulative exposure contradicted these results reporting effect estimates less than null (ranging from 0.6-0.9) when analyzing across tertiles. Furthermore, the two-case control studies did not account for co-exposure to other pesticides, which would be expected to cause inflated effect estimates. Various limitations that could impact the calculated effect estimate were identified for these studies and discussed in Section 3.6. The effect estimates greater than the null were not strengthened by other lines of evidence, as described in Sections 6.2-6.5.

In 7 (5 rat and 2 mouse) of the 15 animal carcinogenicity studies conducted with glyphosate, no tumors were identified for detailed evaluation. Of the remaining 8 studies, 7 observed a statistically significant trend for a particular tumor type; however, the agency determined that these tumors findings are not considered to be related to treatment. Although a statistically significant trend was obtained, closer examination of the incidence data across doses did not demonstrate a monotonic dose responses in several instances. Although the incidence at the highest dose tested (approaching or exceeding 1,000 mg/kg/day for almost all studies) for some of these tumors were statistically significant from concurrent controls using raw (unadjusted) p-values, none of the pairwise comparisons were found to be statistically significant following adjustment for multiple comparisons, except the testicular tumors that were seen in a single study. Furthermore, these high-dose tumors were given less weight. There was no evidence of corroborating pre-neoplastic or related non-neoplastic lesions and tumors showed no evidence of tumor progression to support the biological significance of tumor findings. In a limited number of cases, the agency also considered historical control data to inform the relevance of tumor findings when incidence rates in the concurrent controls were unusually low. Lastly, tumors seen in individual studies were not reproduced in studies of equal quality, including studies in the same animal species and strain at similar or higher doses.

Although positive responses were observed in a limited number of genotoxicity assays evaluating chromosomal and primary DNA damage, the overall weight-of-evidence indicates that there is no convincing evidence that glyphosate induces mutations *in vivo* via the oral route. When administered via i.p. injection the studies were predominantly negative. In the two cases where an increase in micronuclei were reported via this route of administration, the results were contradicted by numerous other studies at similar or higher doses using the same assays and route of administration. Technical glyphosate was negative in all gene mutation studies. There was limited evidence of positive findings in studies evaluating primary DNA damage; however, the endpoints measured in these assays are less specific in regards to detecting permanent DNA changes (mutations) and can be attributed to other factors, such as cytotoxicity or cell culture conditions. Although some positive findings were reported for chromosomal alterations *in vitro*, these findings were limited to a few studies and are not supported by the *in vivo* studies that are the most relevant for human risk assessment.

In summary, considering the entire range of information for the weight-of-evidence, the evidence outlined above to potentially support the “suggestive evidence of carcinogenic potential” descriptor are contradicted by other studies of equal or higher quality and, therefore, the data do not support this cancer classification descriptor.

For the “not likely to be carcinogenic to humans” descriptor, one of the considerations is whether there is “convincing evidence that carcinogenic effects are not likely below a defined dose range”. In the case of glyphosate, effects are not likely below 500 mg/kg/day based on oral studies. Tumor incidences were not increased in animal carcinogenicity at doses <500 mg/kg/day, except for the testicular tumors observed in a single study that were not considered treatment-related. In genotoxicity studies, assays with oral administration were negative except for one instance where an extremely high dose (5,000 mg/kg/day) was administered.

The 2005 EPA Guidelines for Carcinogen Risk Assessment also state that “weighing of the evidence includes addressing not only the likelihood of human carcinogenic effects of the agent but also the conditions under which such effects may be expressed”. Increased tumor incidence was typically observed at doses of 1,000 mg/kg/day or greater. Additionally, the only *in vivo* positive assays seen in the genotoxicity studies were administered via i.p. injection at doses of 200 mg/kg/day and 300 mg/kg/day or orally at 5,000 mg/kg/day. These high doses are not considered relevant to human health risk assessment based on the currently registered use pattern for glyphosate. Maximum potential glyphosate exposure in residential and occupational settings have been estimated at 0.47 mg/kg/day and 7 mg/kg/day, respectively, which are well-below the doses necessary to elicit the effects seen in these animal carcinogenicity and genotoxicity studies. Additionally, non-linear kinetics may also be occurring at the high doses. The carcinogenicity test guidelines (OCSP 870.4200 and OCSP 870.4300) and the 2005 Guidelines for Carcinogen Risk Assessment state that inappropriate toxicokinetics (e.g., overwhelming absorption or detoxification mechanisms) should be avoided. A well-conducted pharmacokinetic study evaluating the toxicokinetic profile of glyphosate is needed to further investigate the toxicokinetic properties between high and low dose levels to ensure that inappropriate toxicokinetics is avoided.

Overall, there is not strong support for the “suggestive evidence of carcinogenic potential” cancer classification descriptor based on the weight-of-evidence, which includes the fact that even small, non-statistically significant changes observed in animal carcinogenicity and epidemiological studies were contradicted by studies of equal or higher quality. The strongest support is for “not likely to be carcinogenic to humans” at the doses relevant to human health risk assessment for glyphosate.

6.7 Proposed Conclusions Regarding the Carcinogenic Potential of Glyphosate

Glyphosate is a non-selective, phosphonomethyl amino acid herbicide registered to control weeds in various agricultural and non-agricultural settings. Labeled uses of glyphosate include over 100 terrestrial food crops as well as other non-agricultural sites, such as greenhouses, aquatic areas, and residential areas. Following the introduction of glyphosate-resistant crops in 1996, glyphosate use increased dramatically; however, glyphosate use has stabilized in recent years due to the increasing number of glyphosate-resistant weed species.

Since its registration in 1974, numerous human and environmental health analyses have been completed for glyphosate, which consider all anticipated exposure pathways. Glyphosate is currently undergoing Registration Review. As part of this process, the hazard and exposure of glyphosate are reevaluated to determine its potential risk to human and environmental health using current practices and policies. The human carcinogenic potential of glyphosate has been evaluated by the agency several times. As part of the current evaluation for Registration Review, the agency has performed a comprehensive analysis of available data from submitted guideline studies and the open literature. This includes epidemiological, animal carcinogenicity, and genotoxicity studies.

An extensive database exists for evaluating the carcinogenic potential of glyphosate, including 23 epidemiological studies, 15 animal carcinogenicity studies, and nearly 90 genotoxicity studies for the active ingredient glyphosate. These studies were evaluated for quality and results were analyzed across studies within each line of evidence. The modified Bradford Hill criteria were then used to evaluate multiple lines of evidence using such concepts as strength, consistency, dose response, temporal concordance and biological plausibility. The available data at this time do not support a carcinogenic process for glyphosate. Overall, animal carcinogenicity and genotoxicity studies were remarkably consistent and did not demonstrate a clear association between glyphosate exposure and outcomes of interest related to carcinogenic potential. In epidemiological studies, there was no evidence of an association between glyphosate exposure and numerous cancer outcomes; however, due to conflicting results and various limitations identified in studies investigating NHL, a conclusion regarding the association between glyphosate exposure and risk of NHL cannot be determined based on the available data. Increases in tumor incidence were not considered treatment-related in any of the animal carcinogenicity studies. In 7 of these studies, no tumors were identified for detailed evaluation. In the remaining studies, tumor incidences were not increased at doses <500 mg/kg/day, except for the testicular tumors observed in a single study. Increased tumor incidences at or exceeding the limit dose (≥ 1000 mg/kg/day) are not considered relevant to human health. Furthermore, data from epidemiological and animal carcinogenicity studies do not reliably demonstrate expected dose-response relationships.

For cancer descriptors, the available data and weight-of-evidence clearly do not support the descriptors “carcinogenic to humans”, “likely to be carcinogenic to humans”, or “inadequate information to assess carcinogenic potential”. For the “suggestive evidence of carcinogenic potential” descriptor, considerations could be looked at in isolation; however, following a thorough integrative weight-of-evidence evaluation of the available data, the database would not support this cancer descriptor. The strongest support is for “not likely to be carcinogenic to humans” at doses relevant to human health risk assessment.

This analysis integrating multiple lines of evidence highlights the need for mechanistic studies to elucidate the MOA/AOP of glyphosate, as well as additional epidemiology studies and updates from the AHS cohort study to further investigate the carcinogenic potential of glyphosate in humans. This evaluation focused on studies on the active ingredient glyphosate; however, additional research could also be performed to determine whether formulation components, such as surfactants, influence the toxicity of glyphosate formulations. The agency has been working on plans to initiate research given these identified data gaps and these plans are described in Section 7.0.

The agency is soliciting advice from the FIFRA SAP on the evaluation and interpretation of the available data for each line of evidence for the active ingredient glyphosate and the weight-of-evidence analysis, as well as how the available data inform cancer classification descriptors according to the agency’s 2005 Guidelines for Carcinogen Risk Assessment.

7.0 Collaborative Research Plan for Glyphosate and Glyphosate Formulations

As previously mentioned, some have believed that glyphosate formulations may be more toxic than glyphosate alone. Glyphosate has been studied in a multitude of studies and there are studies that have been conducted on numerous formulations that contain glyphosate; however, there are relatively few research projects that have attempted to directly compare glyphosate and the formulations in the same experimental design. Furthermore, there are even less instances of studies comparing toxicity across formulations.

The agency has been collaborating with the NTP Division of the National Institute of Environmental Health Sciences to develop a research plan intended to evaluate the role of glyphosate in product formulations and the differences in formulation toxicity. Four objectives were identified that laid out how research by NTP might contribute to these research questions: 1) compare the toxicity of glyphosate vs. formulations, as well as compare formulations vs. formulations, 2) provide publicly available toxicology data on cancer-related endpoints, 3) provide publicly available toxicology data on non-cancer endpoints, and 4) investigate the mechanisms of how glyphosate and formulations cause toxic effects.

As part of the first objective, NTP will investigate the differential biological activity of glyphosate, glyphosate formulations, and the individual components of formulations. . The NTP Laboratory Branch generated preliminary data by exposing human hepatoma cells (HepG2) to five different glyphosate products bought off the shelf. The endpoint in the assay was cell viability, measured by ATP levels. The data, presented in Figure 7.1, demonstrate at-a-glance

that formulations are not equally toxic and that the toxicity is not being driven by the amount of glyphosate in the formulations, at least for the endpoint of cell viability. This observation highlights how informative the data generated from this research can be to the overall research questions.

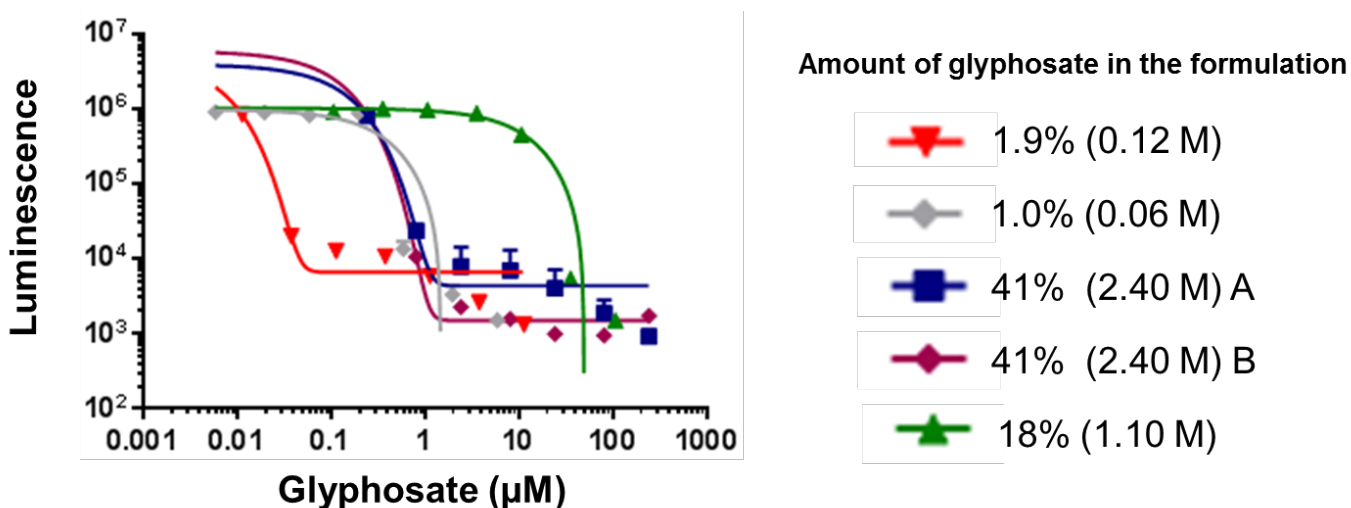


Figure 7.1. Results of HepG2 exposures following 24 hour incubation using different glyphosate formulations.

For the second objective, NTP will contribute to the publicly available knowledge-base describing the biological effects of glyphosate and formulations by conducting guideline studies addressing genotoxicity and studies that evaluate the oxidative stress potential. In order to organize publicly available data on glyphosate and formulations, IARC used 10 key characteristics of carcinogens as a way to help inform their conclusion (Smith *et al.*, 2016). Their review concluded that data were only available for two of these characteristics (genotoxicity and oxidative stress) and little to no information on the remaining characteristics was available. However, when members of a NTP workgroup looked at the available data included in the IARC review, the group did not agree with IARC that the data provided strong or clear evidence for either genotoxicity or induction of oxidative stress given protocol deficiencies that could produce questionable results.

Currently, the publicly available information regarding non-cancer endpoints for glyphosate and glyphosate formulations is limited. To begin to address the third objective, NTP will conduct a screening level analysis of the literature using text mining software, for studies regarding non-cancer endpoints resulting from glyphosate exposure. The resulting scoping report will describe the evidence base for health outcomes investigated in connection to glyphosate, as well as help identify data gaps.

As discussed in Section 6.0, there is a need for mechanistic studies to elucidate the MOA/AOP of glyphosate. Although there are data suggesting glyphosate may be genotoxic or cause oxidative stress, there is little mechanistic information to support these observations. For the last objective, NTP will use *in vitro* screening assays to gain mechanistic information on the effects

of glyphosate and different formulations for a variety of endpoints and allow for direct comparisons among them. The screening approach will also allow for the identification of test substances that would be good candidates for further *in vivo* testing. Since *in vivo* findings in genetic toxicology testing are generally considered as having a greater relevance to humans than *in vitro* findings, it is valuable to confirm the results seen at the cellular level at the whole animal level.

8.0 References

- Alvarez-Moya et al. (2014) "Comparison of the in vivo and in vitro genotoxicity of glyphosate isopropylamine salt in three different organisms". *Genetics and Molecular Biology*, 37, 1, 105-110
- Akanuma, M. (1995). HR-001: Reverse Mutation Test. Kodaira Laboratories. The Institute of Environmental Toxicology, Tokyo, Japan. Laboratory Project ID: IET 94-0142. April 3, 1995. MRID 50017102. Unpublished.
- Andreotti, G., Freeman, L.E., Hou, L., Coble, J., Rusiecki, J., Hoppin, J.A., Silverman, D.T., and Alavanja, M.C. (2009). Agricultural pesticide use and pancreatic cancer risk in the Agricultural Health Study Cohort. *International journal of cancer* 124, 2495-2500.
- Atkinson, C., Strutt, A., Henderson, W., *et al.* (1993a). 104-Week Chronic Feeding/Oncogenicity study in rats with 52-week interim kill. MRID No. 49631701. Unpublished
- Atkinson, C., Martin, T., Hudson, P., and Robb, D. (1993b). Glyphosate: 104 week dietary carcinogenicity study in mice. Inveresk Research International, Tranent, EH33 2NE, Scotland. IRI Project No. 438618. April 7, 1993. MRID 49631702. Unpublished.
- Band, P.R., Abanto, Z., Bert, J., Lang, B., Fang, R., Gallagher, R.P., and Le, N.D. (2011). Prostate cancer risk and exposure to pesticides in British Columbia farmers. *The Prostate* 71, 168-183.
- Baris, D, Garrity, TJ, Telles, JL, Heineman, EF, Olshan, A, Hoar Zahm, S. 2001. *American Journal of Industrial Medicine*. 39: 463-476.
- Benbrook. (2016). Trends in glyphosate herbicide use in the United States and globally. *Environmental Sciences Europe*. 28(3).
- Bolognesi, C., Bonatti, S., Degan, P., Gallerani, E., Peluso, M., Rabboni, R., Roggieri, P., and Abbondandolo, A. (1997). Genotoxic activity of glyphosate and its technical formulation roundup. *Journal of Agricultural and Food Chemistry* 45, 1957-1962.
- Brammer. (2001). Glyphosate Acid: Two Year Dietary Toxicity and Oncogenicity Study in Wistar Rats. Central Toxicology Laboratory, Alderley Park Macclesfield, Cheshire, UK: Syngenta. MRID 49704601. Unpublished.
- Brown, L.M., Blair, A., Gibson, R., Everett, G.D., Cantor, K.P., Schuman, L.M., Burmeister, L.F., Vanlier, S.F., and Dick, F. (1990). Pesticide Exposures and Other Agricultural Risk Factors for Leukemia among Men in Iowa and Minnesota. *Cancer Research* 50, 6585-6591.
- Brown, L.M., Burmeister, L.F., Everett, G.D., and Blair, A. (1993). Pesticide Exposures and Multiple Myeloma in Iowa Men. *Cancer Causes Control* 4, 153-156.

Brayton et al., 2012. Pathology of aging mice and GEM background strains and experimental design. *Vet Path.* 49 (1): 85-105.

Burnett, P., Borders, J.; Kush, J. (1979) Report to Monsanto Company: Two Year Chronic Oral Toxicity Study with CP- 76100 in Albino Rats: IBT No. 8560-08924. (Unpublished study received Jun 24, 1982 under 524-308; prepared by Industrial Bio-Test Laboratories, Inc., submitted by Monsanto Co., Washington, DC; CDL:247746-A; 247745; 247747; 247748; 247749; 247750; 247751; 247752)

Collander R.D. (1996). Glyphosate Acid: An Evaluation of Mutagenic Potential Using *S. typhimurium* and *E. coli*. Central Toxicology Laboratory, Cheshire, UK. Laboratory Project ID: CTL/P/4874 Study No. YV3611. February 16, 1996. MRID 44320617. Unpublished.

Cantor, K.P., Blair, A., Brown, L.M., Burmeister, L.F., and Everett, G. (1993). Pesticides and Other Agricultural Risk Factors for Non-Hodgkin's Lymphoma among Men in Iowa and Minnesota. *Cancer Research* 53, 2421-2421.

Carreón, T., Butler, M.A., Ruder, A.M., Waters, M.A., Davis-King, K.E., Calvert, G.M., Schulte, P.A., Connally, B., Ward, E.M., Sanderson, W.T., *et al.* (2005). Gliomas and Farm Pesticide Exposure in Women: The Upper Midwest Health Study. *Environmental Health Perspectives* 113, 546-551.

Cimino, M.C. (2006). Comparative overview of current international strategies and guidelines for genetic toxicology testing for regulatory purposes. *Environmental and Molecular Mutagenesis* 47 (9): 362-390.

Chang, E.T., and Delzell, E. (2016). Systematic review and meta-analysis of glyphosate exposure and risk of lymphohematopoietic cancers. *Journal of environmental science and health Part B, Pesticides, food contaminants, and agricultural wastes* 51, 402-434.

Chhabra et al. (1990). An over view of prechronic and chronic toxicity/carcinogenicity experimental study designs and criteria used by the National Toxicology Program. *Environ Health Perspect.* 86: 313-321.

Chruscielska et al. (2000). Glyphosate: evaluation of chronic activity and possible far-reaching effects. Part 1. Studies on chronic toxicity. *Pestycydy (Warsaw)*. 3-4: 11-20.

Collins, A.R., Oscoz, A.A., Brunborg, G., Gaivão, I., Giovannelli, L., Kruszewski, M., C.C., Stetina, R. (2008). The Comet assay: topical issues. *Mutagenesis* 23 (3): 143-151.

Cocco, P., Satta, G., Dubois, S., Pili, C., Pilleri, M., Zucca, M., t Mannetje, A.M., Becker, N., Benavente, Y., de Sanjose, S., *et al.* (2013). Lymphoma risk and occupational exposure to pesticides: results of the Epilymph study. *Occupational and environmental medicine* 70, 91-98.

Cooke et al., (2003). Oxidative DNA damage: mechanisms, mutation, and disease. *FASEB J.* 17 (10): 1195-214.

Chruscielska, K. et al. 2000. Glyphosate Evaluation of chronic activity and possible far-reaching effects. Part 2. Studies on mutagenic activity. *Pestycydy*, 2000, (3-4), 21-25. Published.

Dennis, L.K., Lynch, C.F., Sandler, D.P., and Alavanja, M.C. (2010). Pesticide use and cutaneous melanoma in pesticide applicators in the agricultural health study. *Environ Health Perspect* 118, 812-817.

De Roos, A.J., Zahm, S.H., Cantor, K.P., Weisenburger, D.D., Holmes, F.F., Burmeister, L.F., and Blair, A. (2003). Integrative assessment of multiple pesticides as risk factors for non-Hodgkin's lymphoma among men. *Occupational and environmental medicine* 60. 1-9.

De Roos, A. J., et al. (2005). "Cancer incidence among glyphosate-exposed pesticide applicators in the Agricultural Health Study." *Environ Health Perspect* 113(1): 49-54.

Durward, R. (2006). Technical Glyphosate: Micronucleus Test in the Mouse. Safepharm Laboratories Limited, Shardlow Business Park, Shardlow, Derbyshire DE72 2GD, UK, Study No. 2060/014. February 8, 2006. MRID 49957411. Unpublished.

Engel, L.S., Hill, D.A., Hoppin, J.A., Lubin, J.H., Lynch, C.F., Pierce, J., Samanic, C., Sandler, D.P., Blair, A., and Alavanja, M.C. (2005). Pesticide use and breast cancer risk among farmers' wives in the agricultural health study. *Am J Epidemiol* 161, 121-135.

Enemoto, K. (1997), HR-001: 24-Month Oral Chronic Toxicity and Oncogenicity Study in Rats, Vol. 1. The Institute of Environmental Toxicology, Kodaira-shi, Tokyo, Japan, Arysta Life Sciences, Study No.:IET 94-0150. MRID 50017104, 50017105, 5001703. Unpublished.

Eriksson, M., Hardell, L., Carlberg, M., and Akerman, M. (2008). Pesticide exposure as risk factor for non-Hodgkin lymphoma including histopathological subgroup analysis. *International journal of cancer* 123, 1657-1663.

Excel (1997). Combined chronic toxicity/carcinogenicity study of glyphosate technical in Sprague Dawley rats. Pune, India: Indian Institute of Toxicology.

Fisher, RA. (1950) Statistical methods for research workers. Edinburgh, Scotland: Oliver and Boyd.

Flower, K.B., Hoppin, J.A., Lynch, C.F., Blair, A., Knott, C., Shore, D.L., and Sandler, D.P. (2004). Cancer Risk and Parental Pesticide Application in Children of Agricultural Health Study Participants. *Environmental Health Perspectives* 112, 631-635.

Flowers, L.J., and Kier, L.D., Hannah, L.H. (1978) Final Report on Salmonella Mutagenicity Assay of Glyphosate: Test No. LF-78-161. MRID 00078620. Unpublished.

Fontana et al (1998). Incidence rates of lymphomas and environmental measurements of phenoxy herbicides: ecological analysis and case-control study. *Archives of Environmental Health: An International Journal*. 53: 384-387.

- Fox, V. (1998). Glyphosate acid: In vitro cytogenetic assay in human lymphocytes. Central Toxicology Laboratory, Cheshire, UK. Report CTL/P/6050. October 29, 1998. MRID 49961803. Unpublished
- Fox and Mackay (1996). Glyphosate acid: Mouse bone marrow micronucleus test. Central Toxicology Laboratory, Cheshire, UK. March 21, 1996. MRID 44320619. Unpublished.
- Green and Owen, (2011). Herbicide-resistant crops: utilities and limitations for herbicide-resistant weed management. *J. Agric Food Chem.* 59 (11): 5819-29.
- Greim, H., et al. (2015). "Evaluation of carcinogenic potential of the herbicide glyphosate, drawing on tumor incidence data from fourteen chronic/carcinogenicity rodent studies." *Crit Rev Toxicol* 45(3): 185-208.
- George, J., et al. (2010). "Studies on glyphosate-induced carcinogenicity in mouse skin: a proteomic approach." *J Proteomics* 73(5): 951-964.
- Ghisi, N.d.C., de Oliveira, E.C., and Prioli, A.J. (2016). Does exposure to glyphosate lead to an increase in the micronuclei frequency? A systematic and meta-analytic review. *Chemosphere* 145, 42-54.
- Giknis, M. L. A., and Clifford, C. B. (2005). Spontaneous Neoplastic Lesions in the Crl:CD1 (ICR) Mouse in Control Groups from 18 Month to 2 Year Studies. Charles River.
http://www.criver.com/files/pdfs/rms/cd1/rm_rm_r_lesions_crlcd_1_icr_mouse.aspx
- Hardell, L., Eriksson, M., and Nordstrom, M. (2002). Exposure to pesticides as risk factor for non-Hodgkin's lymphoma and hairy cell leukemia: Pooled analysis of two Swedish case-control studies. *Leukemia & Lymphoma* 43, 1043-1049.
- Hardell, L., and Eriksson, M. (1999). A case-control study of non-Hodgkin lymphoma and exposure to pesticides. *Cancer* 85, 1353-1360.
- Haseman, JK. (1995) Data analysis: Statistical analysis and use of historical control data. *Regul Toxicol Pharmacol* 21:52–59.
- Hill AB (1965). The Environment and Disease: Association or Causation? *Proc R Soc Med.* May 1965; 58(5): 295–300.
- Hoar *et al.* (1986). Agricultural herbicide use and risk of lymphoma and soft-tissue sarcoma. *JAMA.* 256:1141-1147.
- Hohenadel, K., Harris, S.A., McLaughlin, J.R., Spinelli, J.J., Pahwa, P., Dosman, J.A., Demers, P.A., and Blair, A. (2011). Exposure to multiple pesticides and risk of non-Hodgkin lymphoma in men from six Canadian provinces. *International journal of environmental research and public health* 8, 2320-2330.
- Hornarvar, N. (2008). Technical Glyphosate: Micronucleus Test of the Mouse. RCC, Cytotest

Cell Research GmbH (RCC-CCR0, In den Leppsteinswiesen 19, 64380 Rossdorf, Germany, Study No. 1158500. June 9, 2008. MRID 49961802. Unpublished.

Hsu and Stedeford (2010). *Cancer Risk Assessment: Chemical Carcinogenesis, Hazard Evaluation, and Risk Quantification*. John Wiley & Sons.

Jensen, J.C., 1991a. Mutagenicity test: Ames Salmonella Assay with Glyphosate, Batch 206-JaK-25-1. Scantox A/S, Lemvig, Denmark. Laboratory No. 12323. October 9, 1991. MRID 49961502. Unpublished.

Jensen, J.C. (1991b). Mutagenicity test: In vitro mammalian cell gene mutation test with glyphosate, batch 206-JaK-25-1. Scantox A/S, Lemvig, Denmark. Laboratory No. 12325. October 9, 1991. MRID 49961504. Unpublished.

Jensen, JC (1991c). Mutagenicity test: Micronucleus test with Glyphosate, batch 206-JAK-25-1. Scantox A/S, Lemvig, Denmark. Report number 12324. December 9, 1991. MRID 49961503.

Kachuri, L., Demers, P.A., Blair, A., Spinelli, J.J., Pahwa, M., McLaughlin, J.R., Pahwa, P., Dosman, J.A., and Harris, S.A. (2013). Multiple pesticide exposures and the risk of multiple myeloma in Canadian men. *International journal of cancer* 133, 1846-1858.

Karipidis et al. 2007. Occupational exposure to ionizing and non-ionizing radiation and risk of non-Hodgkin lymphoma. *Int Arch Occup Environ Health*. 80: 663-670.

Karunanayake, CP, McDuffie, HH, Dosman, JA, Spinelli, JJ, Pahwa, P. 2008. Occupational exposures and non-Hodgkin's lymphoma: Canadian case-control study. *Environmental Health*. 7:44.

Karunanayake, C.P., Spinelli, J.J., McLaughlin, J.R., Dosman, J.A., Pahwa, P., and McDuffie, H.H. (2012). Hodgkin lymphoma and pesticides exposure in men: a Canadian case-control study. *Journal of agromedicine* 17, 30-39.

Kaufman, D.W., Anderson, T.E., and Issaragrisil, S. (2009). Risk factors for leukemia in Thailand. *Annals of hematology* 88, 1079-1088.

Kato et al. (2005). Personal and occupational exposure to organic solvents and risk of non-Hodgkin's lymphoma (NHL) in women (United States). *Cancer Causes & Control*. 16:1215-1224.

Kier, L. D. (2015). "Review of genotoxicity biomonitoring studies of glyphosate-based formulations." *Crit Rev Toxicol* 45(3): 209-218.

Kier and Kirkland (2013). Review of genotoxicity studies of glyphosate and glyphosate-based formulations. *Critical Reviews in Toxicology*. 43: 283-315.

Knezevich, A.L and Hogan, G. K. (1983). A chronic feeding study of glyphosate in mice. Unpublished report prepared by Bio/Dynamic Inc., dated July 21, 1983. Report No. 77-2011.

EPA Accession No. 251007 – 251009, and 251014. EPA Accession no. 251007-09, 251014. Unpublished.

Koller, V.J., Furhacker, M., Nersesyan, A., Misik, M., Eisenbauer, M., and Knasmueller, S. (2012). Cytotoxic and DNA-damaging properties of glyphosate and Roundup in human-derived buccal epithelial cells. *Archives of toxicology* 86, 805-813.

Koureas, M., Tsezou, A., Tsakalof, A., Orfanidou, T., and Hadjichristodoulou, C. (2014). Increased levels of oxidative DNA damage in pesticide sprayers in Thessaly Region (Greece). Implications of pesticide exposure. *The Science of the total environment* 496, 358-364.

Koutros, S., Beane Freeman, L.E., Lubin, J.H., Heltshe, S.L., Andreotti, G., Barry, K.H., DellaValle, C.T., Hoppin, J.A., Sandler, D.P., Lynch, C.F., *et al.* (2013). Risk of total and aggressive prostate cancer and pesticide use in the Agricultural Health Study. *Am J Epidemiol* 177, 59-74.

Kumar, D.P.S. (2001), Carcinogenicity Study with Glyphosate Technical in Swiss Albino Mice, Toxicology Department Rallis Research Centre, Rallis India Limited. Study No. TOXI: 1559.CARCI-M. MRID 49987403. Unpublished.

Landgren, O., Kyle, R.A., Hoppin, J.A., Beane Freeman, L.E., Cerhan, J.R., Katzmann, J.A., Rajkumar, S.V., and Alavanja, M.C. (2009). Pesticide exposure and risk of monoclonal gammopathy of undetermined significance in the Agricultural Health Study. *Blood* 113, 6386-6391.

J. Langsdale *et al.* (2009). Glyphosate. Human Health Assessment Scoping Document in Support of Registration Review. June 3, 2009. D362745.

Lankas, G, P. (1981) A Lifetime Study of Glyphosate in Rats. Report No. 77-2062 prepared by Bio Dynamics, Inc. EPA Accession. No. 247617 – 247621. December 23, 1981. MRID 00093879. Unpublished.

Lee, W.J., Cantor, K.P., Berzofsky, J.A., Zahm, S.H., and Blair, A. (2004a). Non-Hodgkin's lymphoma among asthmatics exposed to pesticides. *International journal of cancer* 111, 298-302.

Lee, W.J., Lijinsky, W., Heineman, E.F., Markin, R.S., Weisenburger, D.D., and Ward, M.H. (2004b). Agricultural pesticide use and adenocarcinomas of the stomach and oesophagus. *Occupational and environmental medicine* 61, 743-749.

Lee, W.J., Colt, J.S., Heineman, E.F., McComb, R., Weisenburger, D.D., Lijinsky, W., and Ward, M.H. (2005). Agricultural pesticide use and risk of glioma in Nebraska, United States. *Occupational and environmental medicine* 62, 786-792.

Lee, W.J., Sandler, D.P., Blair, A., Samanic, C., Cross, A.J., and Alavanja, M.C. (2007). Pesticide use and colorectal cancer risk in the Agricultural Health Study. *International journal of cancer* 121, 339-346.

Li, A.P. (1983a). CHO/HGPRT gene mutation assay with glyphosate. Environmental Health Lab, St. Louis, MO. Study Number T830044. October 20, 1983. MRID 00132681. Unpublished.

Li, A.P. (1983b). In vivo bone marrow cytogenetic study of glyphosate in Sprague Dawley rats. Environmental Health Laboratory, St. Louis, MO. October 20, 1983. MRID 00132683. Unpublished.

Li, A. P. and T. J. Long (1988). An evaluation of the genotoxic potential of glyphosate. *Fundam Appl Toxicol* 10(3): 537-546.

Lioi M.B., et al. (1998a). Genotoxicity and oxidative stress induced by pesticide exposure in bovine lymphocyte cultures in vitro. *Mutation Research- Fundamental and Molecular Mechanisms of Mutagenesis*. 403 (1-2), 13-20.

Lioi M.B., et al. (1998b). Cytogenetic Damage and Induction of Pro-Oxidant State in Human Lymphocytes Exposed In Vitro to Glyphosate, Vinclozolin, Atrazine, and DPX-E9636. *Environmental and Molecular Mutagenesis* 32: 39-46.

Lioi, M.B., Scarfi, M.R., Santoro, A., Barbieri, R., Zeni, O., Salvemini, F., Di Berardino, D., and Ursini, M.V. (1998a). Cytogenetic damage and induction of pro-oxidant state in human lymphocytes exposed in vitro to glyphosate, vinclozolin, atrazine, and DPX-EP636. *Environmental and Molecular Mutagenesis* 32, 39-46.

Lioi, M.B., Scarfi, M.R., Santoro, A., Barbieri, R., Zeni, O., Di Berardino, D., and Ursini, M.V. (1998b). Genotoxicity and oxidative stress induced by pesticide exposure in bovine lymphocyte cultures in vitro. *Mutation Research-Fundamental and Molecular Mechanisms of Mutagenesis* 403, 13-20.

Lowit *et al.* (2015). Literature Review on Neurodevelopmental Effects & FQPA Safety Factor Determination for the Organophosphate Pesticides. September 15, 2015. D331251.

Maizlish, N, Beaumont, J, Singleton, J. 1998. Mortality among California highway workers. *American Journal of Industrial Medicine*. 13: 363-379.

Majeska, J.; Zameck, R.; Matheson, D. (1982a) SC-0224 (Lot No. 7269- 10): Mutagenicity Evaluation in *Salmonella typhimurium*: Report No. T-10847. MRID 00126612. Unpublished.

Majeska, J. (1982b) Morphological transformation of Balb/3T3 cells. Report No. T-10884. MRID 00126616. Unpublished.

Majeska, J. (1982c) Mutagenicity evaluation of bone marrow cytogenetic analysis in rats. Report No. T10884. MRID 00132176. Unpublished.

Majeska, J. (1985a) Mutagenicity Evaluation in *Salmonella typhimurium*: SC-0224: Report No. T-12660. MRID 00155527. Unpublished.

Majeska, J. (1985b) Mutagenicity Evaluation in Mouse Lymphoma Multiple Endpoint Test Forward Mutation Assay: SC-0224: Report No. T-12661. MRID 00155528. Unpublished.

Majeska, J. (1985c) Mutagenicity Evaluation in Chinese Hamster Ovary Cytogenetic Assay: SC-0224: Report No. T-12663. MRID 00155530. Unpublished.

Majeska, J (1987). Mutagenic evaluation in bone marrow micronucleus. Environmental Health Center, Farmington, CT. Study Report No. T12689/SC-0024. April 23, 1987. MRID 40214004. Unpublished.

Mañas, F., Peralta, L., Raviolo, J., Ovando, H.G., Weyers, A., Ugnia, L., Cid, M.G., Larripa, I., and Gorla, N. (2009). Genotoxicity of glyphosate assessed by the comet assay and cytogenetic tests. *Environmental toxicology and pharmacology* 28, 37-41.

Mañas, F., Peralta, L., Ugnia, L., Weyers, A., García Ovando, H., and Gorla, N. (2013). Oxidative stress and comet assay in tissues of mice administered glyphosate and ampa in drinking water for 14 days. *BAG Journal of basic and applied genetics* 24, 67-75.

Marques, M.F.C. (1999). A Micronucleus Study in Mice for Glifosate Tecnico Nufarm. Bioagri Laboratorios Ltda. Study No: RF-G12.79/99. December 27, 1999. MRID 49957412. Unpublished.

Matsumoto (1995). HR-001: In vitro cytogenetics test. The Institute of Environmental Toxicology, Tokyo, Japan. Laboratory Project ID: IET 94-0143. May 29, 1995. MRID 50017106. Unpublished.

McConnell, EE; Solleveld, HA; Swenberg, JA; et al. (1986) Guidelines for combining neoplasms for evaluation of rodent carcinogenesis studies. *J Natl Cancer Inst* 76:283–289.

McDuffie, H.H., Pahwa, P., McLaughlin, J.R., Spinelli, J.J., Fincham, S., Dosman, J.A., Robson, D., Skinnider, L.F., and Choi, N.W. (2001). Non-Hodgkin's lymphoma and specific pesticide exposures in men: Cross-Canada study of pesticides and health. *Cancer Epidemiol Biomarkers Prev* 10, 1155-1163.

McDuffie, HH, Pahwa, P, McLaughlin, JR, Fincham, S, Robson, D, Dosman, JA, Hu, J. 2002. Canadian male farm residents, pesticide safety handling practices, exposure to animals and non-Hodgkin's lymphoma (NHL). *American Journal of Industrial Medicine*. 42: 54-61.

Meek ME, Boobis A, Cote I, Dellarco V, Fotakis G, Munn S, Seed J, Vickers C. 2014. New developments in the evolution and application of the WHO/IPCS framework on mode of action/species concordance analysis. *J Appl Toxicol*. 2014 Jan;34(1):1-18.

Mink, P. J., et al. (2012). "Epidemiologic studies of glyphosate and cancer: a review." *Regul Toxicol Pharmacol* 63(3): 440-452.

Mladinic, M., Berend, S., Vrdoljak, A.L., Kopjar, N., Radic, B., and Zeljezic, D. (2009a). Evaluation of genome damage and its relation to oxidative stress induced by glyphosate in human lymphocytes in vitro. *Environmental and molecular mutagenesis* 50, 800-807.

Mladinic, M., Perkovic, P., and Zeljezic, D. (2009b). Characterization of chromatin instabilities induced by glyphosate, terbuthylazine and carbofuran using cytome FISH assay. *Toxicology letters* 189, 130-137.

Moriya, M. et al. 1983. Further mutagenicity studies on pesticides in bacterial reverse assay systems. *Mutation Research*, 116 (1983), 185-216.

Morton *et al.* (2014). Etiologic heterogeneity among non-Hodgkin lymphoma subtypes: the interlymp non-Hodgkin lymphoma subtypes project. *J Natl Cancer Inst Monogr* 48: 130-144.

Nordstrom, M., Hardell, L., Magnuson, A., Hagberg, H., and Rask-Andersen, A. (1998). Occupational exposures, animal exposure and smoking as risk factors for hairy cell leukaemia evaluated in a case-control study. *Br J Cancer* 77, 2048-2052.

NTP (1992). NTP technical report on the toxicity studies of Glyphosate (CAS No. 1071-83-6) Administered In Dosed Feed To F344/N Rats And B6C3F1 Mice. Toxic Rep Ser 16: 1-d3.

OECD (2015). Guidance Document on Revisions to OECD Genetic Toxicology Test Guidelines. August 31, 2015.

Olsson and Brandt (1988). Risk of non-Hodgkin's lymphoma among men occupationally exposed to organic solvents. *Scandinavian Journal of Work, Environment, & Health*. 14:246-251.

Orsi, L., Delabre, L., Monnereau, A., Delval, P., Berthou, C., Fenaux, P., Marit, G., Soubeyran, P., Huguet, F., Milpied, N., *et al.* (2009). Occupational exposure to pesticides and lymphoid neoplasms among men: results of a French case-control study. *Occupational and environmental medicine* 66, 291-298.

Pahwa, P., Karunanayake, C.P., Dosman, J.A., Spinelli, J.J., McLaughlin, J.R., and Cross-Canada, G. (2011). Soft-tissue sarcoma and pesticides exposure in men: results of a Canadian case-control study. *Journal of occupational and environmental medicine / American College of Occupational and Environmental Medicine* 53, 1279-1286.

Pahwa, P., Karunanayake, C.P., Dosman, J.A., Spinelli, J.J., McDuffie, H.H., and McLaughlin, J.R. (2012). Multiple myeloma and exposure to pesticides: a Canadian case-control study. *Journal of agromedicine* 17, 40-50.

Parajuli, K. R., et al. (2015). "Aminomethylphosphonic acid and methoxyacetic acid induce apoptosis in prostate cancer cells." *Int J Mol Sci* 16(5): 11750-11765.

Parajuli, K. R., et al. (2016). "Aminomethylphosphonic acid inhibits growth and metastasis of human prostate cancer in an orthotopic xenograft mouse model." *Oncotarget* 7(9): 10616-10626.

Pavkov, K.Ll, Turnier, J.C. (1987) Two-Year Chronic Toxicity and Oncogenicity Dietary Study with SC-0224 in Mice. Stauffer Chemical Company. MRID 40214006. Unpublished.

Pavkov, K.Ll, Wyand, S. (1987) Two-Year Chronic Toxicity and Oncogenecity Dietary Study with SC-0224 in Rats. Stauffer Chemical Company. MRID 40214006. Unpublished.

Peluso M. et al. (1998). 32P-Postlabeling detection of DNA adducts in mice treated with herbicide Roundup. *Environ and Mol Mutagen* 31:55-59.

Piesova, E. (2004). The influence of different treatment length on the induction of micronuclei in bovine lymphocytes after exposure to glyphosate. *Folia Veterinaria* 48, 130-134.

Piesova, E. (2005). The effect of glyphosate on the frequency of micronuclei in bovine lymphocytes in vitro. *Acta Veterinaria-Beograd* 55, 101-109.

Portier, C.J., Armstrong, B.K., Baguley, B.C., Baur, X., Belyaev, I., Belle, R., Belpoggi, F., Biggeri, A., Bosland, M.C., Bruzzi, P., *et al.* (2016). Differences in the carcinogenic evaluation of glyphosate between the International Agency for Research on Cancer (IARC) and the European Food Safety Authority (EFSA). *Journal of epidemiology and community health*.

Rank, J., et al. (1993). "Genotoxicity testing of the herbicide Roundup and its active ingredient glyphosate isopropylamine using the mouse bone marrow micronucleus test, Salmonella mutagenicity test, and Allium anaphase-telophase test." *Mutat Res* 300(1): 29-36.

Recore *et al.* (2014). Glyphosate: Tier II incident report. February 6, 2014. D417808.

Reyna, M.S. Gordon, D.E. (1973) 18-Month Carcinogenic Study with CP67573 in Swiss White Mice: IBT No. B569. (Unpublished study, including sponsor's validation report dated Feb 1, 1978, received Jun 21, 1978 under 524-308; prepared by Industrial Bio-Test Laboratories, Inc., submitted by Monsanto Co., Washington, D.C.; CDL:234136-G). MRID 00061113. Unpublished.

Reyna, M.S. Richter, W.R. Gordon, D.E. (1974) Two- Year Chronic Oral Toxicity Study with CP67573 in Albino Rats: IBT No. B564. MRID 00062507. Unpublished.

Ribeiro do Val, R. (2007). Bacterial reverse mutation test (Ames test) for Glifosato Technico Helm. TECAM Tecnologia Ambiental Ltda., Sao Paulo, Brasil. Study Number 3393/2007-2.0AM, Report Number RL3393/2007-2.0 AM-B, December 13, 2007. MRID 50000903. Unpublished.

Rodney, DE (1980). Dominant lethal study in mice. International Research and Development Corp. May 23, 1980. MRID 0004634

Roustan, A., et al. (2014). "Genotoxicity of mixtures of glyphosate and atrazine and their environmental transformation products before and after photoactivation." *Chemosphere* 108: 93-100.

Ruder, A.M., Waters, M.A., Butler, M.A., Carreon, T., Calvert, G.M., Davis-King, K.E., Schulte, P.A., Sanderson, W.T., Ward, E.M., Connally, L.B., *et al.* (2004). Gliomas and farm pesticide exposure in men: the upper midwest health study. *Archives of environmental health* 59, 650-657.

Schinasi, L., and Leon, M.E. (2014). Non-Hodgkin lymphoma and occupational exposure to agricultural pesticide chemical groups and active ingredients: a systematic review and meta-analysis. *International journal of environmental research and public health* 11, 4449-4527.

Shirasu, Y.; Moriya, M.; Ohta, T. (1978) Microbial Mutagenicity Testing on CP67573 (Glyphosate). Apr 25, 1979 under 524-308; prepared by Institute of Environmental Toxicology, Japan, submitted by Monsanto Co., Washington, D.C.; CDL:238233-A. MRID 00078619. Unpublished.

Sivikova, K. and J. Dianovsky (2006). "Cytogenetic effect of technical glyphosate on cultivated bovine peripheral lymphocytes." *Int J Hyg Environ Health* 209(1): 15-20.

Smith *et al.*, (2016). Key characteristics of carcinogens as a basis for organizing data on mechanisms of carcinogenesis. *Environmental Health Perspectives*. 124: 713.

Snedecor, GW; Cochran, WG. (1967) *Statistical methods*, 6th ed. Ames, Iowa: Iowa State University Press.

Sokolowski, A. (2007a). *Salmonella typhimurium* and *Escherichia coli* Reverse Mutation Assay with Glyphosate Technical (NUP-05068). RCC Cytotest Cell Research GmbH, Rossdorf, Germany. Study Number 1061401. March, 16, 2007. MRID 49957406. Unpublished.

Sokolowski, A. (2007b). *Salmonella typhimurium* and *Escherichia coli* Reverse Mutation Assay with Glyphosate Technical (NUP-05070). RCC Cytotest Cell Research GmbH, Rossdorf, Germany. Study Number 1061402. March, 16, 2007. MRID 49957407. Unpublished.

Sokolowski, A. (2007c). *Salmonella typhimurium* and *Escherichia coli* Reverse Mutation Assay with Glyphosate Technical (NUP-05067). RCC Cytotest Cell Research GmbH, Rossdorf, Germany. Study Number 1061403. March, 16, 2007. MRID 49957408. Unpublished.

Sokolowski, A. (2009b). Glyphosate technical- *Salmonella Typhimurium* and *Escherichia Coli* reverse mutation assay. Cytotest Cell Research GmbH (Harlan CCR), Rossdorf, German. Study and Report Number 1264500. December 18, 2009. MRID 49961801. Unpublished.

Sokolowski, A. (2010). *Salmonella typhimurium* and *Escherichia coli* Reverse Mutation Assay with Glyphosate TC spiked with glyphosine. RCC Cytotest Cell Research GmbH, Rossdorf, Germany. Study Number 1332300. April 7, 2010. MRID 500000902. Unpublished.

Sorahan, T. (2015). "Multiple Myeloma and Glyphosate Use: A Re-Analysis of US Agricultural Health Study (AHS) Data." *Int J Environ Res Public Health* 12(2): 1548-1559.

Stout, L. D. and Ruecker, P.A. (1990). Chronic Study of Glyphosate Administered in Feed to Albino Rats. MRID No. 41643801; Historical Controls. MRID 41728700. Unpublished.

Sugimoto, K. (1997), HR-001: 18-Month Oral Oncogenicity Study in Mice, Vol. 1 and 2. The Institute of Environmental Toxicology, 2-772, Suzuki-cho, Kodaira-shi, Tokyo, 187, Japan, Study No.:IET 94-0151. MRID 50017108, 50017109. Unpublished.

Suresh, T.P. (1993b). Mutagenicity-Micronucleus Test In Swiss Albino Mice. Rallis India Limited. Study No: TOXI: 889-MUT.MN. May 6, 1993. MRID 49987407. Unpublished.

Suresh, T.P. (1994). Genetic Toxicology- In vivo mammalian bone marrow cytogenetic test-Chromosomal analysis. Rallis Agrochemical Research Station, Bangalore, India. January 1, 1994. MRID 49987408. Unpublished.

Suresh, T.P. (1996) Combined Chronic Toxicity and Carcinogenicity Study with Glyphosate Technical in Wistar Rats. Toxicology Department Rallis Research Centre, Rallis India Limited, TOXI-1559, 002/1-GPT-CARCI-M. MRID 49987401. Unpublished.

Taddesse-Heath, L.; Chattopadhyay, S.K.; Dillehay, D.; L.; Lander, M.R.; Nagashfar, Z.; Morse III, H.C.; Hartley, J.W. (2000): Lymphomas and high-level expression of murine leukemia viruses in CFW mice Journal of Virology 74:6832-6837

Tarone, RE. (1982). The use of historical control information in testing for a trend in proportions. Biometrics 38:215–220.

Thompson, P.W. (1996). Technical Glyphosate: Reverse Mutation Assay “Ames Test”Using *Salmonella typhimurium* and *Escherichia coli*. Safepharm Laboratories Limited, Derby, UK. Study Number 434/014. February 20, 1996. MRID 49957409. Unpublished.

Wang et al. (2009). Occupational exposure to solvents and risk of non-Hodgkin lymphoma in Connecticut women. American Journal of Epidemiology. 169:176-185.

Ward, J. M. (2006). Lymphomas and leukemias in mice. Experimental and Toxicologic Pathology, 57 (5-6): 377-381.

Weisenburger, D.D. (1992). Pathological Classification of Non-Hodgkin's Lymphoma for Epidemiological Studies. Cancer Research 52, 5456S-5462S.

Wilderman, A.G. and Nazar, R.N. (1982). Significance of plant metabolism in the mutagenicity and toxicity of pesticides. Canadian Journal of Genetics and Cytology 24(4): 437-449.

Williams, G. M., et al. (2000). "Safety Evaluation and Risk Assessment of the Herbicide Roundup and Its Active Ingredient, Glyphosate, for Humans." Regulatory Toxicology and Pharmacology 31(2): 117-165.

Wood, E., Dunster, J., Watson, P., and Brooks, P. (2009a) Glyphosate Technical: Dietary Combined Chronic Toxicity/Carcinogenicity Study in the Rat. Harlan Laboratories Limited,

Shardlow Business Park, Shardlow, Derbyshire DE72 2GD, UK. Study No. 2060-012. April, 23, 2009. MRID 49957404. Unpublished.

Wood, E., Dunster, J., Watson, P., and Brooks, P. (2009b) Glyphosate Technical: Dietary Carcinogenicity Study in the Mouse. Harlan Laboratories Limited, Shardlow Business Park, Shardlow, Derbyshire DE72 2GD, UK. Study No. 2060-011. April, 22, 2009. MRID 49957402. Unpublished.

Wright, N.P. (1996). Technical Glyphosate: Chromosomal aberration test in CHL cells in vitro. Safepharm Laboratories Limited, Derby, UK. Study Number 434/015. March 13, 1996, MRID 49957410. Unpublished.

Yauk *et al.*, (2015). Approaches to identifying germ cell mutagens: Report of the 2013 IWGT workshop on germ cell assays. *Mutat Res Genet Toxicol Environ Mutagen*, 783: 36-54.

Yiin, J.H., Ruder, A.M., Stewart, P.A., Waters, M.A., Carreon, T., Butler, M.A., Calvert, G.M., Davis-King, K.E., Schulte, P.A., Mandel, J.S., *et al.* (2012). The upper midwest health study: a case-control study of pesticide applicators and risk of glioma. *Environ Health* 11, 13.

Zaccaria, C.B. (1996). A micronucleus study in mice for the product Glyphosate. BioAgri. Study No: G.1.2 – 06/96. November 18, 1996. MRID 49961501. Unpublished.

Zahm *et al* (1990). A case-control study of non-Hodgkin's lymphoma and the herbicide 2,4-dichlorophenoxyacetic acid (2,4-D) in Eastern Nebraska. *Epidemiology*. 1:349-356.

Zhang *et al.* (2007). Ultraviolet radiation exposure and risk of non-Hodgkin's lymphoma. *American Journal of Epidemiology*. 165: 1255-1264.

Zoriki-Hosmi, (2007). Hosomi, R. (2007) Mammalian Erythrocyte Micronucleus Test for Glifosato Tecnico Helm. Project Number: RL/3393/2007/3/0MN/B. MRID 50000901. Unpublished.

Appendix A. Journal articles obtained from open literature search

Abstract Only	Cebollero, L. R., et al. (2011). "Glyphosate based herbicides toxicity, a new approach." <i>Toxicology Letters</i> 205, Supplement: S233.
Abstract Only	Monroy, C. M., et al. (2004). "In vitro evaluation of glyphosate-induced DNA damage in fibrosarcoma cells HT1080 and Chinese hamster ovary (CHO) cells." <i>Environ Mol Mutagen</i> 44(3): 216-216.
Abstract Only	Ramos-Morales, P., et al. (2008). "Combined use of multiple biomarkers to evaluate the genotoxic activity of the herbicide Glyphosate." <i>Environ Mol Mutagen</i> 49(7): 577-577.
Abstract Only/Full article already identified	Sorahan, T. (2015). "Multiple Myeloma and Glyphosate Use: A Re-Analysis of US Agricultural Health Study (AHS) Data." <i>Int J Environ Res Public Health</i> 12(2): 1548-1559.
Article not in English	Kwiatkowska, M., et al. (2013). "GLYPHOSATE AND ITS FORMULATIONS - TOXICITY, OCCUPATIONAL AND ENVIRONMENTAL EXPOSURE." <i>Med Pr</i> 64(5): 717-729.
Article not in English	Lawson, R. and E. Estrade-Chapellaz (1999). "Intoxication volontaire par le glufosinate (Basta®)." <i>Annales Françaises d'Anesthésie et de Réanimation</i> 18(9): 1025-1026.
Article not in English	Mañas, F., et al. (2009). "Aberraciones cromosómicas en trabajadores rurales de la Provincia de Córdoba expuestos a plaguicidas." <i>BAG. Journal of basic and applied genetics</i> 20(1): 0-0.
Article not in English	Martinez, A., et al. (2007). "[Cytotoxicity of the herbicide glyphosate in human peripheral blood mononuclear cells]." <i>Biomedica</i> 27(4): 594-604.
Article not in English	Monroy, C. M., et al. (2005). "[Cytotoxicity and genotoxicity of human cells exposed in vitro to glyphosate]." <i>Biomedica</i> 25(3): 335-345.
Article not in English	Pieniazek, D., et al. (2003). "[Glyphosate--a non-toxic pesticide?]." <i>Med Pr</i> 54(6): 579-583.
Article not in English	Saratovskikh, E. A., et al. (2007). "Genotoxicity of the pestiside in Ames test and the possibility to formate the complexeses with DNA." <i>Ekologicheskaya genetika</i> V(3): 46-54.
Article not in English	В статье представлены результаты генотоксико логических, аллергологических и иммунологических исследований, проведенных в рамках медико-биологической оценки безопасности генно-инженерно-моди-фицированной кукурузы линии MON 88017, устойчивой к глифосату и жуку <i>Diabrotica</i> spp. Анализ данных, полученных при изучении уровня повреждений ДНК и уровня хромосомных aberrаций, тяжести активного анафилактического шока и интенсивности гуморального иммунного ответа, состояния гуморального и клеточного звеньев иммунитета, не выявил какого-либо генотоксического, аллергенного, иммуномодулирующего и сенсибилизирующего действия ГМ-кукурузы линии MON 88017 по сравнению с ее традиционным аналогом.
Cancer treatment	Parajuli, K. R., et al. (2015). "Aminomethylphosphonic acid and methoxyacetic acid induce apoptosis in prostate cancer cells." <i>Int J Mol Sci</i> 16(5): 11750-11765.
Cancer treatment	Li, Q., et al. (2013). "Glyphosate and AMPA inhibit cancer cell growth through inhibiting intracellular glycine synthesis." <i>Drug Des Devel Ther</i> 7: 635-643.
Cancer treatment	Parajuli, K. R., et al. (2016). "Aminomethylphosphonic acid inhibits growth and metastasis of human prostate cancer in an orthotopic xenograft mouse model." <i>Oncotarget</i> 7(9): 10616-10626.
Correspondence article	Belle, R., et al. (2012). "LETTER TO THE EDITOR: TOXICITY OF ROUNDUP AND GLYPHOSATE." <i>Journal of Toxicology and Environmental Health-Part B-Critical Reviews</i> 15(4): 233-235.
Correspondence article	Carrasco, A. E. (2011). "Reply to the Letter to the Editor Regarding Our Article (Paganelli et al., 2010)." <i>Chem Res Toxicol</i> 24(5): 610-613.
Correspondence article	de Souza, L. and L. Macedo Oda (2013). "Letter to the editor." <i>Food and Chemical Toxicology</i> 53: 440.
Correspondence article	de Vendomois, J. S., et al. (2010). "Debate on GMOs Health Risks after Statistical Findings in Regulatory Tests." <i>Int J Biol Sci</i> 6(6): 590-598.
Correspondence article	Farmer, D. R., et al. (2005). "Glyphosate results revisited." <i>Environ Health Perspect</i> 113(6): A365-A366.
Correspondence article	Grunewald, W. and J. Bury (2013). "Comment on "Long term toxicity of a Roundup herbicide and a Roundup-tolerant genetically modified maize" by Seralini et al." <i>Food and Chemical Toxicology</i> 53: 447-448.
Correspondence article	Huang, K. and W. Xu (2013). "Reply to letter to the editor." <i>Food and Chemical Toxicology</i> 59: 811-812.
Correspondence article	Korsaeth, A., et al. (2015). "Comments on the recently published study: "Compositional differences in soybeans on the market: Glyphosate accumulates in Roundup Ready GM soybeans", by T. Bøhn, M.

	Cuhra, T. Traavik, M. Sanden, J. Fagan and R. Primicerio (Food Chemistry 2014, 153: 207–215)." Food Chemistry 172: 921-923.
Correspondence article	Ollivier, L. (2013). "A Comment on "Séralini, G.-E., et al., Long term toxicity of a Roundup herbicide and a Roundup-tolerant genetically modified maize. Food Chem. Toxicol. (2012)," http://dx.doi.org/10.1016/j.fct.2012.08.005 ." Food and Chemical Toxicology 53: 458.
Correspondence article	Portier, C. J., et al. (2016). "Differences in the carcinogenic evaluation of glyphosate between the International Agency for Research on Cancer (IARC) and the European Food Safety Authority (EFSA)." J Epidemiol Community Health.
Correspondence article	Roberfroid, M. (2014). "Letter to the editor." Food and Chemical Toxicology 66: 385.
Correspondence article	Sanders, D., et al. (2013). "Comment on "Long term toxicity of a Roundup herbicide and a Roundup-tolerant genetically modified maize" by Seralini et al." Food and Chemical Toxicology 53: 450-453.
Correspondence article	Schorsch, F. (2013). "Serious inadequacies regarding the pathology data presented in the paper by Séralini et al. (2012)." Food and Chemical Toxicology 53: 465-466.
Correspondence article	Wallace Hayes, A. (2014). "Editor in Chief of Food and Chemical Toxicology answers questions on retraction." Food and Chemical Toxicology 65: 394-395.
Effects on cellular processes	Benachour, N. and G.-E. Seralini (2009). "Glyphosate Formulations Induce Apoptosis and Necrosis in Human Umbilical, Embryonic, and Placental Cells." Chem Res Toxicol 22(1): 97-105.
Effects on cellular processes	Chaufan, G., et al. (2014). "Glyphosate commercial formulation causes cytotoxicity, oxidative effects, and apoptosis on human cells: differences with its active ingredient." Int J Toxicol 33(1): 29-38.
Effects on cellular processes	Coalova, I., et al. (2014). "Influence of the spray adjuvant on the toxicity effects of a glyphosate formulation." Toxicology in Vitro 28(7): 1306-1311.
Effects on cellular processes	George, J., et al. (2010). "Studies on glyphosate-induced carcinogenicity in mouse skin: a proteomic approach." J Proteomics 73(5): 951-964.
Effects on cellular processes	George, J. and Y. Shukla (2013). "Emptying of Intracellular Calcium Pool and Oxidative Stress Imbalance Are Associated with the Glyphosate-Induced Proliferation in Human Skin Keratinocytes HaCaT Cells." ISRN Dermatol 2013: 825180.
Effects on cellular processes	Heu, C., et al. (2012). "A step further toward glyphosate-induced epidermal cell death: Involvement of mitochondrial and oxidative mechanisms." Environmental Toxicology and Pharmacology 34(2): 144-153.
Not Relevant to current fit for purpose review	(1934). "The 1933 meeting of the American society of orthodontists at Oklahoma City." International Journal of Orthodontia and Dentistry for Children 20(1): 102-105.
Not Relevant to current fit for purpose review	(1938). "Supplementary report of region III: Reports of state chairmen." The Journal of Pediatrics 12(<HT>6</HT>): 846-850.
Not Relevant to current fit for purpose review	(1939). "Meeting of the Executive Board of the American Academy of Pediatrics." The Journal of Pediatrics 15(2): 294-315.
Not Relevant to current fit for purpose review	(1939). "Proceedings Meeting of the Executive Board of the American Academy of Pediatrics." The Journal of Pediatrics 14(2): 251-254.
Not Relevant to current fit for purpose review	(1967). "AORN Proceedings." AORN Journal 5(1): 82-91.
Not Relevant to current fit for purpose review	(1969). "AORN Proceedings." AORN Journal 9(5): 89-96.
Not Relevant to current fit for purpose review	(1969). "Nurse Recruitment Program—Stage I Winners." AORN Journal 10(<HT>6</HT>): 71-76.
Not Relevant to current fit for purpose review	(1970). "2036. Excretion of heliotrine in urine and bile: Jago, Marjorie, V., Lanigan, G. W., Bingley, J. B., Piercy, D. W. T., Whittem, J. H.& Titchen, D. A. (1969). Excretion of the pyrrolizidine alkaloid heliotrine in the urine and bile of sheep. J. Path. 98, 115." Food and Cosmetics Toxicology 8(5): 607-608.
Not Relevant to current fit for purpose review	(1970). "2037. A round-up of fungal toxins: Krogh, P. (1969). The pathology of mycotoxicoses. J. stored Prod. Res. 5, 259." Food and Cosmetics Toxicology 8(5): 608.
Not Relevant to current fit for purpose review	(1972). "News and comment." American Journal of Orthodontics 62(3): 319-334.
Not Relevant to current fit for purpose review	(1976). "NHI proposals: a wide spectrum." AORN Journal 23(5): 814-818.
Not Relevant to current fit for purpose review	(1977). "The SR periodicals miscellany." Serials Review 3(2): 11-19.

Not Relevant to current fit for purpose review	(1980). "Appendix F — administrative procedures, facilities and standardization of testing." Survey of Ophthalmology 24, Supplement: 594-597.
Not Relevant to current fit for purpose review	(1981). "Calendar." Annals of Emergency Medicine 10(3): 5-6.
Not Relevant to current fit for purpose review	(1981). "Reviews of Books." The Lancet 317(8232): 1239-1240.
Not Relevant to current fit for purpose review	(1982). "Reviews of Books." The Lancet 320(8307): 1076-1078.
Not Relevant to current fit for purpose review	(1983). "Product news." Journal of Molecular Graphics 1(3): 89-91.
Not Relevant to current fit for purpose review	(1984). "24–27 October 1984 International meeting on artificial intelligence: Marseilles, France." Computer Compacts 2(3–4): 121-122.
Not Relevant to current fit for purpose review	(1985). "Newsview." Journal of Allergy and Clinical Immunology 76(2, Part 1): A36-A41.
Not Relevant to current fit for purpose review	(1990). "Regional direct marketing clubs/associations support DM education at local colleges/universities." Journal of Direct Marketing 4(1): 46-52.
Not Relevant to current fit for purpose review	(1990). "Subject index." Toxicology 61(3): 313-316.
Not Relevant to current fit for purpose review	(1990). "Volume contents." Toxicology 61(3): 319-320.
Not Relevant to current fit for purpose review	(1991). "4971677 Fluorescence detection type electrophoresis apparatus: Hideki Kambara, Yoshiko Katayama, Tetsuo Nishikawa, Hachiouji, Japan assigned to Hitachi Ltd." Biotechnology Advances 9(1): 89.
Not Relevant to current fit for purpose review	(1991). "4971903 Pyrophosphate-based method and apparatus for sequencing nucleic acids: Edward Hyman." Biotechnology Advances 9(1): 89.
Not Relevant to current fit for purpose review	(1991). "4971908 Glyphosate-tolerant 5-enolpyruvyl 3-phosphoshikimate synthase: Ganesh M Kishore, Dilip Shah assigned to Monsanto Company." Biotechnology Advances 9(1): 89.
Not Relevant to current fit for purpose review	(1991). "Author Index." Journal of Chromatography A 540: 479-482.
Not Relevant to current fit for purpose review	(1991). "Challenge to US computer fraud & abuse act." Computer Fraud & Security Bulletin 1991(3): 5-6.
Not Relevant to current fit for purpose review	(1991). "Chemical index for volumes 16–17." Fundamental and Applied Toxicology 17(4): 848-850.
Not Relevant to current fit for purpose review	(1991). "Court case round-ups." Computer Fraud & Security Bulletin 1991(3): 5.
Not Relevant to current fit for purpose review	(1991). "FAT attacker!" Computer Fraud & Security Bulletin 1991(3): 5.
Not Relevant to current fit for purpose review	(1991). "FIGO news." International Journal of Gynecology & Obstetrics 35(3): 265-267.
Not Relevant to current fit for purpose review	(1991). "Local direct marketing clubs/associations support undergraduate/graduate level DM education." Journal of Direct Marketing 5(3): 61-65.
Not Relevant to current fit for purpose review	(1991). "Manipulation of molluscan haemocytes in vitro: *S.E. Fryer. Department of Zoology, Oregon State University, Corvallis OR 97331, USA." Developmental & Comparative Immunology 15, Supplement 1: S73.
Not Relevant to current fit for purpose review	(1991). "Preparation of monoclonal antibodies against hemolymph of the kuruma shrimp Penaeus japonicus (Crustacea: Decapoda): J. Rodriguez, V. Boulo, E. Bachère and E. Mialhe. IFREMER, Unité Pathol. Immunol. Génét. Mol., BP 133, 17390 La Tremblade, France." Developmental & Comparative Immunology 15, Supplement 1: S73.
Not Relevant to current fit for purpose review	(1991). "Round-up report from college/university direct marketing centers." Journal of Direct Marketing 5(2): 59-64.
Not Relevant to current fit for purpose review	(1992). "Forthcoming papers." Soil Biology and Biochemistry 24(1): 79.
Not Relevant to current fit for purpose review	(1992). "Subject index volume 85 (1992)." Plant Science 85(2): 235-237.

Not Relevant to current fit for purpose review	(1993). "Media watch." Physiotherapy 79(7): 496-498.
Not Relevant to current fit for purpose review	(1993). "Meetings and Notices." Journal of Equine Veterinary Science 13(9): 481-536.
Not Relevant to current fit for purpose review	(1993). "Meetings and notices." Journal of Equine Veterinary Science 13(8): 432-476.
Not Relevant to current fit for purpose review	(1993). "ROUND-UP Publications." Reproductive Health Matters 1(1): 109-110.
Not Relevant to current fit for purpose review	(1994). "Media Watch." Physiotherapy 80(11): 793-795.
Not Relevant to current fit for purpose review	(1994). "ROUND-UP Research." Reproductive Health Matters 2(4): 121-122.
Not Relevant to current fit for purpose review	(1994). "ROUND-UP Service Delivery." Reproductive Health Matters 2(3): 125-128.
Not Relevant to current fit for purpose review	(1995). "ROUND-UP Law and Policy." Reproductive Health Matters 3(<HT>6</HT>): 164-166.
Not Relevant to current fit for purpose review	(1995). "ROUND-UP Publications." Reproductive Health Matters 3(5): 146-151.
Not Relevant to current fit for purpose review	(1995). "ROUND-UP Service Delivery." Reproductive Health Matters 3(<HT>6</HT>): 167-169.
Not Relevant to current fit for purpose review	(1996). "5410871 Emission control device and method: Masters Ben F; Self James M Gastonia, NC, United States Assigned to Unlimited Technologies Inc." Environment International 22(2): XVI-XVII.
Not Relevant to current fit for purpose review	(1996). "5411697 Method for processing contaminated plastic waste: McGraw Peter S; Drake John; Hane Thomas H Severna Park, MD, United States Assigned to The United States of America as represented by the Secretary of the Navy." Environment International 22(2): XVII.
Not Relevant to current fit for purpose review	(1996). "5411944 Glyphosate-sulfuric acid adduct herbicides and use: Young Donald C Fullerton, CA, United States Assigned to Union Oil Company of California." Environment International 22(2): XVII.
Not Relevant to current fit for purpose review	(1996). "5412544 Method of illuminating and providing emergency egress guidance for hazardous areas: Derrick Donald E; Harris Hollis A; Marion Robert H; Tower William A; Towle L Christophe Hanover, NH, United States Assigned to Loctite Luminescent Systems Inc; The MTL Instruments Group." Environment International 22(2): XVII.
Not Relevant to current fit for purpose review	(1996). "Animal breeding and infertility: M.J. Meredith (Editor), Blackwell Science, Oxford, 1995, 508 pp., £60.00, ISBN 0-632-04038-6." Animal Reproduction Science 44(2): 135-136.
Not Relevant to current fit for purpose review	(1996). "Preliminary program of the ninety-sixth annual session, May 11-15, 1996." American Journal of Orthodontics and Dentofacial Orthopedics 109(2): 196-214.
Not Relevant to current fit for purpose review	(1996). "Roundup of Federal Regulations on Food and Nutrition Issues." Journal of the American Dietetic Association 96(7): 654.
Not Relevant to current fit for purpose review	(1996). "ROUND-UP Research." Reproductive Health Matters 4(8): 149-153.
Not Relevant to current fit for purpose review	(1997). "ROUND-UP Conferences." Reproductive Health Matters 5(10): 180.
Not Relevant to current fit for purpose review	(1997). "ROUND-UP Research." Reproductive Health Matters 5(10): 162-167.
Not Relevant to current fit for purpose review	(1998). "E. coli antiserum." Journal of Equine Veterinary Science 18(8): 507.
Not Relevant to current fit for purpose review	(1998). "EIA in wild horses." Journal of Equine Veterinary Science 18(8): 507.
Not Relevant to current fit for purpose review	(1998). "Exercise-induced endotoxemia." Journal of Equine Veterinary Science 18(8): 506-507.
Not Relevant to current fit for purpose review	(1998). "Flamel Technologies sur les starting-blocks." Biofutur 1998(176): 41.
Not Relevant to current fit for purpose review	(1998). "Legislative roundup: protection of human subjects key issue for Congress." J Natl Cancer Inst 90(2): 97-98.
Not Relevant to current fit for purpose review	(1998). "ROUND-UP Conferences." Reproductive Health Matters <HT>6</HT>(12): 186.

Not Relevant to current fit for purpose review	(1999). "Research poster summaries from the ENA 1999 Annual Meeting." Journal of Emergency Nursing 25(<HT>6</HT>): 446-456.
Not Relevant to current fit for purpose review	(1999). "ROUND-UP Publications." Reproductive Health Matters 7(14): 196-201.
Not Relevant to current fit for purpose review	(2000). "Genome roundup." Nat Biotechnol 18(1): 9.
Not Relevant to current fit for purpose review	(2000). "ROUND-UP Law and Policy." Reproductive Health Matters 8(15): 171-174.
Not Relevant to current fit for purpose review	(2000). "ROUND-UP Research." Reproductive Health Matters 8(16): 190-192.
Not Relevant to current fit for purpose review	(2000). "Trauma news today." International Journal of Trauma Nursing <HT>6</HT>(3): 105-108.
Not Relevant to current fit for purpose review	(2001). "ROUND-UP Research." Reproductive Health Matters 9(18): 196-198.
Not Relevant to current fit for purpose review	(2002). "ROUND-UP Research." Reproductive Health Matters 10(19): 209-210.
Not Relevant to current fit for purpose review	(2003). "New Web site lights childhood obesity with fun." Journal of the American Dietetic Association 103(<HT>6</HT>): 671-672.
Not Relevant to current fit for purpose review	(2003). "ROUND-UP Law and Policy." Reproductive Health Matters 11(22): 199-203.
Not Relevant to current fit for purpose review	(2003). "ROUND-UP Research." Reproductive Health Matters 11(21): 201-204.
Not Relevant to current fit for purpose review	(2004). "Contemporary issues in women's health." International Journal of Gynecology & Obstetrics 87(2): 111-113.
Not Relevant to current fit for purpose review	(2004). "Contents page, CD logo." The Journal of Men's Health & Gender 1(4): 283-284.
Not Relevant to current fit for purpose review	(2004). "Editorial Board." European Journal of Oncology Nursing 8(4): i.
Not Relevant to current fit for purpose review	(2004). "Editorial Board." European Journal of Oncology Nursing 8(3): i.
Not Relevant to current fit for purpose review	(2004). "Editorial Board." European Journal of Oncology Nursing 8, Supplement 1: i.
Not Relevant to current fit for purpose review	(2004). "Editorial Board." European Journal of Oncology Nursing 8, Supplement 2: i.
Not Relevant to current fit for purpose review	(2004). "Research round-up." Complementary Therapies in Nursing and Midwifery 10(4): 262-263.
Not Relevant to current fit for purpose review	(2004). "ROUND-UP Condoms." Reproductive Health Matters 12(23): 176-177.
Not Relevant to current fit for purpose review	(2004). "ROUND-UP Publications." Reproductive Health Matters 12(24): 231-236.
Not Relevant to current fit for purpose review	(2004). "ROUND-UP Service Delivery." Reproductive Health Matters 12(23): 191-200.
Not Relevant to current fit for purpose review	(2004). "Volume Contents Index." The Journal of Men's Health & Gender 1(4): 413-415.
Not Relevant to current fit for purpose review	(2004). "WebWatch." The Journal of Men's Health & Gender 1(2-3): 240.
Not Relevant to current fit for purpose review	(2004). "You've Been Had! How the Media and Environmentalists Turned America into a Nation of Hypochondriacs: Melvin A. Benarde. Rutgers University Press, 2002. xiv + 308 pp. \$28.00. ISBN 0-8135-3050-4." Chemical Health and Safety 11(2): 39-40.
Not Relevant to current fit for purpose review	(2005). "Contents page, CD logo." The Journal of Men's Health & Gender 2(4): 387.
Not Relevant to current fit for purpose review	(2005). "Contents page, CD logo." The Journal of Men's Health & Gender 2(2): 159-160.
Not Relevant to current fit for purpose review	(2005). "Editorial Board." European Journal of Oncology Nursing 9(4): i.

Not Relevant to current fit for purpose review	(2005). "Editorial Board." European Journal of Oncology Nursing 9(3): i.
Not Relevant to current fit for purpose review	(2005). "Editorial Board." European Journal of Oncology Nursing 9(2): i.
Not Relevant to current fit for purpose review	(2005). "Editorial Board." European Journal of Oncology Nursing 9(1): i.
Not Relevant to current fit for purpose review	(2005). "JournalWatch." The Journal of Men's Health & Gender 2(3): 353-356.
Not Relevant to current fit for purpose review	(2005). "News Round-up." The Journal of Men's Health & Gender 2(3): 360-363.
Not Relevant to current fit for purpose review	(2005). "News Round-up." The Journal of Men's Health & Gender 2(1): 116-118.
Not Relevant to current fit for purpose review	(2005). "News Round-up for jmhg vol 2 no 4 (Dec 2005)." The Journal of Men's Health & Gender 2(4): 444.
Not Relevant to current fit for purpose review	(2005). "Research round-up : a brief summary of research publications in CAM." Complementary Therapies in Clinical Practice 11(2): 139-141.
Not Relevant to current fit for purpose review	(2006). "ACFAOM newsletter." The Foot 16(4): 226-227.
Not Relevant to current fit for purpose review	(2006). "Aims & Scope/Editorial board." The Journal of Men's Health & Gender 3(1): iii.
Not Relevant to current fit for purpose review	(2006). "Contents page, CD logo." The Journal of Men's Health & Gender 3(2): 119-120.
Not Relevant to current fit for purpose review	(2006). "Contents page, CD logo." The Journal of Men's Health & Gender 3(4): 315-316.
Not Relevant to current fit for purpose review	(2006). "Editorial Board." European Journal of Oncology Nursing 10(5): i.
Not Relevant to current fit for purpose review	(2006). "Editorial Board." European Journal of Oncology Nursing 10(4): i.
Not Relevant to current fit for purpose review	(2006). "Editorial Board." European Journal of Oncology Nursing 10(3): i.
Not Relevant to current fit for purpose review	(2006). "Editorial Board." European Journal of Oncology Nursing 10(2): i.
Not Relevant to current fit for purpose review	(2006). "Editorial Board." European Journal of Oncology Nursing 10(1): i.
Not Relevant to current fit for purpose review	(2006). "Editorial Board/Publication/Advertising info." Progress in Biophysics and Molecular Biology 92(1): CO2.
Not Relevant to current fit for purpose review	(2006). "News Round-up." The Journal of Men's Health & Gender 3(1): 93-96.
Not Relevant to current fit for purpose review	(2006). "News Round-up for jmhg vol 3 no 3 (September 2006)." The Journal of Men's Health & Gender 3(3): 304-306.
Not Relevant to current fit for purpose review	(2006). "Research." Reproductive Health Matters 14(28): 210-218.
Not Relevant to current fit for purpose review	(2006). "Research problems." Discrete Applied Mathematics 154(3): 604-607.
Not Relevant to current fit for purpose review	(2006). "Table of Contents." The American Journal of Emergency Medicine 24(4): A3-A4.
Not Relevant to current fit for purpose review	(2006). "Volume Contents Index: Volume 3." The Journal of Men's Health & Gender 3(4): 427-431.
Not Relevant to current fit for purpose review	(2006). "WebWatch for jmhg vol 3 no 1 (issue 9)." The Journal of Men's Health & Gender 3(1): 88.
Not Relevant to current fit for purpose review	(2007). "Art exhibit powered by geothermal energy." Renewable Energy Focus 8(6): 14.
Not Relevant to current fit for purpose review	(2007). "Contents." The Journal of Men's Health & Gender 4(2): 113-114.

Not Relevant to current fit for purpose review	(2007). "Contents page." The Journal of Men's Health & Gender 4(1): 1-2.
Not Relevant to current fit for purpose review	(2007). "Editorial Board." European Journal of Oncology Nursing 11, Supplement 2: ii.
Not Relevant to current fit for purpose review	(2007). "Editorial Board." European Journal of Oncology Nursing 11(5): i.
Not Relevant to current fit for purpose review	(2007). "Editorial Board." European Journal of Oncology Nursing 11(4): i.
Not Relevant to current fit for purpose review	(2007). "Editorial Board." European Journal of Oncology Nursing 11(3): i.
Not Relevant to current fit for purpose review	(2007). "Editorial Board." European Journal of Oncology Nursing 11(2): i.
Not Relevant to current fit for purpose review	(2007). "Editorial Board." European Journal of Oncology Nursing 11(1): i.
Not Relevant to current fit for purpose review	(2007). "In brief." Renewable Energy Focus 8(<HT>6</HT>): 18-19.
Not Relevant to current fit for purpose review	(2007). "North America." Renewable Energy Focus 8(5): 6.
Not Relevant to current fit for purpose review	(2007). "US could "eliminate CO2 emissions without nuclear power"." Renewable Energy Focus 8(5): 18.
Not Relevant to current fit for purpose review	(2008). "Editorial Board." European Journal of Oncology Nursing 12(5): i.
Not Relevant to current fit for purpose review	(2008). "Editorial Board." European Journal of Oncology Nursing 12(1): i.
Not Relevant to current fit for purpose review	(2008). "Editorial Board." European Journal of Oncology Nursing 12(4): i.
Not Relevant to current fit for purpose review	(2008). "Editorial Board." European Journal of Oncology Nursing 12(3): i.
Not Relevant to current fit for purpose review	(2008). "Editorial Board." European Journal of Oncology Nursing 12(2): i.
Not Relevant to current fit for purpose review	(2008). "In Brief." Renewable Energy Focus 9(3): 16.
Not Relevant to current fit for purpose review	(2008). "In brief – projects." Renewable Energy Focus 9(<HT>6</HT>): 15.
Not Relevant to current fit for purpose review	(2008). "Swiss FiT on slippery slope?" Renewable Energy Focus 9(5): 12.
Not Relevant to current fit for purpose review	(2008). "US wind heavyweights unite over PTC." Renewable Energy Focus 9(3): 17.
Not Relevant to current fit for purpose review	(2008). "Wärtsilä BioPower plant to use brewery spent grain." Renewable Energy Focus 9(2): 12.
Not Relevant to current fit for purpose review	(2009). "Acknowledgements." European Journal of Agronomy 31(3): iv.
Not Relevant to current fit for purpose review	(2009). "Carbon Trust calls for switch to biomass heating." Renewable Energy Focus 10(2): 15.
Not Relevant to current fit for purpose review	(2009). "Editorial Board." European Journal of Oncology Nursing 13(4): i.
Not Relevant to current fit for purpose review	(2009). "Editorial Board." European Journal of Oncology Nursing 13(3): i.
Not Relevant to current fit for purpose review	(2009). "Editorial Board." European Journal of Oncology Nursing 13(2): i.
Not Relevant to current fit for purpose review	(2009). "Editorial Board." European Journal of Oncology Nursing 13(1): i.
Not Relevant to current fit for purpose review	(2009). "EU renewables in short." Renewable Energy Focus 9(7): 14.

Not Relevant to current fit for purpose review	(2009). "Pickens scraps "mega" wind plans." Renewable Energy Focus 10(5): 18.
Not Relevant to current fit for purpose review	(2009). "Policy still needed for solar growth." Renewable Energy Focus 10(6): 16.
Not Relevant to current fit for purpose review	(2009). "Shell withdraws from wind and solar — focuses on biofuels." Renewable Energy Focus 10(3): 19.
Not Relevant to current fit for purpose review	(2010). "Communications orales du 27e congrès de la SFE." Annales d'Endocrinologie 71(5): 340-353.
Not Relevant to current fit for purpose review	(2010). "Ten GW US wind in 2009." Renewable Energy Focus 11(1): 13.
Not Relevant to current fit for purpose review	(2010). "UK introduces feed-in tariffs." Renewable Energy Focus 11(1): 8.
Not Relevant to current fit for purpose review	(2011). "€51 billion offshore wind market." Renewable Energy Focus 12(6): 12.
Not Relevant to current fit for purpose review	(2011). "December, 2010." The Lancet Neurology 10(3): 209.
Not Relevant to current fit for purpose review	(2011). "Directions & Connections." Journal of the American Medical Directors Association 12(8): A14-A15.
Not Relevant to current fit for purpose review	(2011). "Directions & Connections." Journal of the American Medical Directors Association 12(7): A14-A15.
Not Relevant to current fit for purpose review	(2011). "Directions & Connections." Journal of the American Medical Directors Association 12(9): A10-A11.
Not Relevant to current fit for purpose review	(2011). "Editorial Board." Spectrochimica Acta Part A: Molecular and Biomolecular Spectroscopy 78(5): CO2.
Not Relevant to current fit for purpose review	(2011). "EU: huge renewables growth if projections met." Renewable Energy Focus 12(6): 10.
Not Relevant to current fit for purpose review	(2011). "Table of Contents." Regulatory Toxicology and Pharmacology 61(2): iii.
Not Relevant to current fit for purpose review	(2011). "Table of Contents." The American Journal of Emergency Medicine 29(4): A5-A7.
Not Relevant to current fit for purpose review	(2012). "Contents." The American Journal of Medicine 125(8): A5-A7.
Not Relevant to current fit for purpose review	(2012). "Directions & Connections." Journal of the American Medical Directors Association 13(1): A10-A11.
Not Relevant to current fit for purpose review	(2012). "Directions & Connections." Journal of the American Medical Directors Association 13(2): A14-A15.
Not Relevant to current fit for purpose review	(2012). "Impact sanitaire de la pollution environnementale et solutions pour la dépollution cellulaire." La Revue d'Homéopathie 3(3): 115-117.
Not Relevant to current fit for purpose review	(2012). "Revue de presse." La Revue d'Homéopathie 3(4): 159-160.
Not Relevant to current fit for purpose review	(2013). "December, 2012." The Lancet Neurology 12(3): 230.
Not Relevant to current fit for purpose review	(2015). "60 Seconds." New Scientist 225(3014): 7.
Not Relevant to current fit for purpose review	(2015). "60 Seconds." New Scientist 226(3022): 7.
Not Relevant to current fit for purpose review	(2015). "GLYPHOSATE UNLIKELY TO CAUSE CANCER, EU AGENCY FINDS." Chemical & Engineering News 93(45): 26-26.
Not Relevant to current fit for purpose review	(2016). "EDITORIAL BOARD." NeuroToxicology 52: IFC.
Not Relevant to current fit for purpose review	Abdullahi, A. E. (2002). "Cynodon dactylon control with tillage and glyphosate." Crop Protection 21(10): 1093-1100.
Not Relevant to current fit for purpose review	Acar, A., et al. (2015). "The Investigation of Genotoxic, Physiological and Anatomical Effects of Paraquat Herbicide on Allium cepa L." Cytologia 80(3): 343-351.

Not Relevant to current fit for purpose review	Acquavella, J., et al. (1999). "A case-control study of non-Hodgkin lymphoma and exposure to pesticides." <i>Cancer</i> 86(4): 729-731.
Not Relevant to current fit for purpose review	Aicher, R. H. (1999). "Vicarious Liability." <i>Aesthetic Surgery Journal</i> 19(4): 341.
Not Relevant to current fit for purpose review	Akaza, H. (2007). "Report from the 1st Japanese Urological Association-Japanese Society of Medical Oncology joint conference, 2006: 'A step towards better collaboration between urologists and medical oncologists'." <i>Int J Urol</i> 14(5): 375-383.
Not Relevant to current fit for purpose review	Akert, K. (1959). "Morphologie und physiologie des nervensystems: Paul Glees Georg Thieme, Verlag, Stuttgart, 1957, 445 pp. \$13.50." <i>Electroencephalography and Clinical Neurophysiology</i> 11(1): 191-192.
Not Relevant to current fit for purpose review	Albrecht, B., et al. (1991). "In vivo utilization of N-(phosphonomethyl)-anilines and related substances by <i>Pseudomonas spec. GS</i> ." <i>J Basic Microbiol</i> 31(6): 403-411.
Not Relevant to current fit for purpose review	Altenhoff, A. M. and C. Dessimoz (2009). "Phylogenetic and functional assessment of orthologs inference projects and methods." <i>PLoS Comput Biol</i> 5(1): e1000262.
Not Relevant to current fit for purpose review	Amaro, P. (2012). "O mancozebe é um ótimo exemplo da diversidade de classificação toxicológica dos pesticidas em Portugal." <i>Revista de Ciências Agrárias</i> 35(2): 118-125.
Not Relevant to current fit for purpose review	Andersson, K.-E. (2011). "This Month in Investigative Urology." <i>The Journal of Urology</i> 185(4): 1176.
Not Relevant to current fit for purpose review	Angelini, C. (2012). "Neuromuscular diseases: advances in therapy and diagnosis." <i>The Lancet Neurology</i> 11(1): 15-17.
Not Relevant to current fit for purpose review	Anklam, E. (1999). "The validation of methods based on polymerase chain reaction for the detection of genetically modified organisms in food." <i>Analytica Chimica Acta</i> 393(1-3): 177-179.
Not Relevant to current fit for purpose review	Antonini, A. (2012). "Movement disorders: towards new therapies in Parkinson's disease." <i>The Lancet Neurology</i> 11(1): 7-8.
Not Relevant to current fit for purpose review	Arjo, G., et al. (2013). "Plurality of opinion, scientific discourse and pseudoscience: an in depth analysis of the Seralini et al. study claiming that Roundup (TM) Ready corn or the herbicide Roundup (TM) cause cancer in rats." <i>Transgenic Res</i> 22(2): 255-267.
Not Relevant to current fit for purpose review	Astudillo, O. (2012). "Clinical trials: round-up." <i>The Lancet Oncology</i> 13(10): 976.
Not Relevant to current fit for purpose review	Atkinson, S. (2003). "Product developments in biotechnology and medical applications." <i>Membrane Technology</i> 2003(2): 10-11.
Not Relevant to current fit for purpose review	Avella, P., et al. (2007). "Metric inequalities and the Network Loading Problem." <i>Discrete Optimization</i> 4(1): 103-114.
Not Relevant to current fit for purpose review	Axelrad, J. C., et al. (2002). "Enhanced in vitro toxicity of the herbicide glyphosate to neuroblastoma cells chronically pre-treated with the organophosphate pesticide diazinon." <i>Toxicology</i> 178(1): 62-63.
Not Relevant to current fit for purpose review	Axelrad, J. C., et al. (2003). "The effects of acute pesticide exposure on neuroblastoma cells chronically exposed to diazinon." <i>Toxicology</i> 185(1-2): 67-78.
Not Relevant to current fit for purpose review	Azevedo, L., et al. (2010). "In Vivo Antimutagenic Properties of Transgenic and Conventional Soybeans." <i>J Med Food</i> 13(6): 1402-1408.
Not Relevant to current fit for purpose review	Bababunmi, E. A., et al. (1979). "The uncoupling effect of N-(phosphonomethyl)glycine on isolated rat liver mitochondria." <i>Biochemical Pharmacology</i> 28(<HT>6</HT>): 925-927.
Not Relevant to current fit for purpose review	Badr, A., et al. (2013). "Cytophysiological impacts of Metosulam herbicide on <i>Vicia faba</i> plants." <i>Acta Physiologiae Plantarum</i> 35(6): 1933-1941.
Not Relevant to current fit for purpose review	Barker, W. F. (1995). "The Western Vascular Society: Its first ten years." <i>Journal of Vascular Surgery</i> 22(4): 505-506.
Not Relevant to current fit for purpose review	Baron, R., et al. (2012). "Chronic pain: genes, plasticity, and phenotypes." <i>The Lancet Neurology</i> 11(1): 19-21.
Not Relevant to current fit for purpose review	Bar-Or, A. (2016). "Multiple sclerosis and related disorders: evolving pathophysiologic insights." <i>The Lancet Neurology</i> 15(1): 9-11.
Not Relevant to current fit for purpose review	Bartelink, H., et al. (1993). "Foreword." <i>Radiotherapy and Oncology</i> 29(2): xi-xii.
Not Relevant to current fit for purpose review	Bates, J. A. (2004). "Use of narrative interviewing in everyday information behavior research." <i>Library & Information Science Research</i> 26(1): 15-28.
Not Relevant to current fit for purpose review	Begna, S. H., et al. (2001). "Morphology and yield response to weed pressure by corn hybrids differing in canopy architecture." <i>European Journal of Agronomy</i> 14(4): 293-302.

Not Relevant to current fit for purpose review	Benachour, N., et al. (2007). "Time- and dose-dependent effects of roundup on human embryonic and placental cells." <i>Arch Environ Contam Toxicol</i> 53(1): 126-133.
Not Relevant to current fit for purpose review	Ben-Menachem, E. (2016). "Epilepsy in 2015: the year of collaborations for big data." <i>The Lancet Neurology</i> 15(1): 6-7.
Not Relevant to current fit for purpose review	Berdal, K. G. and A. Holst-Jensen (2001). "Roundup Ready (R) soybean event-specific real-time quantitative PCR assay and estimation of the practical detection and quantification limits in GMO analyses." <i>European Food Research and Technology</i> 213(6): 432-438.
Not Relevant to current fit for purpose review	Berer, M. (1996). "Men." <i>Reproductive Health Matters</i> 4(7): 7-10.
Not Relevant to current fit for purpose review	Berkovic, S. F. (2010). "Epilepsy: insights into causes and treatment dilemmas." <i>The Lancet Neurology</i> 9(1): 9-11.
Not Relevant to current fit for purpose review	Bermel, R. A. and J. A. Cohen (2011). "Multiple sclerosis: advances in understanding pathogenesis and emergence of oral treatment options." <i>The Lancet Neurology</i> 10(1): 4-5.
Not Relevant to current fit for purpose review	Bertheussen, K., et al. (1997). "A new sensitive cell culture test for the assessment of pesticide toxicity." <i>Journal of Environmental Science and Health Part B-Pesticides Food Contaminants and Agricultural Wastes</i> 32(2): 195-211.
Not Relevant to current fit for purpose review	Bertucci, A., et al. (2015). "Detection of unamplified genomic DNA by a PNA-based microstructured optical fiber (MOF) Bragg-grating optofluidic system." <i>Biosens Bioelectron</i> 63: 248-254.
Not Relevant to current fit for purpose review	Bf (1995). "Let's Get Things Moving: Overcoming constipation." <i>Physiotherapy</i> 81(5): 301.
Not Relevant to current fit for purpose review	Bjerrum-Bohr, J. J., et al. (2012). "Hydrodynamics of a quark droplet." <i>Nuclear Physics A</i> 882: 90-106.
Not Relevant to current fit for purpose review	Blumann, W. and D. Fauth (1993). "Expensive NM standards round-up: Network Management Standards: OSI, SNMP and CMOL Protocols by Ulyses Black. Published by McGraw Hill, 1992, £38.95, 336 pp." <i>Computer Communications</i> 16(7): 449.
Not Relevant to current fit for purpose review	Bogler, O. and R. Sawaya (2008). "Foreword." <i>Current Problems in Cancer</i> 32(3): 95-96.
Not Relevant to current fit for purpose review	Bonn, D. (2005). "Web round-up: rare-tumour information." <i>The Lancet Oncology</i> <HT>6</HT>(3): 144.
Not Relevant to current fit for purpose review	Botha, G. M. and C. D. Viljoen (2009). "South Africa: A case study for voluntary GM labelling." <i>Food Chemistry</i> 112(4): 1060-1064.
Not Relevant to current fit for purpose review	Bournat, P. (1997). "Les techniques de transgénèse végétale." <i>Biofutur</i> 1997(172): 3-7.
Not Relevant to current fit for purpose review	Boutet, I., et al. (2004). "Molecular identification and expression of two non-P450 enzymes, monoamine oxidase A and flavin-containing monooxygenase 2, involved in phase I of xenobiotic biotransformation in the Pacific oyster, <i>Crassostrea gigas</i> ." <i>Biochim Biophys Acta</i> 1679(1): 29-36.
Not Relevant to current fit for purpose review	Bowersox, T. W., et al. (1990). "Coppicing success of young <i>Eucalyptus saligna</i> in Hawaii." <i>Biomass</i> 23(2): 137-148.
Not Relevant to current fit for purpose review	Bowman, W. C. (1983). "Drugs and anesthesia—Pharmacology for anesthesiologists." <i>Trends in Pharmacological Sciences</i> 4: 358-359.
Not Relevant to current fit for purpose review	Brandenberger, L. P., et al. (2005). "Preemergence weed control in direct-seeded watermelon." <i>Weed Technology</i> 19(3): 706-712.
Not Relevant to current fit for purpose review	Breaden, A. (1991). "Victoria ASCCN." <i>Confederation of Australian Critical Care Nurses Journal</i> 4(3): 6.
Not Relevant to current fit for purpose review	Briscoe, M. and C. P. Wynne (1988). "State Round-Up Tasmanian Branch of A.S.C.C.N." <i>CNSA Journal</i> 1(1): 3-7.
Not Relevant to current fit for purpose review	Burkhart, C. G. (1981). "Looking at eyelid lesions--a clinical roundup." <i>Geriatrics</i> 36(8): 91-95.
Not Relevant to current fit for purpose review	Burr, T. J., et al. (1995). "SURVIVAL AND TUMORIGENICITY OF AGROBACTERIUM-VITIS IN LIVING AND DECAYING GRAPE ROOTS AND CANES IN SOIL." <i>Plant Disease</i> 79(7): 677-682.
Not Relevant to current fit for purpose review	Bushby, K. (2011). "Neuromuscular diseases: milestones in development of treatments." <i>The Lancet Neurology</i> 10(1): 11-13.
Not Relevant to current fit for purpose review	Butler, T. J., et al. (2002). "Broomsedge (<i>Andropogon virginicus</i>) response to herbicides and burning." <i>Weed Technology</i> 16(1): 18-22.

Not Relevant to current fit for purpose review	Cai, Z., et al. (2012). "3.102 PERINATAL LPS EXPOSURE INCREASES THE RISK FOR DOPAMINERGIC DISORDERS IN ADULT LIFE." <i>Parkinsonism & Related Disorders</i> 18, Supplement 2: S190.
Not Relevant to current fit for purpose review	Caloni, F., et al. (2015). "In vitro effects of glyphosate on cell proliferation and steroid production by bovine granulosa cells." <i>Toxicology Letters</i> 238(2, Supplement): S290.
Not Relevant to current fit for purpose review	Campbell, M. H. and A. R. Gilmour (1979). "EFFECT OF TIME AND RATE OF APPLICATION OF HERBICIDES ON SERRATED TUSsock (NASSELLA-TRICHOTOMA) AND IMPROVED PASTURE SPECIES .1. GLYPHOSATE AND 2,2-DPA." <i>Australian Journal of Experimental Agriculture</i> 19(99): 472-475.
Not Relevant to current fit for purpose review	Cao, G., et al. (2013). "Draft genome sequence of pseudomonas strain p818, isolated from glyphosate-polluted soil." <i>Genome Announc</i> 1(6).
Not Relevant to current fit for purpose review	Casorri, L., et al. (2010). "Testing Round-up Ready soybean along processing chain by traditional and innovative methods." <i>Journal of Biotechnology</i> 150, Supplement: 536.
Not Relevant to current fit for purpose review	Castillo, M., et al. (2014). "Seedbed Preparation Techniques and Weed Control Strategies for Strip-Planting Rhizoma Peanut into Warm-Season Grass Pastures." <i>Crop Science</i> 54(4): 1868-1875.
Not Relevant to current fit for purpose review	Cavusoglu, K., et al. (2011). "Protective effect of Ginkgo biloba L. leaf extract against glyphosate toxicity in Swiss albino mice." <i>J Med Food</i> 14(10): 1263-1272.
Not Relevant to current fit for purpose review	Cebeci, O. and H. Budak (2009). "Global Expression Patterns of Three Festuca Species Exposed to Different Doses of Glyphosate Using the Affymetrix GeneChip Wheat Genome Array." <i>Comp Funct Genomics</i> : 505701.
Not Relevant to current fit for purpose review	Chang, H. S., et al. (2001). "Allergenicity test of genetically modified soybean in Sprague-Dawley rats." <i>Archives of Pharmacal Research</i> 24(3): 256-261.
Not Relevant to current fit for purpose review	Chang, J.-S., et al. (1999). "Transformation of rat fibroblasts by phospholipase C- γ 1 overexpression is accompanied by tyrosine dephosphorylation of paxillin." <i>FEBS Letters</i> 460(1): 161-165.
Not Relevant to current fit for purpose review	Charles, A. (2011). "Headache: new genes, new mechanisms, and new therapies." <i>The Lancet Neurology</i> 10(1): 13-14.
Not Relevant to current fit for purpose review	Charlton, C. G. and G. V. Muthian (2012). "3.101 PRENATAL SENSITIZATION AND POSTNATAL PRECIPITATION IN A MODEL OF PARKINSONISM." <i>Parkinsonism & Related Disorders</i> 18, Supplement 2: S189-S190.
Not Relevant to current fit for purpose review	Chee, P. P. and J. L. Slightom (1991). "TRANSFER AND EXPRESSION OF CUCUMBER MOSAIC-VIRUS COAT PROTEIN GENE IN THE GENOME OF CUCUMIS-SATIVUS." <i>Journal of the American Society for Horticultural Science</i> 116(6): 1098-1102.
Not Relevant to current fit for purpose review	Chen, L., et al. (2009). "Modeling Multi-typed Structurally Viewed Chemicals with the UMLS Refined Semantic Network." <i>Journal of the American Medical Informatics Association</i> 16(1): 116-131.
Not Relevant to current fit for purpose review	Chiò, A. and G. Lauria (2016). "Degenerative neuromuscular diseases: crucial gene and cell machinery discoveries." <i>The Lancet Neurology</i> 15(1): 12-13.
Not Relevant to current fit for purpose review	Chorfa, A., et al. (2013). "Specific Pesticide-Dependent Increases in alpha-Synuclein Levels in Human Neuroblastoma (SH-SY5Y) and Melanoma (SK-MEL-2) Cell Lines." <i>Toxicological Sciences</i> 133(2): 289-297.
Not Relevant to current fit for purpose review	Cichosz, G. and S. K. Wiackowski (2012). "[Genetically modified food--great unknown]." <i>Pol Merkur Lekarski</i> 33(194): 59-63.
Not Relevant to current fit for purpose review	Cooke, R. M. (1997). "Protein NMR extends into new fields of structural biology." <i>Current Opinion in Chemical Biology</i> 1(3): 359-364.
Not Relevant to current fit for purpose review	Cornish-Bowden, A. and M. L. Cardenas (2003). "Metabolic analysis in drug design." <i>Comptes Rendus Biologies</i> 326(5): 509-515.
Not Relevant to current fit for purpose review	Costa, M. d. D. L. d., et al. (2003). "Alterações de neuroimagem no parkinsonismo: estudo de cinco casos." <i>Arquivos De Neuro-Psiquiatria</i> 61(2B): 381-386.
Not Relevant to current fit for purpose review	Costello, J. (2015). "Research roundup." <i>Int J Palliat Nurs</i> 21(8): 410-411.
Not Relevant to current fit for purpose review	Coyle, C. (2003). "A Practical Guide to Intensity – Modulated Radiation Therapy." <i>Clinical Oncology</i> 15(8): 507.
Not Relevant to current fit for purpose review	Croft, S. M., et al. (1974). "Inhibition of chloroplast ribosome formation by N,N-bis(phosphonomethyl)glycine." <i>Biochimica et Biophysica Acta (BBA) - Nucleic Acids and Protein Synthesis</i> 335(2): 211-217.

Not Relevant to current fit for purpose review	Cross, J. R. (1995). "Modern Chinese Acupuncture - A review of acupuncture techniques as practised in China today." <i>Physiotherapy</i> 81(5): 302.
Not Relevant to current fit for purpose review	Cui, J., et al. (2011). "Detecting biological network organization and functional gene orthologs." <i>Bioinformatics</i> 27(20): 2919-2920.
Not Relevant to current fit for purpose review	Culbreth, M. E., et al. (2012). "Comparison of chemical-induced changes in proliferation and apoptosis in human and mouse neuroprogenitor cells." <i>NeuroToxicology</i> 33(6): 1499-1510.
Not Relevant to current fit for purpose review	Cundiff, J. S., et al. (1997). "A linear programming approach for designing a herbaceous biomass delivery system." <i>Bioresource Technology</i> 59(1): 47-55.
Not Relevant to current fit for purpose review	da Costa, M. D. L., et al. (2003). "Neuroimaging abnormalities in parkinsonism: study of five cases." <i>Arquivos De Neuro-Psiquiatria</i> 61(2B): 381-386.
Not Relevant to current fit for purpose review	D'Amico, D. B. (1978). "Critics prompt plans to revise health planning guidelines." <i>AORN Journal</i> 27(4): 702-706.
Not Relevant to current fit for purpose review	Dan, B. and P. Baxter (2014). "Paediatric neurology: a year of DNA technology." <i>The Lancet Neurology</i> 13(1): 16-18.
Not Relevant to current fit for purpose review	Dan, Y. H., et al. (2006). "MicroTom-a high-throughput model transformation system for functional genomics." <i>Plant Cell Rep</i> 25(5): 432-441.
Not Relevant to current fit for purpose review	Daniel, R. K., et al. (1986). "Tissue transplants in primates for upper extremity reconstruction: A preliminary report." <i>The Journal of Hand Surgery</i> 11(1): 1-8.
Not Relevant to current fit for purpose review	Das, M., et al. (2010). "A composite transcriptional signature differentiates responses towards closely related herbicides in <i>Arabidopsis thaliana</i> and <i>Brassica napus</i> ." <i>Plant Mol Biol</i> 72(4-5): 545-556.
Not Relevant to current fit for purpose review	Das, M., et al. (2010). "A composite transcriptional signature differentiates responses towards closely related herbicides in <i>Arabidopsis thaliana</i> and <i>Brassica napus</i> ." <i>Plant Mol Biol</i> 72(4-5): 545-556.
Not Relevant to current fit for purpose review	Datta, R. and H. Daniell (1996). "Transformation of the tobacco chloroplast genome with the <i>arcA</i> gene to confer glyphosate tolerance." <i>Plant Physiol</i> 111(2): 790-790.
Not Relevant to current fit for purpose review	Davis, W. M. and I. W. Waters (1999). "A new-drug roundup." <i>Rn</i> 62(4): 54-60; quiz 62.
Not Relevant to current fit for purpose review	De Marco, A., et al. (1992). "Importance of the type of soil for the induction of micronuclei and the growth of primary roots of <i>Vicia faba</i> treated with the herbicides atrazine, glyphosate and maleic hydrazide." <i>Mutat Res</i> 279(1): 9-13.
Not Relevant to current fit for purpose review	DeAngelis, L. M. (2010). "Neuro-oncology: new hope for patients with gliomas." <i>The Lancet Neurology</i> 9(1): 17-18.
Not Relevant to current fit for purpose review	Defarge, N., et al. (2016). "Co-Formulants in Glyphosate-Based Herbicides Disrupt Aromatase Activity in Human Cells below Toxic Levels." <i>Int J Environ Res Public Health</i> 13(3).
Not Relevant to current fit for purpose review	Dehghan, R., et al. (2015). "Optimization of overlapping activities in the design phase of construction projects." <i>Automation in Construction</i> 59: 81-95.
Not Relevant to current fit for purpose review	DeLuca, T. F., et al. (2006). "Roundup: a multi-genome repository of orthologs and evolutionary distances." <i>Bioinformatics</i> 22(16): 2044-2046.
Not Relevant to current fit for purpose review	Demarco, A., et al. (1995). "INFLUENCE OF SOIL CHARACTERISTICS ON THE CLASTOGENIC ACTIVITY OF MALEIC HYDRAZIDE IN ROOT-TIPS OF VICIA-FABA." <i>Mutation Research-Genetic Toxicology</i> 344(1-2): 5-12.
Not Relevant to current fit for purpose review	Demarco, A., et al. (1992). "IMPORTANCE OF THE TYPE OF SOIL FOR THE INDUCTION OF MICRONUCLEI AND THE GROWTH OF PRIMARY ROOTS OF VICIA-FABA TREATED WITH THE HERBICIDES ATRAZINE, GLYPHOSATE AND MALEIC HYDRAZIDE." <i>Mutat Res</i> 279(1): 9-13.
Not Relevant to current fit for purpose review	Demeke, T. and I. Ratnayaka (2008). "Multiplex qualitative PCR assay for identification of genetically modified canola events and real-time event-specific PCR assay for quantification of the GT73 canola event." <i>Food Control</i> 19(9): 893-897.
Not Relevant to current fit for purpose review	Deming, D. (1989). "Application of bottom-hole temperature corrections in geothermal studies." <i>Geothermics</i> 18(5-<HT>6</HT>): 775-786.
Not Relevant to current fit for purpose review	Desimone, C., et al. (1992). "GENOTOXIC EFFECT INDUCED BY HERBICIDES ATRAZINE GLYPHOSATE IN PLANTS OF VICIA-FABA GROWN IN DIFFERENT SOILS." <i>Science of the Total Environment</i> 123: 233-240.
Not Relevant to current fit for purpose review	Diener, H.-C. (2004). "Important advances in headache." <i>The Lancet Neurology</i> 3(1): 12.

Not Relevant to current fit for purpose review	Dong Kyu Kim, M. D., et al. (2015). "The Factors Associated with the Hypotension Development in Acute Glyphosate-surfactant Herbicide Poisoning." <i>Journal of The Korean Society of Emergency Medicine</i> 26(3): 248-255.
Not Relevant to current fit for purpose review	Donnan, G. A. (2004). "Stroke: prediction, prevention, and outcome." <i>The Lancet Neurology</i> 3(1): 9.
Not Relevant to current fit for purpose review	Dragutinovich, S. (1987). "Stimulus intensity reducers: Are they sensation seekers, extraverts, and strong nervous types?" <i>Personality and Individual Differences</i> 8(5): 693-704.
Not Relevant to current fit for purpose review	Druwe, I., et al. (2015). "Sensitivity of neuroprogenitor cells to chemical-induced apoptosis using a multiplexed assay suitable for high-throughput screening." <i>Toxicology</i> 333: 14-24.
Not Relevant to current fit for purpose review	Duncan, K., et al. (1984). "The complete amino acid sequence of Escherichia coli 5-enolpyruvylshikimate 3-phosphate synthase." <i>FEBS Letters</i> 170(1): 59-63.
Not Relevant to current fit for purpose review	Dunn, S. V. (1999). "Recognising our limits: the murky shores where science meets values and beliefs." <i>Australian Critical Care</i> 12(4): 130.
Not Relevant to current fit for purpose review	Ehlert, A., et al. (2008). "Development of a modular system for detection of genetically modified organisms in food based on ligation-dependent probe amplification." <i>European Food Research and Technology</i> 227(3): 805-812.
Not Relevant to current fit for purpose review	Elgorashi, E. E., et al. (2004). "Isolation of captan from <i>Cyrtanthus suaveolens</i> : the effect of pesticides on the quality and safety of traditional medicine." <i>South African Journal of Botany</i> 70(4): 512-514.
Not Relevant to current fit for purpose review	Elie-Caille, C., et al. (2010). "Morphological damages of a glyphosate-treated human keratinocyte cell line revealed by a micro- to nanoscale microscopic investigation." <i>Cell Biol Toxicol</i> 26(4): 331-339.
Not Relevant to current fit for purpose review	Endo, M. and S. Toki (2013). "Creation of herbicide-tolerant crops by gene targeting." <i>Journal of Pesticide Science</i> 38(1-2): 49-59.
Not Relevant to current fit for purpose review	Engel, K.-H., et al. (2006). "Quantification of DNA from genetically modified organisms in composite and processed foods." <i>Trends in Food Science & Technology</i> 17(9): 490-497.
Not Relevant to current fit for purpose review	Erickson, B. (2015). "GLYPHOSATE LINKED TO CANCER." <i>Chemical & Engineering News</i> 93(13): 5-5.
Not Relevant to current fit for purpose review	Ewald, M., et al. (2014). "From surface to intracellular non-invasive nanoscale study of living cells impairments." <i>Nanotechnology</i> 25(29).
Not Relevant to current fit for purpose review	Farrington, J. W. (1979). "Newsletters." <i>Serials Review</i> 5(2): 55-57.
Not Relevant to current fit for purpose review	Favata, A., et al. (2014). "A nonlinear theory of prestressed elastic stick-and-spring structures." <i>International Journal of Engineering Science</i> 80: 4-20.
Not Relevant to current fit for purpose review	Feng, Z., et al. (2005). "The influence of GFP-actin expression on the adhesion dynamics of HepG2 cells on a model extracellular matrix." <i>Biomaterials</i> 26(26): 5348-5358.
Not Relevant to current fit for purpose review	Feriotto, G., et al. (2002). "Biosensor technology and surface plasmon resonance for real-time detection of genetically modified Roundup Ready soybean gene sequences." <i>J Agric Food Chem</i> 50(5): 955-962.
Not Relevant to current fit for purpose review	Ferrara, G., et al. (2000). "Evaluation of antimutagenic and desmutagenic effects of humic and fulvic acids on root tips of <i>Vicia faba</i> ." <i>Environmental Toxicology</i> 15(5): 513-517.
Not Relevant to current fit for purpose review	Ferrari, M. D. (2013). "Headache: the changing migraine brain." <i>The Lancet Neurology</i> 12(1): 6-8.
Not Relevant to current fit for purpose review	Fiers, T. (2004). "Lab-on-a-Chip. Miniaturised systems for (bio)chemical analysis and synthesis: (E. Oosterbroek, A. van den Berg) Elsevier. ISBN 0-444 51100-8." <i>Clinica Chimica Acta</i> 343(1-2): 245.
Not Relevant to current fit for purpose review	Figenschau, Y., et al. (1997). "A sensitive serum-free colorimetric assay for the detection of cytotoxic effects of pesticides." <i>Journal of Environmental Science and Health Part B-Pesticides Food Contaminants and Agricultural Wastes</i> 32(2): 177-194.
Not Relevant to current fit for purpose review	Fischer, G., et al. (2016). "Superficial Temporal Artery to Middle Cerebral Artery Bypass via a Minimized Approach: Operative Nuances and Problem-Solving Aspects." <i>World Neurosurgery</i> 88: 97-103.
Not Relevant to current fit for purpose review	Flóra, T. and Z. Simon (1982). "Thermische zersetzung des wirkstoffes von herbizid glyphosat." <i>Thermochimica Acta</i> 59(2): 125-132.
Not Relevant to current fit for purpose review	Forgacs, A. L., et al. (2012). "BLTK1 murine Leydig cells: a novel steroidogenic model for evaluating the effects of reproductive and developmental toxicants." <i>Toxicol Sci</i> 127(2): 391-402.
Not Relevant to current fit for purpose review	Forsyth, R. (2005). "Paediatrics: genetic insights and long-term follow-up." <i>The Lancet Neurology</i> 4(1): 8.

Not Relevant to current fit for purpose review	Fox, R. J. and R. A. Rudick (2004). "Multiple sclerosis: disease markers accelerate progress." <i>The Lancet Neurology</i> 3(1): 10.
Not Relevant to current fit for purpose review	Fox, S. (1999). "European Roundup - Positive news for Marimastat - British Biotech's drug shows promise against nonoperable gastric cancer." <i>Genetic Engineering News</i> 19(16): 20-+.
Not Relevant to current fit for purpose review	Fox, S. (2000). "European roundup - Genome research initiative launches in European union - Networking, training and mobility to be focus of integrated projects." <i>Genetic Engineering News</i> 20(21): 41-+.
Not Relevant to current fit for purpose review	Francisco Rossi, L., et al. (2016). "Chaetophractus villosus as a sentinel organism: Baseline values of mitotic index, chromosome aberrations and sister chromatid exchanges." <i>Mutation Research-Genetic Toxicology and Environmental Mutagenesis</i> 796: 40-45.
Not Relevant to current fit for purpose review	Fu, G. M., et al. (2016). "Optimization of liquid-state fermentation conditions for the glyphosate-degradation enzyme production of strain <i>Aspergillus oryzae</i> by ultraviolet mutagenesis." <i>Prep Biochem Biotechnol</i> .
Not Relevant to current fit for purpose review	Garry, V. F., et al. (2002). "Birth defects, season of conception, and sex of children born to pesticide applicators living in the Red River Valley of Minnesota, USA." <i>Environ Health Perspect</i> 110: 441-449.
Not Relevant to current fit for purpose review	Gasnier, C., et al. (2010). "Dig1 protects against cell death provoked by glyphosate-based herbicides in human liver cell lines." <i>J Occup Med Toxicol</i> 5: 29.
Not Relevant to current fit for purpose review	Gasnier, C., et al. (2009). "Glyphosate-based herbicides are toxic and endocrine disruptors in human cell lines." <i>Toxicology</i> 262(3): 184-191.
Not Relevant to current fit for purpose review	Gatty, A. (1990). "South Australian branch ASCCN." <i>Confederation of Australian Critical Care Nurses Journal</i> 3(3): 10.
Not Relevant to current fit for purpose review	Gayen, D., et al. (2013). "Comparative analysis of nutritional compositions of transgenic high iron rice with its non-transgenic counterpart." <i>Food Chemistry</i> 138(2-3): 835-840.
Not Relevant to current fit for purpose review	Giakoumaki, E., et al. (2003). "Combination of amplification and post-amplification strategies to improve optical DNA sensing." <i>Biosensors and Bioelectronics</i> 19(4): 337-344.
Not Relevant to current fit for purpose review	Giuffrida, M. C., et al. (2015). "Isothermal circular-strand-displacement polymerization of DNA and microRNA in digital microfluidic devices." <i>Anal Bioanal Chem</i> 407(6): 1533-1543.
Not Relevant to current fit for purpose review	Goldstein, C. and R. Chervin (2016). "Waking up to sleep research in 2015." <i>The Lancet Neurology</i> 15(1): 15-17.
Not Relevant to current fit for purpose review	Gonzalez, F. R. and A. M. Nardillo (1997). "Retention in multistep programmed-temperature gas chromatography and flow control Linear head pressure programs." <i>Journal of Chromatography A</i> 757(1-2): 109-118.
Not Relevant to current fit for purpose review	Gonzalez, N. V., et al. (2011). "A combination of the cytokinesis-block micronucleus cytome assay and centromeric identification for evaluation of the genotoxicity of dicamba." <i>Toxicology Letters</i> 207(3): 204-212.
Not Relevant to current fit for purpose review	Graus, F. and J. Dalmau (2012). "CNS autoimmunity: new findings and pending issues." <i>The Lancet Neurology</i> 11(1): 17-19.
Not Relevant to current fit for purpose review	Graus, F. and A. Tortosa (2006). "Neuro-oncology: setting new standards of management." <i>The Lancet Neurology</i> 5(1): 8-9.
Not Relevant to current fit for purpose review	Gregg, J. (1995). "Discipline, control, and the school mathematics tradition." <i>Teaching and Teacher Education</i> 11(<HT>6</HT>): 579-593.
Not Relevant to current fit for purpose review	Gress, S., et al. (2015). "Cardiotoxic Electrophysiological Effects of the Herbicide Roundup((R)) in Rat and Rabbit Ventricular Myocardium In Vitro." <i>Cardiovasc Toxicol</i> 15(4): 324-335.
Not Relevant to current fit for purpose review	Grosicka-Maciag, E., et al. (2012). "Dithiocarbamate fungicide zineb induces oxidative stress and apoptosis in Chinese hamster lung fibroblasts." <i>Pesticide Biochemistry and Physiology</i> 102(1): 95-101.
Not Relevant to current fit for purpose review	Gubbiga, N. G., et al. (1996). "Root/rhizome exudation of nicosulfuron from treated johnsongrass (<i>Sorghum halepense</i>) and possible implications for corn (<i>Zea mays</i>)." <i>Weed Science</i> 44(3): 455-460.
Not Relevant to current fit for purpose review	Guchi, T., et al. (2008). "Development of event-specific quantitation method for GA21 maize, which is a GM event without CaMV35S promoter." <i>Journal of the Food Hygienic Society of Japan</i> 49(1): 16-22.
Not Relevant to current fit for purpose review	Guo, B., et al. (2015). "Co-expression of G2-EPSPS and glyphosate acetyltransferase GAT genes conferring high tolerance to glyphosate in soybean." <i>Front Plant Sci</i> 6: 847.
Not Relevant to current fit for purpose review	Guyton, K. Z., et al. (2015). "Carcinogenicity of tetrachlorvinphos, parathion, malathion, diazinon, and glyphosate." <i>Lancet Oncology</i> 16(5): 490-491.
Not Relevant to current fit for purpose review	Hammond, B., et al. (2004). "Results of a 13 week safety assurance study with rats fed grain from glyphosate tolerant corn." <i>Food Chem Toxicol</i> 42(6): 1003-1014.

Not Relevant to current fit for purpose review	Hanberry, B. B., et al. (2013). "Small Mammal Responses to Intensively Established Pine Plantations in Coastal Plain Mississippi." <i>Southern Journal of Applied Forestry</i> 37(1): 53-58.
Not Relevant to current fit for purpose review	Haritou, M., et al. (2000). "Agents facilitating the electric field-induced fusion of intact rabbit erythrocytes." <i>Bioelectrochemistry</i> 52(2): 229-238.
Not Relevant to current fit for purpose review	Harkin, D. G., et al. (2000). "A novel approach to studying the migratory morphology of embryonic mesenchymal cells." <i>Biology of the Cell</i> 92(7): 537-543.
Not Relevant to current fit for purpose review	Harlow, S. D. and O. M. R. Campbell (2000). "Menstrual dysfunction: A missed opportunity for improving reproductive health in developing countries." <i>Reproductive Health Matters</i> 8(15): 142-147.
Not Relevant to current fit for purpose review	Hassannayebi, E. and S. H. Zegordi "Variable and adaptive neighbourhood search algorithms for rail rapid transit timetabling problem." <i>Computers & Operations Research</i> .
Not Relevant to current fit for purpose review	Hatamie, S., et al. (2015). "Curcumin-reduced graphene oxide sheets and their effects on human breast cancer cells." <i>Mater Sci Eng C Mater Biol Appl</i> 55: 482-489.
Not Relevant to current fit for purpose review	Haupt, S. E. (2006). "A quadratic empirical model formulation for dynamical systems using a genetic algorithm." <i>Computers & Mathematics with Applications</i> 51(3-4): 431-440.
Not Relevant to current fit for purpose review	Healy, C., et al. (2008). "Results of a 13-week safety assurance study with rats fed grain from corn rootworm-protected, glyphosate-tolerant MON 88017 corn." <i>Food and Chemical Toxicology</i> 46(7): 2517-2524.
Not Relevant to current fit for purpose review	Heck, G. R., et al. (2005). "Development and characterization of a CP4 EPSPS-Based, glyphosate-tolerant corn event." <i>Crop Science</i> 45(1): 329-339.
Not Relevant to current fit for purpose review	Henderson, N., et al. (2016). "PCR-Based Detection and Quantification of a Transgenic Glyphosate-Tolerant Canola Using a Novel Reference Gene System." <i>Food Analytical Methods</i> 9(2): 353-361.
Not Relevant to current fit for purpose review	Héraux, F. M. G., et al. (2005). "Combining <i>Trichoderma virens</i> -inoculated compost and a rye cover crop for weed control in transplanted vegetables." <i>Biological Control</i> 34(1): 21-26.
Not Relevant to current fit for purpose review	Hernández, A. F., et al. (2006). "Influence of exposure to pesticides on serum components and enzyme activities of cytotoxicity among intensive agriculture farmers." <i>Environmental Research</i> 102(1): 70-76.
Not Relevant to current fit for purpose review	Herrmann, K. M. and L. M. Weaver (1999). "The shikimate pathway." <i>Annual Review of Plant Physiology and Plant Molecular Biology</i> 50: 473-503.
Not Relevant to current fit for purpose review	Heu, C., et al. (2012). "Glyphosate-induced stiffening of HaCaT keratinocytes, a Peak Force Tapping study on living cells." <i>J Struct Biol</i> 178(1): 1-7.
Not Relevant to current fit for purpose review	Hill, H. F. (1962). "Nakajima's Glaucoma Operation." <i>American Journal of Ophthalmology</i> 53(2): 391-392.
Not Relevant to current fit for purpose review	Hohzaki, R. (2006). "Search allocation game." <i>European Journal of Operational Research</i> 172(1): 101-119.
Not Relevant to current fit for purpose review	Hokanson, R., et al. (2007). "Alteration of estrogen-regulated gene expression in human cells induced by the agricultural and horticultural herbicide glyphosate." <i>Hum Exp Toxicol</i> 26(9): 747-752.
Not Relevant to current fit for purpose review	Hothorn, L. A. and R. Oberdoerfer (2006). "Statistical analysis used in the nutritional assessment of novel food using the proof of safety." <i>Regulatory Toxicology and Pharmacology</i> 44(2): 125-135.
Not Relevant to current fit for purpose review	Hour, B. T., et al. (2012). "Herbicide Roundup Intoxication: Successful Treatment with Continuous Renal Replacement Therapy." <i>The American Journal of Medicine</i> 125(8): e1-e2.
Not Relevant to current fit for purpose review	Hsu, M.-H., et al. (2007). "2-(3-Fluorophenyl)-6-methoxyl-4-oxo-1,4-dihydroquinoline-3-carboxylic acid (YJC-1) induces mitotic phase arrest in A549 cells." <i>European Journal of Pharmacology</i> 559(1): 14-20.
Not Relevant to current fit for purpose review	Hu, B.-b., et al. (2014). "Energy efficiency of massive MIMO wireless communication systems with antenna selection." <i>The Journal of China Universities of Posts and Telecommunications</i> 21(<HT>6</HT>): 1-8.
Not Relevant to current fit for purpose review	Hu, T., et al. (2011). "Isolation and characterization of a rice glutathione S-transferase gene promoter regulated by herbicides and hormones." <i>Plant Cell Rep</i> 30(4): 539-549.
Not Relevant to current fit for purpose review	Huang, C. C., et al. (2004). "Development and application of a nested polymerase chain reaction method for the detection of genetically modified soybean in Chinese traditional fermented soy food-sufu." <i>Journal of Food and Drug Analysis</i> 12(3): 266-272.
Not Relevant to current fit for purpose review	Huang, J. L., et al. (2016). "Enhancement of the genotoxicity of benzo[a]pyrene by arecoline through suppression of DNA repair in HEp-2 cells." <i>Toxicology in Vitro</i> 33: 80-87.
Not Relevant to current fit for purpose review	Huang, Y.-T., et al. (2014). "The depletion of securin enhances butein-induced apoptosis and tumor inhibition in human colorectal cancer." <i>Chemico-Biological Interactions</i> 220: 41-50.

Not Relevant to current fit for purpose review	Hübner, P., et al. (1999). "Quantitative competitive PCR for the detection of genetically modified organisms in food." <i>Food Control</i> 10(6): 353-358.
Not Relevant to current fit for purpose review	Hubner, P., et al. (2001). "Validation of PCR methods for quantitation of genetically modified plants in food." <i>J AOAC Int</i> 84(6): 1855-1864.
Not Relevant to current fit for purpose review	Huisman, W., et al. (1997). "Costs of supply chains of <i>Miscanthus giganteus</i> ." <i>Industrial Crops and Products</i> 6(3-4): 353-366.
Not Relevant to current fit for purpose review	Hultberg, M. (2007). "Cysteine turnover in human cell lines is influenced by glyphosate." <i>Environmental Toxicology and Pharmacology</i> 24(1): 19-22.
Not Relevant to current fit for purpose review	Huynh, Q. K., et al. (1988). "Site-directed mutagenesis of <i>Petunia hybrida</i> 5-enolpyruvylshikimate-3-phosphate synthase: Lys-23 is essential for substrate binding." <i>J Biol Chem</i> 263(24): 11636-11639.
Not Relevant to current fit for purpose review	Jackson, E. (1933). "Progressive Ophthalmology." <i>American Journal of Ophthalmology</i> 16(2): 154-155.
Not Relevant to current fit for purpose review	Jacobelli, J., et al. (2004). "New views of the immunological synapse: variations in assembly and function." <i>Current Opinion in Immunology</i> 16(3): 345-352.
Not Relevant to current fit for purpose review	Jank, B. and A. Haslberger (2000). "Recombinant DNA insertion into plant retrotransposons." <i>Trends in Biotechnology</i> 18(8): 326-327.
Not Relevant to current fit for purpose review	Jankovic, J. (2008). "Parkinson's disease and movement disorders: moving forward." <i>The Lancet Neurology</i> 7(1): 9-11.
Not Relevant to current fit for purpose review	Jennings, J. C., et al. (2003). "Determining whether transgenic and endogenous plant DNA and transgenic protein are detectable in muscle from swine fed Roundup Ready soybean meal." <i>J Anim Sci</i> 81(6): 1447-1455.
Not Relevant to current fit for purpose review	Jiang, L. X., et al. (2013). "Glyphosate effects on the gene expression of the apical bud in soybean (<i>Glycine max</i>)." <i>Biochem Biophys Res Commun</i> 437(4): 544-549.
Not Relevant to current fit for purpose review	Johnson, K. A. (2013). "A century of enzyme kinetic analysis, 1913 to 2013." <i>FEBS Letters</i> 587(17): 2753-2766.
Not Relevant to current fit for purpose review	Jones, W. E. (1986). "Genetic recreation of wild horses." <i>Journal of Equine Veterinary Science</i> 6(5): 246-249.
Not Relevant to current fit for purpose review	Kang, M., et al. (1993). "Interactive partitioning criteria set method for multiple objective linear programming." <i>Computers & Operations Research</i> 20(4): 435-446.
Not Relevant to current fit for purpose review	Kang, S. (2014). "Research round-up." <i>The Lancet Psychiatry</i> 1(7): 502.
Not Relevant to current fit for purpose review	Kang, S. (2014). "Research round-up." <i>The Lancet Psychiatry</i> 1(4): 261.
Not Relevant to current fit for purpose review	Kang, S. (2015). "Research round-up." <i>The Lancet Psychiatry</i> 2(1): 17.
Not Relevant to current fit for purpose review	Kang, S. (2016). "Research round-up." <i>The Lancet Psychiatry</i> 3(4): 323.
Not Relevant to current fit for purpose review	Kang, Y., et al. (2011). "Knockout and pullout recombineering for naturally transformable <i>Burkholderia thailandensis</i> and <i>Burkholderia pseudomallei</i> ." <i>Nature Protocols</i> 6(8): 1085-1104.
Not Relevant to current fit for purpose review	Keenan, R. J., et al. (2005). "DNA shuffling as a tool for protein crystallization." <i>Proc Natl Acad Sci U S A</i> 102(25): 8887-8892.
Not Relevant to current fit for purpose review	Kelly, D. (2003). "Research Round-up." <i>European Journal of Oncology Nursing</i> 7(3): 210-212.
Not Relevant to current fit for purpose review	Kiernan, M. C. (2014). "ALS and neuromuscular disease: in search of the Holy Grail." <i>The Lancet Neurology</i> 13(1): 13-14.
Not Relevant to current fit for purpose review	Kim, H. J., et al. (2007). "Engineering and characterization of the isolated C-terminal domain of 5-enolpyruvylshikimate-3-phosphate (EPSP) synthase." <i>Journal of Microbiology and Biotechnology</i> 17(8): 1385-1389.
Not Relevant to current fit for purpose review	Kim, I., et al. (2012). "Cloud computing for comparative genomics with windows azure platform." <i>Evol Bioinform Online</i> 8: 527-534.
Not Relevant to current fit for purpose review	Kim, J.-H., et al. (2015). "Detection of eight genetically modified canola events using two event-specific pentaplex PCR systems." <i>Food Control</i> 51: 183-189.
Not Relevant to current fit for purpose review	Kim, J.-H., et al. (2015). "A simplified and accurate detection of the genetically modified wheat MON71800 with one calibrator plasmid." <i>Food Chemistry</i> 176: 1-6.

Not Relevant to current fit for purpose review	Kim, Y. H., et al. (2013). "Mixtures of glyphosate and surfactant TN20 accelerate cell death via mitochondrial damage-induced apoptosis and necrosis." <i>Toxicol In Vitro</i> 27(1): 191-197.
Not Relevant to current fit for purpose review	Kim, Y. H., et al. (2014). "Heart rate-corrected QT interval predicts mortality in glyphosate-surfactant herbicide-poisoned patients." <i>The American Journal of Emergency Medicine</i> 32(3): 203-207.
Not Relevant to current fit for purpose review	Kira, J.-i. (2008). "MS: prevention of neural damage by early intervention." <i>The Lancet Neurology</i> 7(1): 3-5.
Not Relevant to current fit for purpose review	Kirkham, F. (2004). "Paediatric neurology: genes and the environment." <i>The Lancet Neurology</i> 3(1): 18.
Not Relevant to current fit for purpose review	Klein, E. T. (1977). "J. Lyndon Carman (1905–1977)." <i>American Journal of Orthodontics</i> 72(2): 213-214.
Not Relevant to current fit for purpose review	Knopper, L. D. and D. R. S. Lean (2004). "Carcinogenic and genotoxic potential of turf pesticides commonly used on golf courses." <i>Journal of Toxicology and Environmental Health-Part B-Critical Reviews</i> 7(4): 267-279.
Not Relevant to current fit for purpose review	Kohn, P. M., et al. (1987). "Relationships between psychometric and experimental measures of arousability." <i>Personality and Individual Differences</i> 8(2): 225-231.
Not Relevant to current fit for purpose review	Kordowska, J., et al. (2006). "Phosphorylated l-caldesmon is involved in disassembly of actin stress fibers and postmitotic spreading." <i>Experimental Cell Research</i> 312(2): 95-110.
Not Relevant to current fit for purpose review	Kovalic, D., et al. (2012). "The Use of Next Generation Sequencing and Junction Sequence Analysis Bioinformatics to Achieve Molecular Characterization of Crops Improved Through Modern Biotechnology." <i>Plant Genome</i> 5(3): 149-163.
Not Relevant to current fit for purpose review	Kresge, K. J. (2005). "Women and HIV. Women's research roundup." <i>Beta</i> 17(4): 30-33.
Not Relevant to current fit for purpose review	Kristensen, T. (2004). "Lethal pictures." <i>Network Security</i> 2004(10): 19-20.
Not Relevant to current fit for purpose review	Kristensen, T. (2004). "Microsoft leaves Win2000, XPSP1 users in lurch." <i>Network Security</i> 2004(11): 16-17.
Not Relevant to current fit for purpose review	Kristensen, T. (2005). "More holes than a phishing net." <i>Network Security</i> 2005(1): 18-19.
Not Relevant to current fit for purpose review	Kudtarkar, P., et al. (2010). "Cost-effective cloud computing: a case study using the comparative genomics tool, roundup." <i>Evol Bioinform Online</i> 6: 197-203.
Not Relevant to current fit for purpose review	Kulesa, P. M., et al. (2008). "Neural crest invasion is a spatially-ordered progression into the head with higher cell proliferation at the migratory front as revealed by the photoactivatable protein, KikGR." <i>Developmental Biology</i> 316(2): 275-287.
Not Relevant to current fit for purpose review	Kuramochi, Y., et al. (2006). "Neuregulin activates erbB2-dependent src/FAK signaling and cytoskeletal remodeling in isolated adult rat cardiac myocytes." <i>Journal of Molecular and Cellular Cardiology</i> 41(2): 228-235.
Not Relevant to current fit for purpose review	Kuribara, H., et al. (2002). "Novel reference molecules for quantitation of genetically modified maize and soybean." <i>J AOAC Int</i> 85(5): 1077-1089.
Not Relevant to current fit for purpose review	Kutateladze, T., et al. (2007). "Detection of biotechnological crops in Georgia." <i>Journal of Biotechnology</i> 131(2, Supplement): S37-S38.
Not Relevant to current fit for purpose review	Kwon, K., et al. (2012). "3.103 PREDICTION AND CHARACTERIZATION OF ARABIDOPSIS THALIANA HOMOLOGS OF PARKINSON DISEASE-ASSOCIATED GENES BASED ON SEQUENCE SIMILARITIES AND MOLECULAR MODELING." <i>Parkinsonism & Related Disorders</i> 18, Supplement 2: S190.
Not Relevant to current fit for purpose review	La Mura, M., et al. (2011). "Application of QUIZ for GM quantification in food." <i>Food Chemistry</i> 125(4): 1340-1344.
Not Relevant to current fit for purpose review	Lam, S. H. F., et al. "Use of Intravenous Fat Emulsion in the Emergency Department for the Critically Ill Poisoned Patient." <i>The Journal of Emergency Medicine</i> .
Not Relevant to current fit for purpose review	Lamb, G. C., et al. (2000). "Prolonging the MGA-prostaglandin F2 α interval from 17 to 19 days in an estrus synchronization system for heifers." <i>Theriogenology</i> 53(3): 691-698.
Not Relevant to current fit for purpose review	Larkin, M. (2001). "Websites in brief." <i>The Lancet</i> 358(9284): 850.
Not Relevant to current fit for purpose review	Larkin, M. (2003). "Technology and Public Health." <i>The Lancet Infectious Diseases</i> 3(5): 314-315.

Not Relevant to current fit for purpose review	Larkin, M. (2004). "Sites accelerate global response to HIV." <i>The Lancet Infectious Diseases</i> 4(7): 472-473.
Not Relevant to current fit for purpose review	Larsen, R. J. and D. W. Baggs (1986). "Some psychophysical and personality correlates of the Strelau temperament inventory." <i>Personality and Individual Differences</i> 7(4): 561-565.
Not Relevant to current fit for purpose review	Larsen, R. J. and M. A. Zarate (1991). "Extending reducer/augmenter theory into the emotion domain: The role of affect in regulating stimulation level." <i>Personality and Individual Differences</i> 12(7): 713-723.
Not Relevant to current fit for purpose review	Lee, A. H. (2008). "A pilot intervention for pregnant women in Sichuan, China on passive smoking." <i>Patient Education and Counseling</i> 71(3): 396-401.
Not Relevant to current fit for purpose review	Lee, C.-H., et al. (2008). "The early prognostic factors of glyphosate-surfactant intoxication." <i>The American Journal of Emergency Medicine</i> 26(3): 275-281.
Not Relevant to current fit for purpose review	Lee, D., et al. (2010). "Quantitation using informative zeros (QUIZ): Application for GMO detection and quantification without recourse to certified reference material." <i>Food Chemistry</i> 118(4): 974-978.
Not Relevant to current fit for purpose review	Lee, E.-J., et al. (2004). "G2/M cell cycle arrest and induction of apoptosis by a stilbenoid, 3,4,5-trimethoxy-4'-bromo-cis-stilbene, in human lung cancer cells." <i>Life Sciences</i> 75(23): 2829-2839.
Not Relevant to current fit for purpose review	Lee, H. J., et al. (2010). "Cloning and Characterization of a 5-Enolpyruvyl Shikimate 3-Phosphate Synthase (EPSPS) Gene from Korean Lawn Grass (<i>Zoysia japonica</i>)." <i>Korean Journal of Horticultural Science & Technology</i> 28(4): 648-655.
Not Relevant to current fit for purpose review	Lee, S. R., et al. (2007). "Neuroprotective effects of kobophenol A against the withdrawal of tropic support, nitrosative stress, and mitochondrial damage in SH-SY5Y neuroblastoma cells." <i>Bioorganic & Medicinal Chemistry Letters</i> 17(7): 1879-1882.
Not Relevant to current fit for purpose review	Lees, K. R. (2009). "Stroke: success for extending acute treatment." <i>The Lancet Neurology</i> 8(1): 2-4.
Not Relevant to current fit for purpose review	Lerat, S., et al. (2005). "Real-time polymerase chain reaction quantification of the transgenes for roundup ready corn and roundup ready soybean in soil samples." <i>J Agric Food Chem</i> 53(5): 1337-1342.
Not Relevant to current fit for purpose review	Leva, A. (2015). "Computing systems and the network as a control education arena." <i>IFAC-PapersOnLine</i> 48(29): 265-276.
Not Relevant to current fit for purpose review	Levy, M. L. (2006). "What's in this issue." <i>Primary Care Respiratory Journal</i> 15(<HT>6</HT>): 321-322.
Not Relevant to current fit for purpose review	Li, G., et al. (2015). "A genetic algorithm-based decomposition approach to solve an integrated equipment-workforce-service planning problem." <i>Omega</i> 50: 1-17.
Not Relevant to current fit for purpose review	Li, H., et al. (2014). "Chaetoglobosins from <i>Chaetomium globosum</i> , an endophytic fungus in <i>Ginkgo biloba</i> , and their phytotoxic and cytotoxic activities." <i>J Agric Food Chem</i> 62(17): 3734-3741.
Not Relevant to current fit for purpose review	Li, L., et al. (2015). "A transcriptomic analysis for identifying the unintended effects of introducing a heterologous glyphosate-tolerant EPSP synthase into <i>Escherichia coli</i> ." <i>Molecular Biosystems</i> 11(3): 852-858.
Not Relevant to current fit for purpose review	Li, R. and R. S. El-Mallahk (2000). "A novel evidence of different mechanisms of lithium and valproate neuroprotective action on human SY5Y neuroblastoma cells: caspase-3 dependency." <i>Neuroscience Letters</i> 294(3): 147-150.
Not Relevant to current fit for purpose review	Lian, Y., et al. (2016). "Direct and simultaneous quantification of ATP, ADP and AMP by ¹ H and ³¹ P Nuclear Magnetic Resonance spectroscopy." <i>Talanta</i> 150: 485-492.
Not Relevant to current fit for purpose review	Liang, J., et al. (2009). "The Role of Seminal Vesicle Motion in Target Margin Assessment for Online Image-Guided Radiotherapy for Prostate Cancer." <i>International Journal of Radiation Oncology*Biophysics</i> 73(3): 935-943.
Not Relevant to current fit for purpose review	Liberatore, M., et al. (2010). "Forensic investigation of peer-to-peer file sharing networks." <i>Digital Investigation</i> 7, Supplement: S95-S103.
Not Relevant to current fit for purpose review	Lin, N. and V. F. Garry (2000). "In vitro studies of cellular and molecular developmental toxicity of adjuvants, herbicides, and fungicides commonly used in Red River Valley, Minnesota." <i>J Toxicol Environ Health A</i> 60(6): 423-439.
Not Relevant to current fit for purpose review	Lin, Y.-C., et al. (2011). "744 ZOLEDRONIC ACID INDUCES AUTOPHAGIC CELL DEATH IN HUMAN PROSTATE CANCER CELLS." <i>The Journal of Urology</i> 185(4, Supplement): e298-e299.
Not Relevant to current fit for purpose review	Lin, Y.-K., et al. (2013). "Multiple-objective heuristics for scheduling unrelated parallel machines." <i>European Journal of Operational Research</i> 227(2): 239-253.
Not Relevant to current fit for purpose review	Lipton, R. B. and M. E. Bigal (2005). "Headache: triumphs in translational research." <i>The Lancet Neurology</i> 4(1): 11-12.

Not Relevant to current fit for purpose review	Liu, M. (2008). "Stroke: encouragement and disappointment in clinical trials." <i>The Lancet Neurology</i> 7(1): 5-7.
Not Relevant to current fit for purpose review	Liu, S., et al. (2003). "A prospective study of the association between APOE genotype and the risk of myocardial infarction among apparently healthy men." <i>Atherosclerosis</i> 166(2): 323-329.
Not Relevant to current fit for purpose review	Lluis, M., et al. (2008). "Severe acute poisoning due to a glufosinate containing preparation without mitochondrial involvement." <i>Hum Exp Toxicol</i> 27(6): 519-524.
Not Relevant to current fit for purpose review	Lodge, G. M., et al. (1994). "EFFECTS OF GLYPHOSATE, FLUPROPANATE AND 2,2-DPA ON HYPARRHENIA-HIRTA (L) STAPF (COOLATAI GRASS)." <i>Australian Journal of Experimental Agriculture</i> 34(4): 479-485.
Not Relevant to current fit for purpose review	Lombardo, M. and G. Lombardo (2010). "Wave aberration of human eyes and new descriptors of image optical quality and visual performance." <i>Journal of Cataract & Refractive Surgery</i> 36(2): 313-331.
Not Relevant to current fit for purpose review	Loredano, P., et al. (2010). "CASCAT: The Power of The Combined Protein Engineering Approach: Evolution of A Glycine Oxidase for A Novel Mechanism of Glyphosate Tolerance." <i>Journal of Biotechnology</i> 150: S122-S123.
Not Relevant to current fit for purpose review	Lucentini, J. (2008). "Technology roundup." <i>Scientist</i> : 76-82.
Not Relevant to current fit for purpose review	Machado, A. F. L., et al. (2008). "Anatomical characterization of the leaf, stem and rhizome of <i>Digitaria insularis</i> ." <i>Planta Daninha</i> 26(1): 1-8.
Not Relevant to current fit for purpose review	Mahmood, A. (2009). "Renewables in Asia – a roundup." <i>Renewable Energy Focus</i> 9(7): 19.
Not Relevant to current fit for purpose review	Main, C. L., et al. (2004). "Sulfentrazone persistence in southern soils: Bioavailable concentration and effect on a rotational cotton crop." <i>Weed Technology</i> 18(2): 346-352.
Not Relevant to current fit for purpose review	Makowski, M., et al. (2009). "Oestrogen receptor alpha polymorphisms in women with breast cancer and their clinical significance." <i>Przegląd Menopauzalny</i> 8(1): 40-44.
Not Relevant to current fit for purpose review	Malatesta, M., et al. (2008). "Hepatoma tissue culture (HTC) cells as a model for investigating the effects of low concentrations of herbicide on cell structure and function." <i>Toxicol In Vitro</i> 22(8): 1853-1860.
Not Relevant to current fit for purpose review	Malecot, M., et al. (2013). "Specific proteomic response of <i>Unio pictorum</i> mussel to a mixture of glyphosate and microcystin-LR." <i>J Proteome Res</i> 12(11): 5281-5292.
Not Relevant to current fit for purpose review	Malhotra, R. C., et al. (2010). "Glyphosate–surfactant herbicide-induced reversible encephalopathy." <i>Journal of Clinical Neuroscience</i> 17(11): 1472-1473.
Not Relevant to current fit for purpose review	Mariana, A., et al. (2009). "The impact of simultaneous intoxication with agrochemicals on the antioxidant defense system in rat." <i>Pesticide Biochemistry and Physiology</i> 94(2-3): 93-99.
Not Relevant to current fit for purpose review	Marques, M. R., et al. (2007). "The inhibition of 5-enolpyruvylshikimate-3-phosphate synthase as a model for development of novel antimicrobials." <i>Current Drug Targets</i> 8(3): 445-457.
Not Relevant to current fit for purpose review	Marrelli, M., et al. (2013). "A comparative study of phytochemical composition of genetically and non-genetically modified soybean (<i>Glycine max</i> L.) and evaluation of antitumor activity." <i>Nat Prod Res</i> 27(6): 574-578.
Not Relevant to current fit for purpose review	Marriott, G. and J. W. Walker (1999). "Caged peptides and proteins: new probes to study polypeptide function in complex biological systems." <i>Trends in Plant Science</i> 4(8): 330-334.
Not Relevant to current fit for purpose review	Mastaglia, F. L. (2005). "Neuromuscular disorders: molecular and therapeutic insights." <i>The Lancet Neurology</i> 4(1): 6-7.
Not Relevant to current fit for purpose review	Mavunganidze, Z., et al. (2014). "The impact of tillage system and herbicides on weed density, diversity and yield of cotton (<i>Gossipium hirsutum</i> L.) and maize (<i>Zea mays</i> L.) under the smallholder sector." <i>Crop Protection</i> 58: 25-32.
Not Relevant to current fit for purpose review	Mayeux, R. and P. S. G. Hyslop (2008). "Alzheimer's disease: advances in trafficking." <i>The Lancet Neurology</i> 7(1): 2-3.
Not Relevant to current fit for purpose review	McAlister, B. G. and J. van Staden (1995). "Effect of artificially induced stress conditions on the growth of the medicinal plant <i>Hypoxis hemerocallidea</i> ." <i>South African Journal of Botany</i> 61(2): 85-89.
Not Relevant to current fit for purpose review	McCleneghan, J. S. (1994). "The 1993 newspaper science reporter: Contributing, creative, and responsible." <i>The Social Science Journal</i> 31(4): 467-477.
Not Relevant to current fit for purpose review	McDowell, L. M., et al. (2004). "Rotational-echo double-resonance NMR-restrained model of the ternary complex of 5-enolpyruvylshikimate-3-phosphate synthase." <i>J Biomol NMR</i> 28(1): 11-29.
Not Relevant to current fit for purpose review	McKeone, D. H. (1973). "Liquid crystals for display applications." <i>Optics & Laser Technology</i> 5(1): 39-40.

Not Relevant to current fit for purpose review	Meacham, M. (1976). "Development of school libraries around the world." <i>International Library Review</i> 8(4): 453-459.
Not Relevant to current fit for purpose review	Merlis, J. K. (1959). "Fundamentals of clinical neurophysiology: Paul O. Chatfield Charles C. Thomas, Springfield, Illinois, 1957, 392 pp. \$8.50." <i>Electroencephalography and Clinical Neurophysiology</i> 11(1): 192.
Not Relevant to current fit for purpose review	Mesnager, R., et al. (2013). "Ethoxylated adjuvants of glyphosate-based herbicides are active principles of human cell toxicity." <i>Toxicology</i> 313(2-3): 122-128.
Not Relevant to current fit for purpose review	Milán, M. and S. M. Cohen (1999). "Notch Signaling Is Not Sufficient to Define the Affinity Boundary between Dorsal and Ventral Compartments." <i>Molecular Cell</i> 4(<HT>6</HT>): 1073-1078.
Not Relevant to current fit for purpose review	Miller, J. W., et al. (1991). "Phthalocyanine Photodynamic Therapy of Experimental Iris Neovascularization." <i>Ophthalmology</i> 98(11): 1711-1719.
Not Relevant to current fit for purpose review	Miller, S. P. and D. M. Ferriero (2013). "Paediatric neurology: improved care of the developing brain." <i>The Lancet Neurology</i> 12(1): 16-18.
Not Relevant to current fit for purpose review	Milligan, A. L., et al. (2003). "A field assessment of the role of selective herbicides in the restoration of British moorland dominated by <i>Molinia</i> ." <i>Biological Conservation</i> 109(3): 369-379.
Not Relevant to current fit for purpose review	Mink, P. J., et al. (2011). "Epidemiologic studies of glyphosate and non-cancer health outcomes: a review." <i>Regul Toxicol Pharmacol</i> 61(2): 172-184.
Not Relevant to current fit for purpose review	Minshull, J., et al. (2005). "Predicting enzyme function from protein sequence." <i>Current Opinion in Chemical Biology</i> 9(2): 202-209.
Not Relevant to current fit for purpose review	Mo, Q. and X. Zhuang (2012). "Matrix splitting with symmetry and dyadic framelet filter banks over algebraic number fields." <i>Linear Algebra and its Applications</i> 437(10): 2650-2679.
Not Relevant to current fit for purpose review	Monsen, E. R. (1998). "ADA'S Nutrition Campaigns Link Dietitians and the Public's Health." <i>Journal of the American Dietetic Association</i> 98(7): 746.
Not Relevant to current fit for purpose review	Monsen, E. R. (1998). "Dietetics Professionals Fight Obesity." <i>Journal of the American Dietetic Association</i> 98(10): 1098.
Not Relevant to current fit for purpose review	Monsen, E. R. (1999). "The Past Dietetics Century and the Future Dietetics Century." <i>Journal of the American Dietetic Association</i> 99(12): 1498.
Not Relevant to current fit for purpose review	Monsen, E. R. (1999). "What's Fresh in this Month's Journal?" <i>Journal of the American Dietetic Association</i> 99(3): 274.
Not Relevant to current fit for purpose review	Moreano, F., et al. (2006). "Ligation-dependent probe amplification for the simultaneous event-specific detection and relative quantification of DNA from two genetically modified organisms." <i>European Food Research and Technology</i> 222(5-6): 479-485.
Not Relevant to current fit for purpose review	Morgensztern, D. and R. Govindan (2010). "Best of the month: a roundup of articles published in recent months." <i>J Thorac Oncol</i> 5(8): 1305-1307.
Not Relevant to current fit for purpose review	Morgensztern, D. and R. Govindan (2011). "A Roundup of Recently Published Articles Relevant to Thoracic Oncology." <i>Journal of Thoracic Oncology</i> <HT>6</HT>(7): 1295-1297.
Not Relevant to current fit for purpose review	Morgensztern, D. and R. Govindan (2011). "A roundup of recently published articles relevant to thoracic oncology." <i>J Thorac Oncol</i> 6(7): 1295-1297.
Not Relevant to current fit for purpose review	Morris, R. T., et al. (2011). "RiceRBP: A database of experimentally identified RNA-binding proteins in <i>Oryza sativa</i> L." <i>Plant Science</i> 180(2): 204-211.
Not Relevant to current fit for purpose review	Muntoni, F. and J. H. Cross (2015). "Paediatric neurology: from molecular mechanisms to targeted treatments." <i>The Lancet Neurology</i> 14(1): 16-18.
Not Relevant to current fit for purpose review	Mylynen, P. and K. Vähäkangas (2013). "Placental transfer and metabolism: An overview of the experimental models utilizing human placental tissue." <i>Toxicology in Vitro</i> 27(1): 507-512.
Not Relevant to current fit for purpose review	Nam, H.-J., et al. (2008). "The ERK-RSK1 activation by growth factors at G2 phase delays cell cycle progression and reduces mitotic aberrations." <i>Cellular Signalling</i> 20(7): 1349-1358.
Not Relevant to current fit for purpose review	Naydenova, E., et al. (2007). "Synthesis, cytotoxicity and clastogenicity of novel alpha-aminophosphonic acids." <i>Amino Acids</i> 33(4): 695-702.
Not Relevant to current fit for purpose review	Naydenova, E. D., et al. (2008). "Novel N-(phosphonomethyl) glycine derivatives: Design, characterization and biological activity." <i>Eur J Med Chem</i> 43(6): 1199-1205.
Not Relevant to current fit for purpose review	Neumann, G., et al. (2011). "Assessment of the genetic stability of GMOs with a detailed examination of MON810 using Scorpion probes." <i>European Food Research and Technology</i> 233(1): 19-30.
Not Relevant to current fit for purpose review	Newton, F. H. (1962). "Appraising New Techniques." <i>American Journal of Ophthalmology</i> 53(2): 392.

Not Relevant to current fit for purpose review	Norrving, B. (2006). "Stroke: the catch-up of underinvestigated topics." <i>The Lancet Neurology</i> 5(1): 10-11.
Not Relevant to current fit for purpose review	Novis, P. M., et al. (2009). "Identification and Characterization of Freshwater Algae from a Pollution Gradient Using rbcL Sequencing and Toxicity Testing." <i>Arch Environ Contam Toxicol</i> 57(3): 504-514.
Not Relevant to current fit for purpose review	Ocaña, K. A. C. S., et al. (2013). "Designing a parallel cloud based comparative genomics workflow to improve phylogenetic analyses." <i>Future Generation Computer Systems</i> 29(8): 2205-2219.
Not Relevant to current fit for purpose review	Oeding, L. and G. Ottaviani (2013). "Eigenvectors of tensors and algorithms for Waring decomposition." <i>Journal of Symbolic Computation</i> 54: 9-35.
Not Relevant to current fit for purpose review	Oguchi, T., et al. (2008). "Development of event-specific quantitation method for GA21 maize, which is a gm event without CaMV35S promoter." <i>Shokuhin Eiseigaku Zasshi</i> 49(1): 16-22.
Not Relevant to current fit for purpose review	Okamoto, T. (2015). "SecondDEP: Resilient Computing that Prevents Shellcode Execution in Cyber-Attacks." <i>Procedia Computer Science</i> 60: 691-699.
Not Relevant to current fit for purpose review	Olanow, C. W. (2006). "Movement disorders: a step in the right direction." <i>The Lancet Neurology</i> 5(1): 3-5.
Not Relevant to current fit for purpose review	Omran, D. (1993). "Doesn't quite live up to expectations: Enterprise-Wide Computing: How to Implement and Manage LANs by Thomas W. Madron. Published by John Wiley, 1991, £25.95, 372 pp." <i>Computer Communications</i> 16(7): 449-450.
Not Relevant to current fit for purpose review	Onori, R., et al. (2013). "Traceability of genetically modified Roundup Ready soybean: A case study on sampling and analytical uncertainty along processing chain." <i>Food Control</i> 34(2): 494-501.
Not Relevant to current fit for purpose review	O'Rourke, K. (2004). "Genome roundup." <i>J Am Vet Med Assoc</i> 224(4): 491-492.
Not Relevant to current fit for purpose review	Ortiz, S., et al. (2006). "Acute cytotoxicity in mammal cells exposed in vitro to a glyphosate-based formulation." <i>Environ Mol Mutagen</i> 47(6): 473-473.
Not Relevant to current fit for purpose review	Park, S., et al. (2015). "The effects of nonyl phenoxypolyethoxyl ethanol on cell damage pathway gene expression in SK-N-SH cells." <i>Korean Journal of Internal Medicine</i> 30(6): 873-883.
Not Relevant to current fit for purpose review	Pasqualone, A., et al. (2007). "Detection of soft wheat in semolina and durum wheat bread by analysis of DNA microsatellites." <i>J Agric Food Chem</i> 55(9): 3312-3318.
Not Relevant to current fit for purpose review	Patterson, P. (1980). "Nurses hold own in federal battle for education funds." <i>AORN Journal</i> 32(1): 116-118.
Not Relevant to current fit for purpose review	Patton, M. L. and R. F. Grove (1992). "Statolith hair movements and the regulation of tonic gravity reflexes in the lobster, <i>Homarus americanus</i> ." <i>Comparative Biochemistry and Physiology Part A: Physiology</i> 101(2): 259-268.
Not Relevant to current fit for purpose review	Perisse, P., et al. (2011). "Seed and seedling morphology of <i>Dicliptera squarrosa</i> Nees (Acanthaceae) as a character identification source, and its relationship with survival structures." <i>Phyton-International Journal of Experimental Botany</i> 80: 73-78.
Not Relevant to current fit for purpose review	Perret, R. (2012). "Wanted dead or alive? Western genre items in the 21st century United States library." <i>Library Collections, Acquisitions, and Technical Services</i> 36(1-2): 39-52.
Not Relevant to current fit for purpose review	Perumal, L., et al. (2016). "Analysis of thin plates with holes by using exact geometrical representation within XFEM." <i>Journal of Advanced Research</i> 7(3): 445-452.
Not Relevant to current fit for purpose review	Pervaiz, S. (2013). "Conference roundup MAC 2011." <i>Mitochondrion</i> 13(3): 153-154.
Not Relevant to current fit for purpose review	Petersen, R. C. (2010). "Alzheimer's disease: progress in prediction." <i>The Lancet Neurology</i> 9(1): 4-5.
Not Relevant to current fit for purpose review	Pörtl, D., et al. (2012). "3.104 CELL-DEATH PATHWAYS DOWNSTREAM OF MITOCHONDRIA IN HUMAN DOPAMINERGIC NEURON DEGENERATION." <i>Parkinsonism & Related Disorders</i> 18, Supplement 2: S190.
Not Relevant to current fit for purpose review	Posch, K. C. and R. Posch (1990). "Approaching encryption at ISDN speed using partial parallel modulus multiplication." <i>Microprocessing and Microprogramming</i> 29(3): 177-184.
Not Relevant to current fit for purpose review	Potrebic, O., et al. (2009). "Acute glyphosate-surfactant poisoning with neurological sequels and fatal outcome." <i>Vojnosanitetski Pregled</i> 66(9): 758-762.
Not Relevant to current fit for purpose review	Powell, S. L. (2000). "Herbal Supplements: What Works, What Doesn't. 1999. From National Health Video, Inc. 12021 Wilshire Blvd., Suite 550, Los Angeles, CA 90025, 1-800-543-6803, videotape, 15 minutes, \$89.00." <i>Journal of Nutrition Education</i> 32(5): 292-293.
Not Relevant to current fit for purpose review	Printz, C. (2013). "A roundup of news and information from our community." <i>Clin Transl Sci</i> 6(2): 83-84.

Not Relevant to current fit for purpose review	Printz, C. (2014). "In the news: a roundup of news and information from our community." Clin Transl Sci 7(4): 287-288.
Not Relevant to current fit for purpose review	Printz, C. (2015). "In the news: a roundup of news and information from our community." Clin Transl Sci 8(1): 1-2.
Not Relevant to current fit for purpose review	Qin, F., et al. (2012). "Composition of Transgenic Soybean Seeds with Higher gamma-Linolenic Acid Content Is Equivalent to That of Conventional Control." J Agric Food Chem 60(9): 2200-2204.
Not Relevant to current fit for purpose review	Radenski, A. and L. Ehwerhemuepha (2014). "Speeding-up codon analysis on the cloud with local MapReduce aggregation." Information Sciences 263: 175-185.
Not Relevant to current fit for purpose review	Radtke, P. J., et al. (2010). "An evaluation of overhead laser scanning to estimate herbage removals in pasture quadrats." Agricultural and Forest Meteorology 150(12): 1523-1528.
Not Relevant to current fit for purpose review	Rao, C. D., et al. (2015). "An enzyme-linked immuno focus assay for rapid detection and enumeration, and a newborn mouse model for human non-polio enteroviruses associated with acute diarrhea." Journal of Virological Methods 224: 47-52.
Not Relevant to current fit for purpose review	Rappold, P., et al. (2012). "3.106 PARAQUAT NEUROTOXICITY IS MEDIATED BY THE DOPAMINE TRANSPORTER AND ORGANIC CATION TRANSPORTER-3." Parkinsonism & Related Disorders 18, Supplement 2: S190-S191.
Not Relevant to current fit for purpose review	Rau, F. J. (1995). "Video review roundup." Adolescent and Pediatric Gynecology 8(3): 165.
Not Relevant to current fit for purpose review	Rau, F. J. (1997). "Comprehensive Review of Colposcopy 1996." Journal of Pediatric and Adolescent Gynecology 10(1): 49.
Not Relevant to current fit for purpose review	Rau, F. J. and K. M. Mulchahey (1997). "Milner-Fenwick Video Counseling Library: Colposcopy and Treatment of Dysplasia." Journal of Pediatric and Adolescent Gynecology 10(2): 103-104.
Not Relevant to current fit for purpose review	Rau, F. J. and E. E. Yordan (1996). "Sexual Assault: The Collection of Evidence for Forensic Analysis." Journal of Pediatric and Adolescent Gynecology 9(3): 169.
Not Relevant to current fit for purpose review	Reeve, J. C. (1995). "Core Textbook of Respiratory Care Practice." Physiotherapy 81(5): 301.
Not Relevant to current fit for purpose review	Reis, L. F., et al. (2006). "GMOs: building the future on the basis of past experience." An Acad Bras Cienc 78(4): 667-686.
Not Relevant to current fit for purpose review	Richard, S., et al. (2005). "Differential effects of glyphosate and roundup on human placental cells and aromatase." Environ Health Perspect 113(6): 716-720.
Not Relevant to current fit for purpose review	Richter, L. (1979). "Summary and guideline to the status reports." Euromicro Newsletter 5(3): 111-112.
Not Relevant to current fit for purpose review	Ring, M. (2007). "Driven Out by Hitler, a Dental Historian Enriches America: The Story of Curt Proskauer." Alpha Omegan 100(1): 25-29.
Not Relevant to current fit for purpose review	Robinson, D. J. (1995). "Study Package." Physiotherapy 81(5): 302.
Not Relevant to current fit for purpose review	Rogoli, R. P., et al. (2008). "Resposta de plantas de beterraba (Beta vulgaris) e de cenoura (Daucus carota) à deriva simulada de glyphosate e clomazone." Planta Daninha 26(2): 451-456.
Not Relevant to current fit for purpose review	Rosculete, E., et al. (2015). "Detection and quantification of genetically modified soybean in food and feed traded in the Romanian market." Journal of Biotechnology 208, Supplement: S73.
Not Relevant to current fit for purpose review	Roskelley, C. D., et al. (1995). "A hierarchy of ECM-mediated signalling regulates tissue-specific gene expression." Current Opinion in Cell Biology 7(5): 736-747.
Not Relevant to current fit for purpose review	Rouquié, D., et al. (2010). "Investigation of endogenous soybean food allergens by using a 2-dimensional gel electrophoresis approach." Regulatory Toxicology and Pharmacology 58(3, Supplement): S47-S53.
Not Relevant to current fit for purpose review	Ryd, W. and B. Hagmar (1977). "In vitro Effects of Cytochalasin B on TA3 Tumor Cells." Beiträge zur Pathologie 161(2): 131-141.
Not Relevant to current fit for purpose review	Sakamoto, Y., et al. (2008). "A 104-week feeding study of genetically modified soybeans in F344 rats." Journal of the Food Hygienic Society of Japan 49(4): 272-282.
Not Relevant to current fit for purpose review	Sakr, J., et al. (2014). "First comprehensive GMOs testing in Lebanon: Screening, identification and quantification of GM soybean imports." Food Control 36(1): 146-152.
Not Relevant to current fit for purpose review	Sanfilippo, J. S. (1998). ""Get A Life"-My Old Kentucky Home." Journal of Pediatric and Adolescent Gynecology 11(1): 1.
Not Relevant to current fit for purpose review	Sarraj, A. and J. C. Grotta (2014). "Stroke: new horizons in treatment." The Lancet Neurology 13(1): 2-3.

Not Relevant to current fit for purpose review	Sauer, N. J., et al. (2016). "Oligonucleotide-mediated genome editing provides precision and function to engineered nucleases and antibiotics in plants." <i>Plant Physiol.</i>
Not Relevant to current fit for purpose review	Sbruzzi, F. A., et al. (2013). "Transgenic and conventional Brazilian soybeans don't cause or prevent preneoplastic colon lesions or oxidative stress in a 90-day in vivo study." <i>Revista De Nutricao-Brazilian Journal of Nutrition</i> 26(4): 443-453.
Not Relevant to current fit for purpose review	Schapira, A. H. V. (2010). "Movement disorders: advances in cause and treatment." <i>The Lancet Neurology</i> 9(1): 6-7.
Not Relevant to current fit for purpose review	Scherlen, A. (2008). "Local to Global: The Importance of State-Level Journals to Library Literature." <i>Serials Review</i> 34(2): 129-136.
Not Relevant to current fit for purpose review	Schoenen, J. and G. Coppola (2010). "Headache: spreading from molecules to patients." <i>The Lancet Neurology</i> 9(1): 11-12.
Not Relevant to current fit for purpose review	Scotter, E. L. and C. E. Shaw (2013). "Neuromuscular disease: new insights and avenues for therapy." <i>The Lancet Neurology</i> 12(1): 13-15.
Not Relevant to current fit for purpose review	Selvapandiyan, A., et al. (1996). "Evidence for the shikimate-3-phosphate interacting site in the N-terminal domain of 5-enolpyruvyl shikimate-3-phosphate synthase of <i>Bacillus subtilis</i> ." <i>Biochemistry and Molecular Biology International</i> 40(3): 603-610.
Not Relevant to current fit for purpose review	Selvapandiyan, A. and R. K. Bhatnagar (1994). "CLONING OF GENES ENCODING FOR C-P LYASE FROM PSEUDOMONAS ISOLATES PG2982 AND GLC11 - IDENTIFICATION OF A CRYPTIC ALLELE ON THE CHROMOSOME OF PSEUDOMONAS-AERUGINOSA." <i>Current Microbiology</i> 29(5): 255-261.
Not Relevant to current fit for purpose review	Selvapandiyan, A., et al. (1995). "Point mutation of a conserved arginine (104) to lysine introduces hypersensitivity to inhibition by glyphosate in the 5-enolpyruvylshikimate-3-phosphate synthase of <i>Bacillus subtilis</i> ." <i>FEBS Lett</i> 374(2): 253-256.
Not Relevant to current fit for purpose review	Shi, G., et al. (2011). "MultiMSOAR 2.0: an accurate tool to identify ortholog groups among multiple genomes." <i>PLoS One</i> 6(6): e20892.
Not Relevant to current fit for purpose review	Shields, R. and K. Pollock (1974). "The adhesion of BHK and PyBHK cells to the substratum." <i>Cell</i> 3(1): 31-38.
Not Relevant to current fit for purpose review	Shuttleworth, W. A. and J. N. Evans (1994). "Site-directed mutagenesis and NMR studies of histidine-385 mutants of 5-enolpyruvylshikimate-3-phosphate synthase." <i>Biochemistry</i> 33(23): 7062-7068.
Not Relevant to current fit for purpose review	Shuttleworth, W. A. and J. N. Evans (1996). "The H385N mutant of 5-enolpyruvylshikimate-3-phosphate synthase: kinetics, fluorescence, and nuclear magnetic resonance studies." <i>Arch Biochem Biophys</i> 334(1): 37-42.
Not Relevant to current fit for purpose review	Shuttleworth, W. A. and J. N. S. Evans (1994). "SITE-DIRECTED MUTAGENESIS AND NMR-STUDIES OF HISTIDINE-385 MUTANTS OF 5-ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE SYNTHASE." <i>Biochemistry</i> 33(23): 7062-7068.
Not Relevant to current fit for purpose review	Shuttleworth, W. A. and J. N. S. Evans (1996). "The H385N Mutant of 5-Enolpyruvylshikimate-3-phosphate Synthase: Kinetics, Fluorescence, and Nuclear Magnetic Resonance Studies." <i>Archives of Biochemistry and Biophysics</i> 334(1): 37-42.
Not Relevant to current fit for purpose review	Shuttleworth, W. A., et al. (1992). "Over-production of 5-enolpyruvylshikimate-3-phosphate synthase in <i>Escherichia coli</i> : use of the T7 promoter." <i>Protein Eng</i> 5(5): 461-466.
Not Relevant to current fit for purpose review	Shuttleworth, W. A., et al. (1999). "Site-directed mutagenesis of putative active site residues of 5-enolpyruvylshikimate-3-phosphate synthase." <i>Biochemistry</i> 38(1): 296-302.
Not Relevant to current fit for purpose review	Sienkiewicz, Z. (2007). "Rapporteur report: Roundup, discussion and recommendations." <i>Progress in Biophysics and Molecular Biology</i> 93(1-3): 414-420.
Not Relevant to current fit for purpose review	Sikorski, J. A. and K. J. Gruys (1997). "Understanding Glyphosate's molecular mode of action with EPSP synthase: Evidence favoring an allosteric inhibitor model." <i>Accounts of Chemical Research</i> 30(1): 2-8.
Not Relevant to current fit for purpose review	Simonetti, E., et al. (2015). "An Interlaboratory Comparative Study on the Quantitative Determination of Glyphosate at Low Levels in Wheat Flour." <i>J AOAC Int</i> 98(6): 1760-1768.
Not Relevant to current fit for purpose review	Singh, N. and A. Srivastava (2014). "Biomonitoring of Genotoxic Effect of Glyphosate and Pendimethalin in <i>Vigna mungo</i> Populations." <i>Cytologia</i> 79(2): 173-180.
Not Relevant to current fit for purpose review	Singh, S. K. and S. Kumar (2011). "Novel adaptive color space transform and application to image compression." <i>Signal Processing: Image Communication</i> 26(10): 662-672.
Not Relevant to current fit for purpose review	Sinha, R. P., et al. (2003). "Wavelength-dependent induction of a mycosporine-like amino acid in a rice-field cyanobacterium, <i>Nostoc commune</i> : role of inhibitors and salt stress." <i>Photochemical & Photobiological Sciences</i> 2(2): 171-176.

Not Relevant to current fit for purpose review	Sliseris, J., et al. (2013). "Optimal Design of GFRP-Plywood Variable Stiffness Plate." <i>Procedia Engineering</i> 57: 1060-1069.
Not Relevant to current fit for purpose review	Smith, S. O. (1993). "Magic angle spinning NMR methods for internuclear distance measurements." <i>Current Opinion in Structural Biology</i> 3(5): 755-759.
Not Relevant to current fit for purpose review	Smitt, P. S. (2004). "Neuro-oncology: diagnosis in the spotlight." <i>The Lancet Neurology</i> 3(1): 14.
Not Relevant to current fit for purpose review	Somasiri, A., et al. (2000). "Phosphatidylinositol 3-Kinase is required for adherens junction-dependent mammary epithelial cell spheroid formation." <i>Differentiation</i> 66(2-3): 116-125.
Not Relevant to current fit for purpose review	Sorensen, P. S. (2005). "Multiple sclerosis: pathophysiology revisited." <i>The Lancet Neurology</i> 4(1): 9-10.
Not Relevant to current fit for purpose review	Spaeth, S. C. and T. R. Sinclair (1983). "Variation in nitrogen accumulation and distribution among soybean cultivars." <i>Field Crops Research</i> 7: 1-12.
Not Relevant to current fit for purpose review	Spector, L. S. (1990). "Not-so-open skies." <i>Space Policy</i> <HT>6</HT>(1): 9-18.
Not Relevant to current fit for purpose review	Sperling, R. A. and K. A. Johnson (2012). "Dementia: new criteria but no new treatments." <i>The Lancet Neurology</i> 11(1): 4-5.
Not Relevant to current fit for purpose review	Sroka, J., et al. (2002). "Phenotype modulation in non-adherent and adherent sublines of Walker carcinosarcoma cells: the role of cell-substratum contacts and microtubules in controlling cell shape, locomotion and cytoskeletal structure." <i>The International Journal of Biochemistry & Cell Biology</i> 34(7): 882-899.
Not Relevant to current fit for purpose review	Stanton, T. L., et al. (1990). "The Effects of Wet on Growing Heifer Brewers Grain Performance." <i>The Professional Animal Scientist</i> <HT>6</HT>(2): 31-34.
Not Relevant to current fit for purpose review	Stauffer, M. E., et al. (2001). "Shikimate-3-phosphate binds to the isolated N-terminal domain of 5-enolpyruvylshikimate-3-phosphate synthase." <i>Biochemistry</i> 40(13): 3951-3957.
Not Relevant to current fit for purpose review	Stobiecka, M., et al. (2007). "Piezoelectric sensor for determination of genetically modified soybean roundup ready((R)) in samples not amplified by PCR." <i>Sensors</i> 7(8): 1462-1479.
Not Relevant to current fit for purpose review	Stoessl, A. J. (2011). "Movement disorders: new insights into Parkinson's disease." <i>The Lancet Neurology</i> 10(1): 5-7.
Not Relevant to current fit for purpose review	Stovner, L. J. (2008). "Headache: new concepts, models, and treatments." <i>The Lancet Neurology</i> 7(1): 11-12.
Not Relevant to current fit for purpose review	Struyker-Boudier, H. A. J. (1983). "Clinical hypertension and hypotension." <i>Trends in Pharmacological Sciences</i> 4: 359.
Not Relevant to current fit for purpose review	Subramanian, J., et al. (2012). "A roundup of articles published in recent months." <i>J Thorac Oncol</i> 7(3): 626-628.
Not Relevant to current fit for purpose review	Sutou, S. and H. Shindo (1975). "Concanavalin a induces endoreduplication in cultured mammalian cells." <i>Biochemical and Biophysical Research Communications</i> 67(1): 79-84.
Not Relevant to current fit for purpose review	Suzuki, N., et al. (1996). "Purification and characterization of a cytosolic isozyme of 3-Deoxy-D-arabino-heptulosonate 7-phosphate synthase from cultured carrot cells." <i>Journal of Plant Physiology</i> 149(1-2): 19-22.
Not Relevant to current fit for purpose review	Swaney, S. (1995). "Spasmodic Torticollis." <i>Physiotherapy</i> 81(5): 301.
Not Relevant to current fit for purpose review	Tai, A., et al. (2008). "Estimate of Radiobiologic Parameters From Clinical Data for Biologically Based Treatment Planning for Liver Irradiation." <i>International Journal of Radiation Oncology*Biophysics</i> 70(3): 900-907.
Not Relevant to current fit for purpose review	Tai, H.-K., et al. (2011). "A one-time inducible transposon for terminating selectable markers in transgenic plants." <i>Botanical Studies</i> 52(4): 375-381.
Not Relevant to current fit for purpose review	Takabatake, R., et al. (2010). "Development and evaluation of event-specific quantitative pcr method for genetically modified soybean MON89788." <i>Journal of Biotechnology</i> 150, Supplement: 482-483.
Not Relevant to current fit for purpose review	Talbot, K. (2007). "Neuromuscular disorders: therapeutic advances." <i>The Lancet Neurology</i> <HT>6</HT>(1): 18-19.
Not Relevant to current fit for purpose review	Tan, A. and G. D. Badhwar (1997). "Detecting the dynamical state of the atmosphere from the orbital decay of the ODERACS spheres." <i>Journal of Atmospheric and Solar-Terrestrial Physics</i> 59(4): 431-437.
Not Relevant to current fit for purpose review	Tan, J. H., et al. (2013). "An interactive lung field segmentation scheme with automated capability." <i>Digital Signal Processing</i> 23(3): 1022-1031.

Not Relevant to current fit for purpose review	Tan, S., et al. (2006). "Herbicide inhibitors of amino acid biosynthesis and herbicide-tolerant crops." <i>Amino Acids</i> 30(2): 195-204.
Not Relevant to current fit for purpose review	Tardieu, M. (2010). "Paediatric neurology: brain development at an interface between genetics, the environment, and the immune system." <i>The Lancet Neurology</i> 9(1): 13-14.
Not Relevant to current fit for purpose review	Taverniers, I., et al. (2004). "Cloned plasmid DNA fragments as calibrators for controlling GMOs: different real-time duplex quantitative PCR methods." <i>Anal Bioanal Chem</i> 378(5): 1198-1207.
Not Relevant to current fit for purpose review	Thakur, K. T., et al. (2015). "CNS infections in 2014: guns, germs, and will." <i>The Lancet Neurology</i> 14(1): 20-22.
Not Relevant to current fit for purpose review	Thongprakaisang, S., et al. (2013). "Glyphosate induces human breast cancer cells growth via estrogen receptors." <i>Food Chem Toxicol</i> 59: 129-136.
Not Relevant to current fit for purpose review	Tong, X. H., et al. (2009). "Physiological and molecular mechanisms of glyphosate tolerance in an in vitro selected cotton mutant." <i>Pesticide Biochemistry and Physiology</i> 94(2-3): 100-106.
Not Relevant to current fit for purpose review	Tonkin, J. (1991). "South Australia CACCN." <i>Confederation of Australian Critical Care Nurses Journal</i> 4(3): 6.
Not Relevant to current fit for purpose review	Toyota, A., et al. (2006). "Quantification of genetically modified soybeans using a combination of a capillary-type real-time PCR system and a plasmid reference standard." <i>Biosci Biotechnol Biochem</i> 70(4): 821-827.
Not Relevant to current fit for purpose review	Trojano, M. and C. Tortorella (2015). "MS and related disorders: looking for markers of phenotypes." <i>The Lancet Neurology</i> 14(1): 11-13.
Not Relevant to current fit for purpose review	Trusler, C. S., et al. (2007). "Italian ryegrass (<i>Lolium multiflorum</i>) management options in winter wheat in Oklahoma." <i>Weed Technology</i> 21(1): 151-158.
Not Relevant to current fit for purpose review	Truta, E., et al. (2011). "Evaluation of Roundup-induced toxicity on genetic material and on length growth of barley seedlings." <i>Acta Biol Hung</i> 62(3): 290-301.
Not Relevant to current fit for purpose review	Turk, J. V. and D. L. Snedeker (1986). "Intro public relations texts: A round-up review." <i>Public Relations Review</i> 12(4): 48-55.
Not Relevant to current fit for purpose review	Tyshko, N. V., et al. (2008). "Medical and biological safety assessment of genetically modified maize event MON 88017. Report 2. Genotoxicologic, immunologic and allergologic examinations." <i>Voprosy pitaniya</i> 77(5): 13-17.
Not Relevant to current fit for purpose review	Ugarte, R. (2014). "Interaction between glyphosate and mitochondrial succinate dehydrogenase." <i>Computational and Theoretical Chemistry</i> 1043: 54-63.
Not Relevant to current fit for purpose review	Underwood, M. (1991). "Western Australia ASCCN." <i>Confederation of Australian Critical Care Nurses Journal</i> 4(3): 6.
Not Relevant to current fit for purpose review	Uppal, R. K. and M. J. Gooding (2013). "Gibberellin-responsive and -insensitive dwarfing alleles on wheat performance in contrasting tillage systems." <i>Field Crops Research</i> 141: 55-62.
Not Relevant to current fit for purpose review	Urda, A., et al. (2012). "Final voluntary assessment for Traumatology and Orthopaedic Surgery medical residents: A report on the results and a look at the future." <i>Revista Española de Cirugía Ortopédica y Traumatología (English Edition)</i> 56(3): 188-196.
Not Relevant to current fit for purpose review	Valencia, E., et al. (1999). "Yield and botanical composition of rhizoma peanut-grass swards treated with herbicides." <i>Agronomy Journal</i> 91(6): 956-961.
Not Relevant to current fit for purpose review	Van Caneghem, T. (2002). "EARNINGS MANAGEMENT INDUCED BY COGNITIVE REFERENCE POINTS." <i>The British Accounting Review</i> 34(2): 167-178.
Not Relevant to current fit for purpose review	van Duijn, G., et al. (1999). "Detection methods for genetically modified crops." <i>Food Control</i> 10(6): 375-378.
Not Relevant to current fit for purpose review	van Haselen, R. and R. Jütte (2013). "The placebo effect and its ramifications for clinical practice and research. Villa La Collina at Lake Como, Italy, 4-6 May 2012." <i>Complementary Therapies in Medicine</i> 21(2): 85-93.
Not Relevant to current fit for purpose review	Van Kampen, J. and J. D. Hundleby (1994). "An investigation of the associations between augmenting-reducing, reactivity, sensation seeking and body image." <i>Personality and Individual Differences</i> 16(3): 373-377.
Not Relevant to current fit for purpose review	Van Loon, A. J. (1991). "Glaciotectionic landforms and structures: J.S. Aber, D.G. Croot and M.M. Fenton. Kluwer Academic Publishers Group, Dordrecht, 1989, ix + 200 pp., Dfl.155.-, US\$79.-, £49.-, ISBN 0-7923-0100-5 (hardcover)." <i>Sedimentary Geology</i> 75(1-2): 164-165.
Not Relevant to current fit for purpose review	Venancio, V. P., et al. (2012). "Conventional (MG-BR46 Conquista) and Transgenic (BRS Valiosa RR) Soybeans Have No Mutagenic Effects and May Protect Against Induced-DNA Damage In Vivo." <i>Nutrition and Cancer-an International Journal</i> 64(5): 725-731.

Not Relevant to current fit for purpose review	Vyn, T. J., et al. (2002). "Corn response to potassium placement in conservation tillage." <i>Soil and Tillage Research</i> 67(2): 159-169.
Not Relevant to current fit for purpose review	Walker, D. C., et al. (2004). "The epitheliome: agent-based modelling of the social behaviour of cells." <i>Biosystems</i> 76(1-3): 89-100.
Not Relevant to current fit for purpose review	Wallace, K., et al. (2015). "A multiplexed assay for determination of neurotoxicant effects on spontaneous network activity and viability from microelectrode arrays." <i>NeuroToxicology</i> 49: 79-85.
Not Relevant to current fit for purpose review	Wander, J. G. N. and H. J. Bouwmeester (1998). "Effects of nitrogen fertilization on dill (<i>Anethum graveolens</i> L.) seed and carvone production." <i>Industrial Crops and Products</i> 7(2-3): 211-216.
Not Relevant to current fit for purpose review	Wang, F., et al. (2008). "The microtubule plus end-binding protein EB1 is involved in Sertoli cell plasticity in testicular seminiferous tubules." <i>Experimental Cell Research</i> 314(1): 213-226.
Not Relevant to current fit for purpose review	Wang, G., et al. (2006). "Methyl protodioscin induces G2/M cell cycle arrest and apoptosis in HepG2 liver cancer cells." <i>Cancer Letters</i> 241(1): 102-109.
Not Relevant to current fit for purpose review	Wang, G., et al. (2012). "Damage to DNA caused by UV-B radiation in the desert cyanobacterium <i>Scytonema javanicum</i> and the effects of exogenous chemicals on the process." <i>Chemosphere</i> 88(4): 413-417.
Not Relevant to current fit for purpose review	Wang, G., et al. (2012). "3.105 GLYPHOSATE INDUCED CELL DEATH THROUGH APOPTOTIC AND AUTOPHAGIC MECHANISMS." <i>Parkinsonism & Related Disorders</i> 18, Supplement 2: S190.
Not Relevant to current fit for purpose review	Wang, M., et al. (2015). "Gene editing by co-transformation of TALEN and chimeric RNA/DNA oligonucleotides on the rice OsEPSPS gene and the inheritance of mutations." <i>PLoS One</i> 10(4): e0122755.
Not Relevant to current fit for purpose review	Wang, S.-A., et al. (2009). "Heat Shock Protein 90 Is Important for Sp1 Stability during Mitosis." <i>Journal of Molecular Biology</i> 387(5): 1106-1119.
Not Relevant to current fit for purpose review	Wang, X., et al. (2005). "Effects of Phe-to-Trp mutation and fluorotryptophan incorporation on the solution structure of cardiac troponin C, and analysis of its suitability as a potential probe for in situ NMR studies." <i>Protein Science</i> 14(9): 2447-2460.
Not Relevant to current fit for purpose review	Wang, X., et al. (2011). "Construction of a reference plasmid molecule containing eight targets for the detection of genetically modified crops." <i>Appl Microbiol Biotechnol</i> 90(2): 721-731.
Not Relevant to current fit for purpose review	Weng, H., et al. (2005). "Novel reference gene, High-mobility-group protein I/Y, used in qualitative and real-time quantitative polymerase chain reaction detection of transgenic rapeseed cultivars." <i>J AOAC Int</i> 88(2): 577-584.
Not Relevant to current fit for purpose review	White, A. K. and W. W. Metcalf (2004). "Two C-P lyase Operons in <i>Pseudomonas stutzeri</i> and their roles in the oxidation of phosphonates, phosphite, and hypophosphite." <i>Journal of Bacteriology</i> 186(14): 4730-4739.
Not Relevant to current fit for purpose review	Willenborg, C. J. and E. N. Johnson (2013). "Influence of seeding date and seeding rate on cow cockle, a new medicinal and industrial crop." <i>Industrial Crops and Products</i> 49: 554-560.
Not Relevant to current fit for purpose review	Williams, M. J., et al. (2002). "No-till establishment of rhizoma peanut." <i>Agronomy Journal</i> 94(6): 1350-1354.
Not Relevant to current fit for purpose review	Williamson, P. R., et al. (2016). "CNS infections in 2015: emerging catastrophic infections and new insights into neuroimmunological host damage." <i>The Lancet Neurology</i> 15(1): 17-19.
Not Relevant to current fit for purpose review	Wilson, R. (1970). "AORN RECRUITMENT PROGRAM ROUND-UP." <i>AORN Journal</i> 11(1): 66-68.
Not Relevant to current fit for purpose review	Wiznitzer, M. (2006). "Paediatric neurology: clinical advances with low technology." <i>The Lancet Neurology</i> 5(1): 14-15.
Not Relevant to current fit for purpose review	Wong, P. K. Y., et al. (1973). "Rapid, selective procedure for isolation of spontaneous temperature-sensitive mutants of Moloney leukemia virus." <i>Virology</i> 51(2): 424-431.
Not Relevant to current fit for purpose review	Wood, E. J. (1985). "From cells to atoms: an illustrated introduction to molecular biology: by A. R. Rees and M. J. E. Sternberg, Blackwell, 1984. £6.80 (viii + 94 pages) ISBN 0 632 00888 1." <i>Trends in Biochemical Sciences</i> 10(5): 214-215.
Not Relevant to current fit for purpose review	Wood, E. J. (1985). "Lecture notes on biochemistry: by J. K. Candish, Blackwell, 1984. £9.80 (vi + 290 pages) ISBN 0 632 01253 6." <i>Trends in Biochemical Sciences</i> 10(5): 214-215.
Not Relevant to current fit for purpose review	Wurz, A., et al. (1999). "Quantitative analysis of genetically modified organisms (GMO) in processed food by PCR-based methods." <i>Food Control</i> 10(<HT>6</HT>): 385-389.
Not Relevant to current fit for purpose review	Xia, J., et al. (2015). "Retraction of a study on genetically modified corn: Expert investigations should speak louder during controversies over safety." <i>Bioscience Trends</i> 9(2): 134-137.

Not Relevant to current fit for purpose review	Xu, J., et al. (2002). "Cloning of genomic DNA of rice 5-enolpyruvylshikimate 3-phosphate synthase gene and chromosomal localization of the gene." <i>Sci China C Life Sci</i> 45(3): 251-259.
Not Relevant to current fit for purpose review	Xu, J. W., et al. (2002). "Isolation of rice EPSP synthase cDNA and its sequence - Analysis and copy number determination." <i>Acta Botanica Sinica</i> 44(2): 188-192.
Not Relevant to current fit for purpose review	Xu, W. T., et al. (2011). "Establishment and evaluation of event-specific qualitative and quantitative PCR method for genetically modified soybean DP-356043-5." <i>European Food Research and Technology</i> 233(4): 685-695.
Not Relevant to current fit for purpose review	Yan, H.-Q., et al. (2011). "Novel AroA from <i>Pseudomonas putida</i> Confers Tobacco Plant with High Tolerance to Glyphosate." <i>PLoS One</i> 6(5).
Not Relevant to current fit for purpose review	Yang, L. T., et al. (2005). "Screening and construct-specific detection methods of transgenic Huafan No 1 tomato by conventional and real-time PCR." <i>Journal of the Science of Food and Agriculture</i> 85(13): 2159-2166.
Not Relevant to current fit for purpose review	Yang, R., et al. (2007). "Event-specific qualitative and quantitative PCR detection of roundup ready event GT73 based on the 3'-integration junction." <i>Plant Cell Rep</i> 26(10): 1821-1831.
Not Relevant to current fit for purpose review	Yang, Z. and X. Dong (2015). "Earnings roundup in private and public bank holding companies." <i>Advances in Accounting</i> 31(1): 96-99.
Not Relevant to current fit for purpose review	Yao, P., et al. (2015). "Improvement of glycine oxidase by DNA shuffling, and site-saturation mutagenesis of F247 residue." <i>Int J Biol Macromol</i> 79: 965-970.
Not Relevant to current fit for purpose review	Yilmaz, R., et al. "Verification of GMO analytical methods: DNA isolation and quantitative detection methods for Roundup Ready soy flour." <i>New Biotechnology</i> 29, Supplement: S196-S197.
Not Relevant to current fit for purpose review	You, Y., et al. (2012). "Effect of intravenous fat emulsion therapy on glyphosate-surfactant-induced cardiovascular collapse." <i>The American Journal of Emergency Medicine</i> 30(9): 2097.e2091-2097.e2092.
Not Relevant to current fit for purpose review	Yousef, M. I., et al. (1996). "A sensitive sperm-motility test for the assessment of cytotoxic effect of pesticides." <i>Journal of Environmental Science and Health Part B-Pesticides Food Contaminants and Agricultural Wastes</i> 31(1): 99-115.
Not Relevant to current fit for purpose review	Zanis, D. A., et al. (1997). "A comparison of three methods of measuring the type and quantity of services provided during substance abuse treatment." <i>Drug and Alcohol Dependence</i> 49(1): 25-32.
Not Relevant to current fit for purpose review	Zanoli, L. M., et al. (2013). "Peptide nucleic acid molecular beacons for the detection of PCR amplicons in droplet-based microfluidic devices." <i>Anal Bioanal Chem</i> 405(2-3): 615-624.
Not Relevant to current fit for purpose review	Zawahir, S., et al. (2009). "Acute intentional self-poisoning with a herbicide product containing fenoxaprop-P-ethyl, ethoxysulfuron, and isoxadifen ethyl: a prospective observational study." <i>Clinical Toxicology</i> 47(8): 792-797.
Not Relevant to current fit for purpose review	Zhan, T., et al. (2013). "Improving Glyphosate Oxidation Activity of Glycine Oxidase from <i>Bacillus cereus</i> by Directed Evolution." <i>PLoS One</i> 8(11).
Not Relevant to current fit for purpose review	Zhang, H., et al. (2008). "Development of one novel multiple-target plasmid for duplex quantitative PCR analysis of roundup ready soybean." <i>Journal of Biotechnology</i> 136, Supplement: S250.
Not Relevant to current fit for purpose review	Zhang, J., et al. (1998). "A transformation technique from RGB signals to the Munsell system for color analysis of tobacco leaves." <i>Computers and Electronics in Agriculture</i> 19(2): 155-166.
Not Relevant to current fit for purpose review	Zhang, M., et al. (2007). "Detection of Roundup Ready soy in highly processed products by triplex nested PCR." <i>Food Control</i> 18(10): 1277-1281.
Not Relevant to current fit for purpose review	Zhang, Q., et al. (2014). "Characterization of cytochalasins from the endophytic <i>Xylaria</i> sp. and their biological functions." <i>J Agric Food Chem</i> 62(45): 10962-10969.
Not Relevant to current fit for purpose review	Zheng, Q. and D. C. Chang (1991). "High-efficiency gene transfection by in situ electroporation of cultured cells." <i>Biochimica et Biophysica Acta (BBA) - Gene Structure and Expression</i> 1088(1): 104-110.
Not Relevant to current fit for purpose review	Zhou, H., et al. (1995). "Glyphosate-tolerant CP4 and GOX genes as a selectable marker in wheat transformation." <i>Plant Cell Rep</i> 15(3-4): 159-163.
Not Relevant to current fit for purpose review	Zhu, X., et al. (2012). "A high-throughput fluorescence resonance energy transfer (FRET)-based endothelial cell apoptosis assay and its application for screening vascular disrupting agents." <i>Biochemical and Biophysical Research Communications</i> 418(4): 641-646.
Not Relevant to current fit for purpose review	Ziegelberger, G., et al. (2006). "International commission on non-ionizing radiation protection." <i>Progress in Biophysics and Molecular Biology</i> 92(1): 1-3.
Not Relevant to current fit for purpose review	Zulet, A., et al. (2015). "Fermentation and alternative oxidase contribute to the action of amino acid biosynthesis-inhibiting herbicides." <i>J Plant Physiol</i> 175: 102-112.

Not Relevant to current fit for purpose review	정원중, et al. (2006). "Prognostic Predictors of Outcome for Poisoning by Glyphosate-containing Herbicides, Based on Initial Findings." <i>Journal of The Korean Society of Emergency Medicine</i> 17(6): 630-636.
Not Relevant to current fit for purpose review	Boyle, W. S. and J. O. Evans (1974). "EFFECTS OF GLYPHOSATE AND ETHEPHON ON MEIOTIC CHROMOSOMES OF SECALE-CEREALE L." <i>Journal of Heredity</i> 65(4): 250-250.
Not Relevant to current fit for purpose review	Sherwood, M. M. and W. C. Davison (1957). "Correspondence." <i>The Journal of Pediatrics</i> 51(4): 486-487.
Not Relevant to current fit for purpose review	Belle, R., et al. (2007). "[Sea urchin embryo, DNA-damaged cell cycle checkpoint and the mechanisms initiating cancer development]." <i>J Soc Biol</i> 201(3): 317-327.
Not Relevant to current fit for purpose review	Gehin, A., et al. (2005). "Vitamins C and E reverse effect of herbicide-induced toxicity on human epidermal cells HaCaT: a biochemometric approach." <i>Int J Pharm</i> 288(2): 219-226.
Not Relevant to current fit for purpose review	Gehin, A., et al. (2006). "Glyphosate-induced antioxidant imbalance in HaCaT: The protective effect of vitamins C and E." <i>Environmental Toxicology and Pharmacology</i> 22(1): 27-34.
Not Relevant to current fit for purpose review	Lueken, A., et al. (2004). "Synergistic DNA damage by oxidative stress (induced by H2O2) and nongenotoxic environmental chemicals in human fibroblasts." <i>Toxicol Lett</i> 147(1): 35-43.
Not Relevant to current fit for purpose review	Baurand, P. E., et al. (2015). "Genotoxicity assessment of pesticides on terrestrial snail embryos by analysis of random amplified polymorphic DNA profiles." <i>J Hazard Mater</i> 298: 320-327.
Not Relevant to current fit for purpose review	Guilherme, S., et al. (2009). "Tissue specific DNA damage in the European eel (<i>Anguilla anguilla</i>) following a short-term exposure to a glyphosate-based herbicide." <i>Toxicology Letters</i> 189: S212-S212.
Not Relevant to current fit for purpose review	Marc, J., et al. (2004). "Glyphosate-based pesticides affect cell cycle regulation." <i>Biol Cell</i> 96(3): 245-249.
Not Relevant to current fit for purpose review	Marc, J., et al. (2003). "Embryonic cell cycle for risk assessment of pesticides at the molecular level." <i>Environmental Chemistry Letters</i> 1(1): 8-12.
Not Relevant to current fit for purpose review	Nwani, C. D., et al. (2014). "Induction of micronuclei and nuclear lesions in <i>Channa punctatus</i> following exposure to carbosulfan, glyphosate and atrazine." <i>Drug Chem Toxicol</i> 37(4): 370-377.
Not Relevant to current fit for purpose review	Owczarek, M., et al. (1999). "Evaluation of toxic and genotoxic activity of some pesticides in a soil-plant system." <i>Human and Environmental Exposure to Xenobiotics</i> : 755-762.
Not Relevant to current fit for purpose review	Perez-Iglesias, J. M., et al. (2016). "Effects of glyphosate on hepatic tissue evaluating melanomacrophages and erythrocytes responses in neotropical anuran <i>Leptodactylus latinasus</i> ." <i>Environ Sci Pollut Res Int</i> .
Not Relevant to current fit for purpose review	Poletta, G. L., et al. (2009). "Genotoxicity of the herbicide formulation Roundup (glyphosate) in broad-snouted caiman (<i>Caiman latirostris</i>) evidenced by the Comet assay and the Micronucleus test." <i>Mutat Res</i> 672(2): 95-102.
Not Relevant to current fit for purpose review	Siddiqui, S., et al. (2012). "Glyphosate, alachor and maleic hydrazide have genotoxic effect on <i>Trigonella foenum-graecum</i> L." <i>Bull Environ Contam Toxicol</i> 88(5): 659-665.
Not Relevant to current fit for purpose review	Song, H.-Y., et al. (2012). "Cellular Toxicity of Surfactants Used as Herbicide Additives." <i>J Korean Med Sci</i> 27(1): 3-9.
Not Relevant to current fit for purpose review	Unver, T., et al. (2010). "Genome-wide profiling and analysis of <i>Festuca arundinacea</i> miRNAs and transcriptomes in response to foliar glyphosate application." <i>Mol Genet Genomics</i> 283(4): 397-413.
Relevant- Cancer Epi	Acquavella, J. F., et al. (2006). "Exposure misclassification in studies of agricultural pesticides - Insights from biomonitoring." <i>Epidemiology</i> 17(1): 69-74.
Relevant- Cancer Epi	Acquavella, J. F., et al. (2005). "Implications for epidemiologic research on variation by pesticide in studies of farmers and their families." <i>Scandinavian Journal of Work Environment & Health</i> 31: 105-109.
Relevant- Cancer Epi	Baker, B. A., et al. (2005). "Farm Family Exposure Study: methods and recruitment practices for a biomonitoring study of pesticide exposure." <i>Journal of Exposure Analysis and Environmental Epidemiology</i> 15(6): 491-499.
Relevant- Cancer Epi	Chang, E. T. and E. Delzell (2016). "Systematic review and meta-analysis of glyphosate exposure and risk of lymphohematopoietic cancers." <i>J Environ Sci Health B</i> 51(6): 402-434.
Relevant- Cancer Epi	De Roos, A. J., et al. (2005). "Cancer incidence among glyphosate-exposed pesticide applicators in the Agricultural Health Study." <i>Environ Health Perspect</i> 113(1): 49-54.
Relevant- Cancer Epi	Firth, H. M., et al. (2007). "Chemical exposure among NZ farmers." <i>International Journal of Environmental Health Research</i> 17(1): 33-43.

Relevant- Cancer Epi	Lash, T. L. (2007). "Bias analysis applied to Agricultural Health Study publications to estimate non-random sources of uncertainty." <i>J Occup Med Toxicol</i> 2: 15.
Relevant- Cancer Epi	Mink, P. J., et al. (2012). "Epidemiologic studies of glyphosate and cancer: a review." <i>Regul Toxicol Pharmacol</i> 63(3): 440-452.
Relevant- Cancer Epi	Sorahan, T. (2012). "Multiple myeloma and glyphosate use: A re-analysis of US Agricultural Health Study data." <i>Toxicology Letters</i> 211, Supplement: S169.
Relevant- Carcinogenicity	Greim, H., et al. (2015). "Evaluation of carcinogenic potential of the herbicide glyphosate, drawing on tumor incidence data from fourteen chronic/carcinogenicity rodent studies." <i>Crit Rev Toxicol</i> 45(3): 185-208.
Relevant- Carcinogenicity	Williams, G. M., et al. (2000). "Safety Evaluation and Risk Assessment of the Herbicide Roundup and Its Active Ingredient, Glyphosate, for Humans." <i>Regulatory Toxicology and Pharmacology</i> 31(2): 117-165.
Relevant- Genotoxicity	Alvarez-Moya et al. (2014) "Comparison of the in vivo and in vitro genotoxicity of glyphosate isopropylamine salt in three different organisms". <i>Genetics and Molecular Biology</i> , 37, 1, 105-110
Relevant- Genotoxicity	Chan, P. and J. Mahler (1992). "NTP technical report on the toxicity studies of Glyphosate (CAS No. 1071-83-6) Administered In Dosed Feed To F344/N Rats And B6C3F1 Mice." <i>Toxic Rep Ser</i> 16: 1-d3.
Relevant- Genotoxicity	Bakry, F. A., et al. (2015). "Glyphosate herbicide induces genotoxic effect and physiological disturbances in <i>Bulinus truncatus</i> snails." <i>Pestic Biochem Physiol</i> 123: 24-30.
Relevant- Genotoxicity	Bolognesi, C., et al. (1997). "Genotoxic activity of glyphosate and its technical formulation roundup." <i>J Agric Food Chem</i> 45(5): 1957-1962.
Relevant- Genotoxicity	Bolognesi, C., et al. (2009). "Biomonitoring of genotoxic risk in agricultural workers from five colombian regions: association to occupational exposure to glyphosate." <i>J Toxicol Environ Health A</i> 72(15-16): 986-997.
Relevant- Genotoxicity	Da Silva, F. R., et al. (2014). "Genotoxic assessment in tobacco farmers at different crop times." <i>Science of the Total Environment</i> 490: 334-341.
Relevant- Genotoxicity	Dimitrov, B. D., et al. (2006). "Comparative genotoxicity of the herbicides Roundup, Stomp and Reglone in plant and mammalian test systems." <i>Mutagenesis</i> 21(6): 375-382.
Relevant- Genotoxicity	El-Shenawy, N. S. (2009). "Oxidative stress responses of rats exposed to Roundup and its active ingredient glyphosate." <i>Environ Toxicol Pharmacol</i> 28(3): 379-385.
Relevant- Genotoxicity	Fortes, C., et al. (2016). "Occupational Exposure to Pesticides With Occupational Sun Exposure Increases the Risk for Cutaneous Melanoma." <i>J Occup Environ Med</i> 58(4): 370-375.
Relevant- Genotoxicity	Ghisi Nde, C., et al. (2016). "Does exposure to glyphosate lead to an increase in the micronuclei frequency? A systematic and meta-analytic review." <i>Chemosphere</i> 145: 42-54.
Relevant- Genotoxicity	Heydens, W. F., et al. (2008). "Genotoxic potential of glyphosate formulations: mode-of-action investigations." <i>J Agric Food Chem</i> 56(4): 1517-1523.
Relevant- Genotoxicity	Kale, P. G., et al. (1995). "MUTAGENICITY TESTING OF 9 HERBICIDES AND PESTICIDES CURRENTLY USED IN AGRICULTURE." <i>Environ Mol Mutagen</i> 25(2): 148-153.
Relevant- Genotoxicity	Kier, L. D. (2015). "Review of genotoxicity biomonitoring studies of glyphosate-based formulations." <i>Crit Rev Toxicol</i> 45(3): 209-218.
Relevant- Genotoxicity	Koller, V. J., et al. (2012). "Cytotoxic and DNA-damaging properties of glyphosate and Roundup in human-derived buccal epithelial cells." <i>Arch Toxicol</i> 86(5): 805-813.
Relevant- Genotoxicity	Li, A. P. and T. J. Long (1988). "An evaluation of the genotoxic potential of glyphosate." <i>Fundam Appl Toxicol</i> 10(3): 537-546.
Relevant- Genotoxicity	Lioi, M. B., et al. (1998). "Genotoxicity and oxidative stress induced by pesticide exposure in bovine lymphocyte cultures in vitro." <i>Mutat Res</i> 403(1-2): 13-20.
Relevant- Genotoxicity	Manas, F., et al. (2009). "Genotoxicity of AMPA, the environmental metabolite of glyphosate, assessed by the Comet assay and cytogenetic tests." <i>Ecotoxicol Environ Saf</i> 72(3): 834-837.
Relevant- Genotoxicity	Manas, F., et al. (2009). "Genotoxicity of glyphosate assessed by the comet assay and cytogenetic tests." <i>Environmental Toxicology and Pharmacology</i> 28(1): 37-41.
Relevant- Genotoxicity	Mandel, J. S., et al. (2005). "Biomonitoring for farm families in the farm family exposure study." <i>Scandinavian Journal of Work Environment & Health</i> 31: 98-104.
Relevant- Genotoxicity	Mladinic, M., et al. (2009). "Evaluation of genome damage and its relation to oxidative stress induced by glyphosate in human lymphocytes in vitro." <i>Environ Mol Mutagen</i> 50(9): 800-807.
Relevant- Genotoxicity	Mladinic, M. and D. Zeljezic (2008). "Assessment of oxidative DNA damage by glyphosate applying hOGG1 modified comet and micronucleus assay." <i>Toxicology Letters</i> 180: S170-S171.

Relevant- Genotoxicity	Paz-y-Mino, C., et al. (2011). "Baseline determination in social, health, and genetic areas in communities affected by glyphosate aerial spraying on the northeastern Ecuadorian border." <i>Rev Environ Health</i> 26(1): 45-51.
Relevant- Genotoxicity	Paz-y-Mino, C., et al. (2007). "Evaluation of DNA damage in an Ecuadorian population exposed to glyphosate." <i>Genetics and Molecular Biology</i> 30(2): 456-460.
Relevant- Genotoxicity	Peluso, M., et al. (1998). " ³² P-postlabeling detection of DNA adducts in mice treated with the herbicide Roundup." <i>Environ Mol Mutagen</i> 31(1): 55-59.
Relevant- Genotoxicity	Piesova, E. (2005). "The effect of glyphosate on the frequency of micronuclei in bovine lymphocytes in vitro." <i>Acta Veterinaria-Beograd</i> 55(2-3): 101-109.
Relevant- Genotoxicity	Prasad, S., et al. (2009). "Clastogenic effects of glyphosate in bone marrow cells of swiss albino mice." <i>J Toxicol</i> 2009: 308985.
Relevant- Genotoxicity	Rank, J., et al. (1993). "Genotoxicity testing of the herbicide Roundup and its active ingredient glyphosate isopropylamine using the mouse bone marrow micronucleus test, Salmonella mutagenicity test, and Allium anaphase-telophase test." <i>Mutat Res</i> 300(1): 29-36.
Relevant- Genotoxicity	Roustan, A., et al. (2014). "Genotoxicity of mixtures of glyphosate and atrazine and their environmental transformation products before and after photoactivation." <i>Chemosphere</i> 108: 93-100.
Relevant- Genotoxicity	Silva Kahl, V. F., et al. (2016). "Telomere measurement in individuals occupationally exposed to pesticide mixtures in tobacco fields." <i>Environ Mol Mutagen</i> 57(1): 74-84.
Relevant- Genotoxicity	Sivikova, K. and J. Dianovsky (2006). "Cytogenetic effect of technical glyphosate on cultivated bovine peripheral lymphocytes." <i>Int J Hyg Environ Health</i> 209(1): 15-20.
Relevant- Genotoxicity	Vigfusson, N. V. and E. R. Vyse (1980). "The effect of the pesticides, Dexon, Captan and Roundup, on sister-chromatid exchanges in human lymphocytes in vitro." <i>Mutat Res</i> 79(1): 53-57.
Retracted Article	Séralini, G.-E., et al. (2014). "Retraction notice to "Long term toxicity of a Roundup herbicide and a Roundup-tolerant genetically modified maize" [<i>Food Chem. Toxicol.</i> 50 (2012) 4221–4231]." <i>Food and Chemical Toxicology</i> 63: 244.

Appendix B

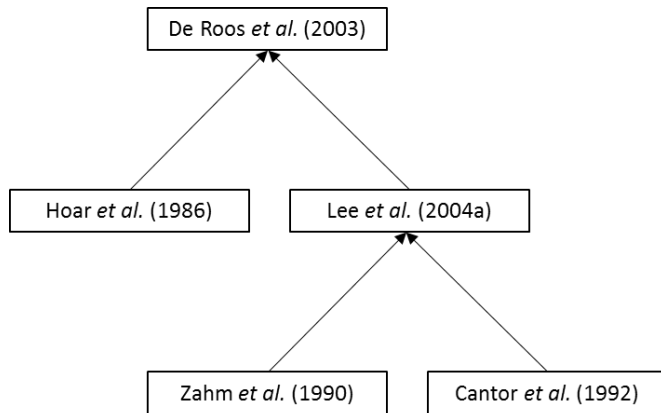


Figure B.1. Visual representation of studies included in De Roos *et al.* (2003).

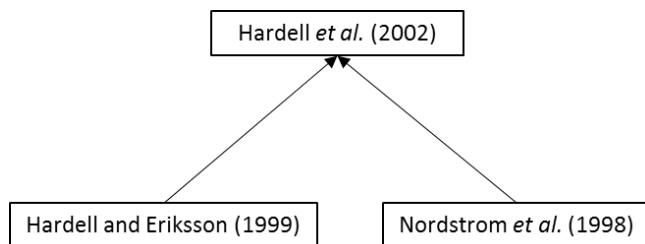


Figure B.2. Visual representation of studies included in Hardell *et al.* (2002).

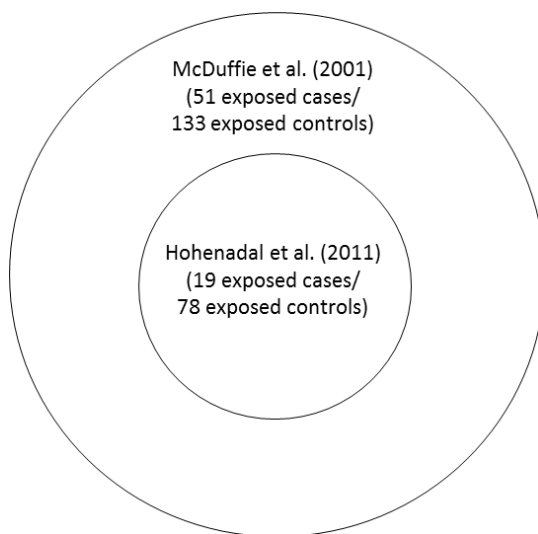


Figure B.3. Visual representation of the association between McDuffie *et al.* (2001) and the follow-up analysis by Hohenadal *et al.* (2011).

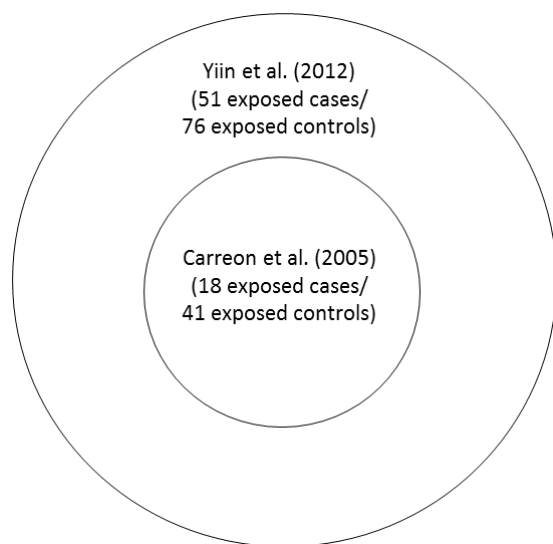


Figure B.4. Visual representation of the association between Carreon *et al.* (2005), which investigated gliomas in women only, and Yiin *et al.* (2012), which investigated both sexes.

Appendix C

Table B.1. Design Characteristics of Epidemiological Studies Evaluated for Study Quality.

Study	Location	Study Years	Case Population	Control Population	Total Number of Subjects	Number of Glyphosate Exposed Cases	Proxy Use
Alavanja <i>et al.</i> (2003)	USA: Iowa and North Carolina	Enrollment (1993-1997) through 2001	Males enrolled in AHS; licensed private and commercial applicators	Males enrolled in AHS; licensed private and commercial applicators	566 cases 54,766 controls	not reported	No
Andreotti <i>et al.</i> (2009)	USA: Iowa and North Carolina	Enrollment (1993-1997) through 2004	Participants enrolled in AHS; licensed private and commercial applicators and spouses	Participants enrolled in AHS; licensed private and commercial applicators and spouses	93 cases (64 applicators, 29 spouses) 82,503 controls (52,721 applicators, 29,782 spouses)	55 cases 48,461 controls	No
Band <i>et al.</i> (2011)	Canada: British Columbia	1983-1990	Male residents in British Columbia identified as cancer patients in British Columbia Cancer Registry (excluding farmers that worked all outside British Columbia)	Male residents in British Columbia identified as cancer patients in British Columbia Cancer Registry (excluding farmers that worked all outside British Columbia) with other cancer sites excluding lung cancer and cancers of unknown primary site	1,153 cases 3,999 controls	25 cases 60 controls	Yes (included in adjustment)
Brown <i>et al.</i> (1990)	USA: Iowa and Minnesota	Iowa: 1981-1983; Minnesota: 1980-1982 Initial interview 1981-1984 and supplemental interviews (Iowa only) in 1987	White males (30 years or older) residing in Iowa or Minnesota diagnosed with leukemia	White males without lymphatic or hematopoietic cancer selected by random digit dialing (< age 65), Medicare records (age > 65) and state death certificate files (deceased controls) - frequency matched for 5-year age group, vital status, and state of residence	Initial: 578 cases; 1245 controls Supplemental: 92 cases; 211 controls	15 cases 49 controls	Yes (not evaluated)
Brown <i>et al.</i> (1993)	USA: Iowa	Iowa: 1981-1983; Interview 1981-1984	White males (30 years or older) residing in Iowa diagnosed with multiple myeloma	White males without lymphatic or hematopoietic cancer selected by random digit dialing (< age 65), Medicare records (age >	173 cases 650 controls	11 cases 40 controls	Yes (not evaluated)

Table B.1. Design Characteristics of Epidemiological Studies Evaluated for Study Quality.							
Study	Location	Study Years	Case Population	Control Population	Total Number of Subjects	Number of Glyphosate Exposed Cases	Proxy Use
				65) and state death certificate files (deceased controls) - frequency matched for 5-year age group, vital status, and state of residence			
Cocco <i>et al.</i> (2013)	Czech Republic, France, Germany, Italy, Ireland, and Spain	1998-2004	Adult patients first diagnosed with lymphoma residing in the referral area of the participating centers	Controls from Germany and Italy were randomly selected by sampling from the general population, matched to cases on sex, 5-year age-group, and residence area. The rest of the centers used matched hospital controls, with eligibility criteria limited to diagnoses other than cancer, infectious diseases, and immunodeficient diseases	2,348 cases 2,462 controls	4 cases 2 controls	No
De Roos <i>et al.</i> (2003)	USA: Nebraska, Iowa, Minnesota, and Kansas	Nebraska: 1983-1986 Iowa: 1981-1983 Minnesota: 1980-1982 Kansas: 1979-1981	White males diagnosed with NHL in one of the 4 states (21 years or older in Nebraska and Kansas; 30 years or older in Iowa and Minnesota)	Males living in same geographic area obtained by random digit dialing, Medicare records and state mortality files - frequency matched for race, sex, age, and vital status	870 cases 2,569 controls	36 cases 61 controls	Yes (not significant in covariate analysis)
De Roos <i>et al.</i> (2005)	USA: Iowa and North Carolina	Enrollment (1993-1997) through 2001	Participants enrolled in AHS; licensed private and commercial applicators and spouses	Participants enrolled in AHS; licensed private and commercial applicators and spouses	54,315 subjects included in this analysis	All cancers – 358 cases Lung – 26 cases Oral cavity – 10 cases Colon – 15 cases Rectum – 14 cases Pancreas – 7 cases Kidney – 9 cases Bladder – 17 cases Prostate – 145 cases Melanoma – 14 cases All lymphohematopoietic cancers – 36 cases NHL – 17 cases Leukemia – 9 cases	No

Table B.1. Design Characteristics of Epidemiological Studies Evaluated for Study Quality.							
Study	Location	Study Years	Case Population	Control Population	Total Number of Subjects	Number of Glyphosate Exposed Cases	Proxy Use
						Multiple myeloma – 6 cases (13,280 subjects not exposed to glyphosate used for comparison population)	
Engel <i>et al.</i> (2005)	USA: Iowa and North Carolina	Enrollment (1993-1997) through 2000	Wives of applicators enrolled in AHS study with no history of breast cancer	Wives of applicators enrolled in AHS study with no history of breast cancer	309 cases 30,145 controls	82 cases; 10,016 controls	No
Eriksson <i>et al.</i> (2008)	Sweden	1999-2002	Patients (18-74 years of age) residing in Sweden and diagnosed with NHL	Swedish residents randomly selected living in same health service regions as cases - frequency matched for age (in 10 years) and sex	910 cases 1,016 controls	29 cases 18 controls	No
Flower <i>et al.</i> (2004)	USA: Iowa	1993-1997	Children (born after 1975) of participants enrolled in AHS study who were diagnosed with childhood cancer up to 19 years of age	Children (born after 1975) of participants enrolled in AHS study not diagnosed with childhood cancer up to 19 years of age	50 cases out of 17,357 total study population	Maternal use: 13 cases of 6075 total exposed Paternal use: 6 cases of 3231 total exposed	No
Hardell <i>et al.</i> (2002)	Sweden	NHL: 1987-1990 HCL: 1987-1992	NHL: Male residents of one of four northern or three middle counties in Sweden age 25 years and older diagnosed with NHL; identified from regional cancer registries HCL: Living male residents of Sweden age 25 years and older diagnosed with HCL; identified from the Swedish Cancer Registry	NHL: Two male controls for each case matched by age, year of death if deceased, and county HCL: Four male controls for each case matched by age and county	515 cases 1,141 controls	8 cases 8 controls	Yes (not evaluated)
Kachuri <i>et al.</i> (2013)	Canada: Alberta, British Columbia, Manitoba, Ontario,	1991–1994	Men aged ≥ 19 years (≥ 30 years in analysis) - pulled from hospital records in Quebec,	Men aged ≥ 19 years (30 years in analysis) - pulled from provincial health insurance records in	342 cases 1,357 controls	32 cases 121 controls	Yes (included in adjustment)

Table B.1. Design Characteristics of Epidemiological Studies Evaluated for Study Quality.							
Study	Location	Study Years	Case Population	Control Population	Total Number of Subjects	Number of Glyphosate Exposed Cases	Proxy Use
	Quebec, and Saskatchewan		cancer registries in all other provinces	Alberta, Saskatchewan, Manitoba, and Quebec; computerized telephone listings in Ontario; voter lists in British Columbia			
Karunanayake <i>et al.</i> (2012)	Canada: Alberta, British Columbia, Manitoba, Ontario, Quebec, and Saskatchewan	1991–1994	Men aged ≥ 19 years - pulled from hospital records in Quebec, cancer registries in all other provinces	Men aged ≥ 19 years - pulled from provincial health insurance records in Alberta, Saskatchewan, Manitoba, and Quebec; computerized telephone listings in Ontario; voter lists in British Columbia	316 cases 1,506 controls	38 cases 133 controls	No
Koureas <i>et al.</i> (2014)	Greece	2010	Inhabitants of the city of Larissa; Eligibility criteria for pesticide sprayers were 1) to personally apply pesticides systematically, and 2) to have recently applied pesticides (no longer than 7 days between last application and sampling).	The rural residents group were occupied in administrative services, public order services, health services, education or trade. Inclusion criteria for this group: absence of any involvement in agricultural activities either as a primary or secondary occupation by participant or any member of household. Also recruited urban residents (mainly blood donors) from the city of Larissa.	80 pesticide sprayers, 85 rural residents, and 121 individuals	Not reported	No
Koutros <i>et al.</i> (2013)	USA: Iowa and North Carolina	Enrollment (1993-1997) through 2007	Males enrolled in AHS; licensed private and commercial applicators	Males enrolled in AHS; licensed private and commercial applicators	1,962 incident cases (including 919 aggressive prostate cancers) among 54,412 applicators	1464 cases 42,420 controls	No
Landgren <i>et al.</i> (2009)	USA: Iowa and North Carolina	Exposure information: enrollment (1993-1997) and 5-year follow-up interview	Males enrolled in AHS; licensed private and commercial applicators	Males enrolled in AHS; licensed private and commercial applicators	678 participants	27 cases out of 570 total exposed	No

Table B.1. Design Characteristics of Epidemiological Studies Evaluated for Study Quality.							
Study	Location	Study Years	Case Population	Control Population	Total Number of Subjects	Number of Glyphosate Exposed Cases	Proxy Use
		Blood samples: 2006-2007 (Iowa) and 2008 (North Carolina)					
Lee <i>et al.</i> (2004b)	USA: Nebraska	1988-1993	White residents of 1 of 66 Nebraska counties age 21 years or older with a newly confirmed case of adenocarcinoma of the stomach or Cases identified from the Nebraska Cancer Registry (1988–1990) or from discharge diagnosis and pathology records from 14 Nebraska hospitals (1991–1993)	Frequency matched by age and sex to the combined distribution of glioma, stomach, and esophageal cancer cases from a control group from a previous study (1986–1987) that selected controls from the general population by random digit dialing for those under 65 years, Health Care Financing Administration Medicare files for those over 65 years, mortality records for deceased and matched by race, sex, vital status (or year of death if deceased)	Stomach: 170 cases Esophagus: 137 cases 502 Controls	12 cases 46 controls	Yes (analysis showed differences)
Lee <i>et al.</i> (2005)	USA: Nebraska	1988-1993	White residents of 1 of 66 Nebraska counties age 21 years or older with confirmed adult glioma. Cases identified from Nebraska Cancer Registry or from participating hospitals in Lincoln and Omaha, Nebraska	Frequency matched by age, sex, and vital status to the combined distribution of glioma, stomach, and esophageal cancer cases from a control group from a previous study (1986–1987) that selected controls from the general population by random digit dialing for those under 65 years, Medicare files for those over 65 years, mortality records for deceased and matched by race, sex, vital status (or year of death if deceased), and 5-year age groups to the overall case distribution. Additional	251 glioma cases 498 controls	17 cases 32 controls	Yes (analysis showed differences, included in adjustment)

Table B.1. Design Characteristics of Epidemiological Studies Evaluated for Study Quality.							
Study	Location	Study Years	Case Population	Control Population	Total Number of Subjects	Number of Glyphosate Exposed Cases	Proxy Use
				younger controls were brought into the study through random digit dialing and from death certificates			
Lee <i>et al.</i> (2007)	USA: Iowa and North Carolina	1993-97; follow-up to 2002	Agricultural Health Study participants: private and commercial applicators licensed in Iowa or North Carolina with no history of colorectal cancer at enrollment. Followed through 2002 for incident colorectal cancer	Agricultural Health Study participants: private and commercial applicators licensed in Iowa or North Carolina with no history of colorectal cancer at enrollment. Followed through 2002 for incident colorectal cancer	56,813 licensed pesticide applicators 305 incident colorectal cancer cases (212 colon, 93 rectum) 56,508 controls	Colon - 151 cases; 49 controls Rectum - 74 cases; 18 controls Colorectal - 225 cases; 67 controls	No
McDuffie <i>et al.</i> (2001)	Canada: Alberta, British Columbia, Manitoba, Ontario, Quebec, and Saskatchewan	1991-1994	Male residents of six Canadian provinces age 19 years and older diagnosed with STS, HD, NHL, or MM; this study only evaluated those with NHL. Cases were identified from Canadian Cancer Registries; in Quebec, hospital ascertainment was used	Random control subject selection using Health Insurance records, computerized telephone listings, and voters' lists; males 19 years and older from same six Canadian provinces as cases matched by age (within 2 years)	517 cases 1506 controls	Univariate analysis: 51 cases; 133 controls (multivariate analysis also conducted - glyphosate exposed numbers not reported)	No
Orsi <i>et al.</i> (2009)	France	2000-2004	Men aged 20–75 years living in the catchment areas of the main hospitals in Brest, Caen, Nantes, Lille, Toulouse, and Bordeaux, with no history of immunosuppression or taking immunosuppressant drugs. Cases ascertained from hospital records.	Patients from the same hospital catchment area as the cases. Patients were hospitalized for orthopedic or rheumatological conditions (89.3%), gastrointestinal or genitourinary tract diseases (4.8%), cardiovascular diseases (1.1%), skin and subcutaneous tissue disease (1.8%), and infections (3.0%), excluding patients admitted for cancer or a disease directly related to	491 cases 456 controls	NHL: 12 cases 24 controls HL: 6 cases 15 controls Lymphoproliferative syndromes: 4 cases 18 controls Multiple myeloma: 5 cases; 18 controls Lymphoid neoplasms: 27 cases; 24 controls	No

Table B.1. Design Characteristics of Epidemiological Studies Evaluated for Study Quality.							
Study	Location	Study Years	Case Population	Control Population	Total Number of Subjects	Number of Glyphosate Exposed Cases	Proxy Use
				occupation, smoking, or alcohol abuse			
Pahwa <i>et al.</i> (2011)	Canada (Alberta, British Columbia, Manitoba, Ontario, Quebec, and Saskatchewan)	1991-1994	Men aged ≥ 19 years - pulled from hospital records in Quebec, cancer registries in all other provinces	Men aged ≥ 19 years - pulled from provincial health insurance records in Alberta, Saskatchewan, Manitoba, and Quebec; computerized telephone listings in Ontario; voter lists in British Columbia	342 cases 1,506 age/resident matched controls	32 cases 133 controls	No
Pahwa <i>et al.</i> (2012)	Canada (Alberta, British Columbia, Manitoba, Ontario, Quebec, and Saskatchewan)	1991-1994	Men aged ≥ 19 years - pulled from hospital records in Quebec, cancer registries in all other provinces	Men aged ≥ 19 years - pulled from provincial health insurance records in Alberta, Saskatchewan, Manitoba, and Quebec; computerized telephone listings in Ontario; voter lists in British Columbia	342 cases 1506 age/resident matched controls	32 cases 133 controls	No
Yiin <i>et al.</i> (2012)	USA: Upper Midwest Health Study (Iowa, Michigan, Minnesota and Wisconsin)	1995-1997	Age 18–80 (at ascertainment or diagnosis in 1995 through January 1997) residing in counties where the largest population center had fewer than 250,000 residents. Referral by physicians or through state cancer registries with cases verified by histological evaluation.	Controls age 18–64 randomly selected from state driver's license/nondriver ID records, and those age 65–80 were selected from Health Care Financing Administration's (HCFA) Medicare data within 10-year age group strata, with the proportion/stratum determined by the age distribution of glioma cases in that state from 1992 to 1994. Controls were frequency-matched within a state but not by county of residence. Selected even if they had a self-reported history of cancer other than glioma.	798 glioma cases; 1,175 controls	12 cases 19 controls	Yes (analysis showed no differences)

Appendix D. List of studies assigned a low quality ranking and not evaluated in detail

As described in Section 3.2, if studies did not collect exposure information on glyphosate from all subjects, did not assess an outcome (e.g., biomonitoring studies), and/or did not provide a quantitative measure of an association between glyphosate and a cancer outcome, then these studies were assigned a low quality ranking and were not further evaluated in detail. These studies included the following 32 studies:

Acquavella *et al.* 2006; Andre *et al.*, 2007; Baker *et al.* 2005; Benedetti *et al.*, 2013; Bolognesi *et al.*, 2002; Bolognesi *et al.*, 2004; Bolognesi *et al.* 2009; Bortoli *et al.*, 2009; Costa *et al.*, 2006; Da Silva *et al.* 2014; Dennis *et al.* 2010; Firth *et al.* 2007; Gomez-Arroyo *et al.*, 2013; Gregio D'Arce *et al.*, 2000; El-Zaemey *et al.*, 2013; Fortes *et al.*, 2016; Fritschi *et al.*, 2005; Hernandez *et al.*, 2006; Kaufman *et al.* 2009; Khayat *et al.*, 2013; Lebailly *et al.*, 2003; Mandel *et al.* 2005; Martinez-Valenzuela *et al.*, 2009; Monge *et al.*, 2007; Pastor *et al.*, 2003; Paz-y Mino *et al.*, 2007; Paz-y Mino *et al.* 2011; Ruder *et al.* 2004; Shaham *et al.*, 2001; Silva Kahl *et al.* 2016; Simoniello *et al.*, 2008; Vlastos *et al.*, 2006.

Appendix E

Chronic Dietary Exposure

The agency uses Dietary Exposure Evaluation Model- Food Consumption Intake Database (DEEM-FCID; version 3.16), which incorporates consumption data from United States Department of Agriculture (USDA) National Health and Nutrition Examination Survey, What We Eat in America (NHANES/WWEIA; 2003-2008) to calculate potential chronic dietary exposures. In an unrefined chronic dietary analysis, several conservative assumptions are used to generate high end estimates of potential exposure. These assumptions include tolerance-level residues for all registered commodities, 100% crop treated, and drinking water values from a direct application to water scenario, as well as DEEM default processing factors. For glyphosate, the highest exposure value for any population subgroup in an unrefined chronic dietary analysis would be 0.23 mg/kg/day for children 1-2 years old (Table E.1; DEEM inputs and results attached below).

Table E.1. Chronic dietary exposure estimates	
Population Subgroup	Exposure (mg/kg/day)
General U.S. Population	0.091515
All Infants (< 1 year old)	0.142826
Children 1-2 years old	0.230816
Children 3-5 years old	0.214117
Children 6-12 years old	0.149269
Youth 13-19 years old	0.089636
Adults 20-49 years old	0.076396
Adults 50-99 years old	0.062987
Females 13-49 years old	0.071057

Post-application Incidental Oral and Dermal Exposure

Glyphosate has residential uses, including application to turf, which would result in the highest potential post-application exposures; therefore, there is potential for children to be exposed via incidental oral and dermal routes from playing on treated lawns. For this assessment, the agency evaluates exposures from hand-to-mouth behavior, object-to-mouth behavior, incidental soil ingestion, and dermal contact using the 2012 Standard Operating Procedures for Residential Pesticide Exposure Assessment²⁹. Incidental oral exposures from hand-to-mouth, object-to-mouth, and incidental soil ingestion are considered inter-related and, therefore, not combined. To calculate high end estimates of exposures, the following is assumed according to the 2012 SOP to be health-protective: 1) maximum label rates are applied to the turf, 2) exposures are assumed to occur every day to the residue values on the day of application (i.e., no dissipation), and 3) individuals engage in post-application activities for the maximum amount of time represented by data for children spending time outdoors and not specifically engaged in activities

²⁹ Available: <http://www2.epa.gov/pesticide-science-and-assessing-pesticide-risks/standard-operating-procedures-residential-pesticide>

on turf, when in actuality children do not spend all of their outdoor time on turf. The highest exposure from incidental oral scenarios using the maximum application rate for turf applications would be 0.16 mg/kg/day from hand-to-mouth behaviors by children (1 to <2 years old). Dermal post-application to children 1 to <2 years old would be 0.08 mg/kg/day.

Table E.2. Post-application Exposure Estimates for Application of Glyphosate to Turf ¹ .			
Lifestage	Post-application Exposure Scenario		Exposure (mg/kg/day)
Children 1 to <2 year old	Turf – sprays	Hand-to-Mouth	0.16
		Object-to-Mouth	0.005
		Incidental Soil Ingestion	0.0003
		Dermal (high contact activities)	0.08

¹ Based on Roundup® Weed & Grass Super Concentrate, EPA Reg. No. 71995-25.

DEEM-FCID™ Chronic Residue File.

Filename: C:\Users\tbloem\Documents\work\glyphosate\registration review\417300C.R08
 Chemical: Glyphosate
 RfD(Chronic): 1 mg/kg bw/day NOEL(Chronic): 100 mg/kg bw/day
 RfD(Acute): 0 mg/kg bw/day NOEL(Acute): 0 mg/kg bw/day
 Date created/last modified: 06-09-2016/10:37:44 Program ver. 3.16, 03-08-d
 Comment: THIS R98 FILE WAS GENERATED USING THE CONVERT TO R98 UTILITY VERSION 1.1.2.

EPA Code	Crop Grp	Commodity Name	Def Res (ppm)	Adj.Factors #1	Adj.Factors #2	Comment
0101050000	1AB	Beet, garden, roots Full comment: P 7F2016	0.200000	1.000	1.000	P 7F20
0101050001	1AB	Beet, garden, roots-babyfood Full comment: P 7F2016	0.200000	1.000	1.000	P 7F20
0101052000	1A	Beet, sugar Full comment: P 7F04886	10.000000	1.000	1.000	P 7F04
0101052001	1A	Beet, sugar-babyfood Full comment: P 7F04886	10.000000	1.000	1.000	P 7F04
0101053000	1A	Beet, sugar, molasses Full comment: P 7F04886	10.000000	1.000	1.000	P 7F04
0101053001	1A	Beet, sugar, molasses-babyfood Full comment: P 7F04886	10.000000	1.000	1.000	P 7F04
0101067000	1AB	Burdock	0.200000	1.000	1.000	
0101078000	1AB	Carrot Full comment: P 8E3676 7F2016	5.000000	1.000	1.000	P 8E36
0101078001	1AB	Carrot-babyfood Full comment: P 8E3676 7F2016	5.000000	1.000	1.000	P 8E36
0101079000	1AB	Carrot, juice Full comment: P 8E3676 7F2016	5.000000	1.000	1.000	P 8E36
0101084000	1AB	Celeriac	0.200000	1.000	1.000	
0101100000	1AB	Chicory, roots	0.200000	1.000	1.000	
0101168000	1AB	Ginseng, dried Full comment: P 7F2016	0.200000	1.000	1.000	P 7F20
0101190000	1AB	Horseradish Full comment: P 8E3676	0.200000	1.000	1.000	P 8E36
0101250000	1AB	Parsley, turnip rooted	0.200000	1.000	1.000	

0101251000	1AB Parsnip	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0101251001	1AB Parsnip-babyfood	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0101314000	1AB Radish, roots	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0101316000	1AB Radish, Oriental, roots	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0101327000	1AB Rutabaga	0.200000	1.000	1.000	
0101331000	1AB Salsify, roots	0.200000	1.000	1.000	
0101388000	1AB Turnip, roots	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103015000	1CD Arrowroot, flour	0.200000	1.000	1.000	
0103015001	1CD Arrowroot, flour-babyfood	0.200000	1.000	1.000	
0103017000	1CD Artichoke, Jerusalem	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103082000	1CD Cassava	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103082001	1CD Cassava-babyfood	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103139000	1CD Dasheen, corm	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103166000	1CD Ginger	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103166001	1CD Ginger-babyfood	0.200000	1.000	1.000	
0103167000	1CD Ginger, dried	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103296000	1C Potato, chips	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103297000	1C Potato, dry (granules/ flakes)	0.200000	6.500	1.000	
0103297001	1C Potato, dry (granules/ flakes)-b	0.200000	6.500	1.000	
0103298000	1C Potato, flour	0.200000	6.500	1.000	P 7F20
	Full comment: P 7F2016				
0103298001	1C Potato, flour-babyfood	0.200000	6.500	1.000	P 7F20
	Full comment: P 7F2016				
0103299000	1C Potato, tuber, w/peel	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103299001	1C Potato, tuber, w/peel-babyfood	0.200000	1.000	1.000	
0103300000	1C Potato, tuber, w/o peel	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103300001	1C Potato, tuber, w/o peel-babyfood	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103366000	1CD Sweet potato	3.000000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103366001	1CD Sweet potato-babyfood	3.000000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103371000	1CD Tanier, corm	0.200000	1.000	1.000	
0103387000	1CD Turmeric	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103406000	1CD Yam, true	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0103407000	1CD Yam bean	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016				
0200051000	2 Beet, garden, tops	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0200101000	2 Chicory, tops	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016 & 8E2122				
0200140000	2 Dasheen, leaves	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0200315000	2 Radish, tops	0.200000	1.000	1.000	
0200317000	2 Radish, Oriental, tops	0.200000	1.000	1.000	
0200332000	2 Salsify, tops	0.200000	1.000	1.000	
0301165000	3A Garlic, bulb	0.200000	1.000	1.000	P 8E36

	Full comment: P 8E3676				
0301165001	3A Garlic, bulb-babyfood	0.200000	1.000	1.000	P 8E36
	Full comment: P 8E3676				
0301237000	3A Onion, bulb	0.200000	1.000	1.000	P 8E36
	Full comment: P 8E3676				
0301237001	3A Onion, bulb-babyfood	0.200000	1.000	1.000	P 8E36
	Full comment: P 8E3676				
0301238000	3A Onion, bulb, dried	0.200000	9.000	1.000	P 8E36
	Full comment: P 8E3676				
0301238001	3A Onion, bulb, dried-babyfood	0.200000	9.000	1.000	P 8E36
	Full comment: P 8E3676				
0301338000	3A Shallot, bulb	0.200000	1.000	1.000	
0302103000	3B Chive, fresh leaves	0.200000	1.000	1.000	P 9E60
	Full comment: P 9E6003				
0302198000	3B Leek	0.200000	1.000	1.000	P 8E36
	Full comment: P 8E3676				
0302239000	3B Onion, green	0.200000	1.000	1.000	P 8E36
	Full comment: P 8E3676				
0302338500	3B Shallot, fresh leaves	0.200000	1.000	1.000	
0401005000	4A Amaranth, leafy	0.200000	1.000	1.000	
0401018000	4A Arugula	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0401104000	4A Chrysanthemum, garland	0.200000	1.000	1.000	
0401133000	4A Cress, garden	0.200000	1.000	1.000	
0401134000	4A Cress, upland	0.200000	1.000	1.000	
0401138000	4A Dandelion, leaves	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0401150000	4A Endive	0.200000	1.000	1.000	P 7F20
	Full comment: P 7F2016 & 8E2122				
0401204000	4A Lettuce, head	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0401205000	4A Lettuce, leaf	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0401248000	4A Parsley, leaves	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0401313000	4A Radicchio	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0401355000	4A Spinach	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0401355001	4A Spinach-babyfood	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0402076000	4B Cardoon	0.200000	1.000	1.000	
0402085000	4B Celery	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0402085001	4B Celery-babyfood	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0402086000	4B Celery, juice	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0402087000	4B Celtuce	0.500000	1.000	1.000	
0402152000	4B Fennel, Florence	0.500000	1.000	1.000	
0402322000	4B Rhubarb	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0402367000	4B Swiss chard	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501061000	5A Broccoli	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501061001	5A Broccoli-babyfood	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501062000	5A Broccoli, Chinese	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501064000	5A Brussels sprouts	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501069000	5A Cabbage	0.200000	1.000	1.000	P 8E21

	Full comment: P 8E2122				
0501071000	5A Cabbage, Chinese, napa	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501072000	5A Cabbage, Chinese, mustard	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501083000	5A Cauliflower	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0501196000	5A Kohlrabi	0.500000	1.000	1.000	
0502063000	5B Broccoli raab	0.200000	1.000	1.000	
0502070000	5B Cabbage, Chinese, bok choy	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0502117000	5B Collards	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0502194000	5B Kale	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0502229000	5B Mustard greens	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0502318000	5B Rape greens	0.200000	1.000	1.000	
0502389000	5B Turnip, greens	0.200000	1.000	1.000	P 8E21
	Full comment: P 8E2122				
0600347000	6 Soybean, seed	20.000000	1.000	1.000	P 5F15
	Full comment: P 5F1536				
0600349000	6 Soybean, soy milk	20.000000	1.000	1.000	P 5F15
	Full comment: P 5F1536				
0600349001	6 Soybean, soy milk-babyfood or in	20.000000	1.000	1.000	
0600350000	6 Soybean, oil	20.000000	1.000	1.000	P 5F15
	Full comment: P 5F1536				
0600350001	6 Soybean, oil-babyfood	20.000000	1.000	1.000	P 5F15
	Full comment: P 5F1536				
0601043000	6A Bean, snap, succulent	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0601043001	6A Bean, snap, succulent-babyfood	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0601257000	6A Pea, edible podded, succulent	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0601349500	6AB Soybean, vegetable	5.000000	1.000	1.000	
0602031000	6B Bean, broad, succulent	5.000000	1.000	1.000	
0602033000	6B Bean, cowpea, succulent	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0602037000	6B Bean, lima, succulent	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0602255000	6B Pea, succulent	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0602255001	6B Pea, succulent-babyfood	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0602259000	6B Pea, pigeon, succulent	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603030000	6C Bean, black, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603032000	6C Bean, broad, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603034000	6C Bean, cowpea, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603035000	6C Bean, great northern, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603036000	6C Bean, kidney, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603038000	6C Bean, lima, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603039000	6C Bean, mung, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				
0603040000	6C Bean, navy, seed	5.000000	1.000	1.000	P 2E41
	Full comment: P 2E4118				

0603041000	6C	Bean, pink, seed	5.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603042000	6C	Bean, pinto, seed	5.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603098000	6C	Chickpea, seed	8.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603098001	6C	Chickpea, seed-babyfood	8.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603099000	6C	Chickpea, flour	8.000000	1.000	1.000	
0603182000	6C	Guar, seed	8.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603182001	6C	Guar, seed-babyfood	8.000000	1.000	1.000	
0603203000	6C	Lentil, seed	8.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603256000	6C	Pea, dry	8.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603256001	6C	Pea, dry-babyfood	8.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
0603258000	6C	Pea, pigeon, seed	8.000000	1.000	1.000	
0603348000	6C	Soybean, flour	20.000000	1.000	1.000	P 5F15
		Full comment: P 5F1536				
0603348001	6C	Soybean, flour-babyfood	20.000000	1.000	1.000	P 5F15
		Full comment: P 5F1536				
0801374000	8A	Tomatillo	0.100000	1.000	1.000	
0801375000	8A	Tomato	0.100000	1.000	1.000	
0801375001	8A	Tomato-babyfood	0.100000	1.000	1.000	
0801376000	8A	Tomato, paste	0.100000	5.400	1.000	
0801376001	8A	Tomato, paste-babyfood	0.100000	5.400	1.000	
0801377000	8A	Tomato, puree	0.100000	3.300	1.000	
0801377001	8A	Tomato, puree-babyfood	0.100000	3.300	1.000	
0801378000	8A	Tomato, dried	0.100000	14.300	1.000	
0801378001	8A	Tomato, dried-babyfood	0.100000	14.300	1.000	
0801379000	8A	Tomato, juice	0.100000	1.500	1.000	
0801380000	8A	Tomato, Tree	0.100000	1.000	1.000	
0802148000	8BC	Eggplant	0.100000	1.000	1.000	
0802234000	8BC	Okra	0.500000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
0802270000	8B	Pepper, bell	0.100000	1.000	1.000	
0802270001	8B	Pepper, bell-babyfood	0.100000	1.000	1.000	
0802271000	8B	Pepper, bell, dried	0.100000	1.000	1.000	
0802271001	8B	Pepper, bell, dried-babyfood	0.100000	1.000	1.000	
0802272000	8BC	Pepper, nonbell	0.100000	1.000	1.000	
0802272001	8BC	Pepper, nonbell-babyfood	0.100000	1.000	1.000	
0802273000	8BC	Pepper, nonbell, dried	0.100000	1.000	1.000	
0901075000	9A	Cantaloupe	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
0901187000	9A	Honeydew melon	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
0901399000	9A	Watermelon	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
0901400000	9A	Watermelon, juice	0.500000	1.000	1.000	
0902021000	9B	Balsam pear	0.500000	1.000	1.000	
0902088000	9B	Chayote, fruit	0.500000	1.000	1.000	
0902102000	9B	Chinese waxgourd	0.500000	1.000	1.000	
0902135000	9B	Cucumber	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
0902308000	9B	Pumpkin	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
0902309000	9B	Pumpkin, seed	0.500000	1.000	1.000	
0902356000	9B	Squash, summer	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
0902356001	9B	Squash, summer-babyfood	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				

0902357000	9B	Squash, winter	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
0902357001	9B	Squash, winter-babyfood	0.500000	1.000	1.000	P 3E28
		Full comment: P 3E2845				
1001106000	10A	Citron	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1001107000	10A	Citrus hybrids	0.500000	1.000	1.000	
1001108000	10A	Citrus, oil	0.500000	1.000	1.000	
1001240000	10A	Orange	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1001241000	10A	Orange, juice	0.500000	1.800	1.000	P 4F43
		Full comment: P 4F4338				
1001241001	10A	Orange, juice-babyfood	0.500000	1.800	1.000	P 4F43
		Full comment: P 4F4338				
1001242000	10A	Orange, peel	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1001369000	10A	Tangerine	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1001370000	10A	Tangerine, juice	0.500000	2.300	1.000	P 4F43
		Full comment: P 4F4338				
1002197000	10B	Kumquat	0.500000	1.000	1.000	
1002199000	10B	Lemon	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1002200000	10B	Lemon, juice	0.500000	2.000	1.000	P 4F43
		Full comment: P 4F4338				
1002200001	10B	Lemon, juice-babyfood	0.500000	2.000	1.000	P 4F43
		Full comment: P 4F4338				
1002201000	10B	Lemon, peel	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1002206000	10B	Lime	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1002207000	10B	Lime, juice	0.500000	2.000	1.000	P 4F43
		Full comment: P 4F4338				
1002207001	10B	Lime, juice-babyfood	0.500000	2.000	1.000	P 4F43
		Full comment: P 4F4338				
1003180000	10C	Grapefruit	0.500000	1.000	1.000	P 4F43
		Full comment: P 4F4338				
1003181000	10C	Grapefruit, juice	0.500000	2.100	1.000	P 4F43
		Full comment: P 4F4338				
1003307000	10C	Pummelo	0.500000	1.000	1.000	
1100007000	11	Apple, fruit with peel	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100008000	11	Apple, peeled fruit	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100008001	11	Apple, peeled fruit-babyfood	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100009000	11	Apple, dried	0.200000	8.000	1.000	P 6F18
		Full comment: P 6F1861				
1100009001	11	Apple, dried-babyfood	0.200000	8.000	1.000	P 6F18
		Full comment: P 6F1861				
1100010000	11	Apple, juice	0.200000	1.300	1.000	P 6F18
		Full comment: P 6F1861				
1100010001	11	Apple, juice-babyfood	0.200000	1.300	1.000	P 6F18
		Full comment: P 6F1861				
1100011000	11	Apple, sauce	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100011001	11	Apple, sauce-babyfood	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100129000	11	Crabapple	0.200000	1.000	1.000	
1100173500	11	Goji berry	0.100000	1.000	1.000	
1100210000	11	Loquat	0.200000	1.000	1.000	
1100266000	11	Pear	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				

1100266001	11	Pear-babyfood	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100267000	11	Pear, dried	0.200000	6.250	1.000	P 6F18
		Full comment: P 6F1861				
1100268000	11	Pear, juice	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100268001	11	Pear, juice-babyfood	0.200000	1.000	1.000	P 6F18
		Full comment: P 6F1861				
1100310000	11	Quince	0.200000	1.000	1.000	
1201090000	12A	Cherry	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1201090001	12A	Cherry-babyfood	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1201091000	12A	Cherry, juice	0.200000	1.500	1.000	P 2600
		Full comment: P 260044				
1201091001	12A	Cherry, juice-babyfood	0.200000	1.500	1.000	P 2600
		Full comment: P 260044				
1202012000	12B	Apricot	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202012001	12B	Apricot-babyfood	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202013000	12B	Apricot, dried	0.200000	6.000	1.000	P 2600
		Full comment: P 260044				
1202014000	12B	Apricot, juice	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202014001	12B	Apricot, juice-babyfood	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202230000	12B	Nectarine	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202260000	12B	Peach	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202260001	12B	Peach-babyfood	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202261000	12B	Peach, dried	0.200000	7.000	1.000	P 2600
		Full comment: P 260044				
1202261001	12B	Peach, dried-babyfood	0.200000	7.000	1.000	
1202262000	12B	Peach, juice	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1202262001	12B	Peach, juice-babyfood	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1203285000	12C	Plum	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1203285001	12C	Plum-babyfood	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1203286000	12C	Plum, prune, fresh	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1203286001	12C	Plum, prune, fresh-babyfood	0.200000	1.000	1.000	P 2600
		Full comment: P 260044				
1203287000	12C	Plum, prune, dried	0.200000	5.000	1.000	P 2600
		Full comment: P 260044				
1203287001	12C	Plum, prune, dried-babyfood	0.200000	5.000	1.000	
1203288000	12C	Plum, prune, juice	0.200000	1.400	1.000	P 2600
		Full comment: P 260044				
1203288001	12C	Plum, prune, juice-babyfood	0.200000	1.400	1.000	P 2600
		Full comment: P 260044				
1301055000	13A	Blackberry	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1301056000	13A	Blackberry, juice	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1301056001	13A	Blackberry, juice-babyfood	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1301058000	13A	Boysenberry	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				

1301208000	13A	Loganberry	0.200000	1.000	1.000	
1301320000	13A	Raspberry	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1301320001	13A	Raspberry-babyfood	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1301321000	13A	Raspberry, juice	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1301321001	13A	Raspberry, juice-babyfood	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1302057000	13B	Blueberry	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1302057001	13B	Blueberry-babyfood	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1302136000	13B	Currant	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1302137000	13B	Currant, dried	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1302149000	13B	Elderberry	0.200000	1.000	1.000	
1302174000	13B	Gooseberry	0.200000	1.000	1.000	
1302191000	13B	Huckleberry	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1303227000	13C	Mulberry	0.200000	1.000	1.000	
1304175000	13D	Grape	0.200000	1.000	1.000	P 5F15
		Full comment: P 5F1560				
1304176000	13D	Grape, juice	0.200000	1.200	1.000	P 5F15
		Full comment: P 5F1560				
1304176001	13D	Grape, juice-babyfood	0.200000	1.200	1.000	P 5F15
		Full comment: P 5F1560				
1304179000	13D	Grape, wine and sherry	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1304195000	13D	Kiwifruit, fuzzy	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2929				
1307130000	13G	Cranberry	0.200000	1.000	1.000	P 0E24
		Full comment: P 0E2421				
1307130001	13G	Cranberry-babyfood	0.200000	1.000	1.000	P 0E24
		Full comment: P 0E2421				
1307131000	13G	Cranberry, dried	0.200000	1.000	1.000	P 0E24
		Full comment: P 0E2421				
1307132000	13G	Cranberry, juice	0.200000	1.100	1.000	P 0E24
		Full comment: P 0E2421				
1307132001	13G	Cranberry, juice-babyfood	0.200000	1.100	1.000	P 0E24
		Full comment: P 0E2421				
1307359000	13G	Strawberry	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1307359001	13G	Strawberry-babyfood	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1307360000	13G	Strawberry, juice	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1307360001	13G	Strawberry, juice-babyfood	0.200000	1.000	1.000	P 3E29
		Full comment: P 3E2930				
1400003000	14	Almond	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1400003001	14	Almond-babyfood	1.000000	1.000	1.000	
1400004000	14	Almond, oil	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1400004001	14	Almond, oil-babyfood	1.000000	1.000	1.000	
1400059000	14	Brazil nut	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1400068000	14	Butternut	1.000000	1.000	1.000	
1400081000	14	Cashew	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1400092000	14	Chestnut	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				

1400155000	14	Hazelnut	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1400156000	14	Hazelnut, oil	1.000000	1.000	1.000	
1400185000	14	Hickory nut	1.000000	1.000	1.000	
1400213000	14	Macadamia nut	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1400269000	14	Pecan	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1400278000	14	Pine nut	1.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1400282000	14	Pistachio	1.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1400391000	14	Walnut	1.000000	1.000	1.000	P 7F18
		Full comment: P 7F1893				
1500025000	15	Barley, pearled barley	30.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
1500025001	15	Barley, pearled barley-babyfood	30.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
1500026000	15	Barley, flour	30.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
1500026001	15	Barley, flour-babyfood	30.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
1500027000	15	Barley, bran	30.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
1500065000	15	Buckwheat	30.000000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500066000	15	Buckwheat, flour	30.000000	1.000	1.000	
1500120000	15	Corn, field, flour	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500120001	15	Corn, field, flour-babyfood	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500121000	15	Corn, field, meal	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500121001	15	Corn, field, meal-babyfood	5.000000	1.000	1.000	
1500122000	15	Corn, field, bran	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500123000	15	Corn, field, starch	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500123001	15	Corn, field, starch-babyfood	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500124000	15	Corn, field, syrup	5.000000	1.500	1.000	P 8F36
		Full comment: P 8F3673				
1500124001	15	Corn, field, syrup-babyfood	5.000000	1.500	1.000	P 8F36
		Full comment: P 8F3673				
1500125000	15	Corn, field, oil	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500125001	15	Corn, field, oil-babyfood	5.000000	1.000	1.000	P 8F36
		Full comment: P 8F3673				
1500126000	15	Corn, pop	0.100000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500127000	15	Corn, sweet	3.500000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500127001	15	Corn, sweet-babyfood	3.500000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500226000	15	Millet, grain	30.000000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500231000	15	Oat, bran	30.000000	1.000	1.000	P 6E46
		Full comment: P 6E4645				
1500232000	15	Oat, flour	30.000000	1.000	1.000	P 6E46
		Full comment: P 6E4645				
1500232001	15	Oat, flour-babyfood	30.000000	1.000	1.000	P 6E46
		Full comment: P 6E4645				
1500233000	15	Oat, groats/rolled oats	30.000000	1.000	1.000	P 6E46

		Full comment: P 6E4645				
1500233001	15	Oat, groats/rolled oats-babyfood	30.000000	1.000	1.000	P 6E46
		Full comment: P 6E4645				
1500323000	15	Rice, white	0.100000	1.000	1.000	
1500323001	15	Rice, white-babyfood	0.100000	1.000	1.000	
1500324000	15	Rice, brown	0.100000	1.000	1.000	
1500324001	15	Rice, brown-babyfood	0.100000	1.000	1.000	
1500325000	15	Rice, flour	0.100000	1.000	1.000	
1500325001	15	Rice, flour-babyfood	0.100000	1.000	1.000	
1500326000	15	Rice, bran	0.100000	1.000	1.000	
1500326001	15	Rice, bran-babyfood	0.100000	1.000	1.000	
1500328000	15	Rye, grain	30.000000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500329000	15	Rye, flour	30.000000	1.000	1.000	
1500344000	15	Sorghum, grain	30.000000	1.000	1.000	
1500345000	15	Sorghum, syrup	30.000000	1.000	1.000	
1500381000	15	Triticale, flour	30.000000	1.000	1.000	
1500381001	15	Triticale, flour-babyfood	30.000000	1.000	1.000	
1500401000	15	Wheat, grain	30.000000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500401001	15	Wheat, grain-babyfood	30.000000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500402000	15	Wheat, flour	30.000000	1.000	1.000	
1500402001	15	Wheat, flour-babyfood	30.000000	1.000	1.000	
1500403000	15	Wheat, germ	30.000000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500404000	15	Wheat, bran	30.000000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1500405000	15	Wild rice	0.100000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1800002000	18	Alfalfa, seed	0.500000	1.000	1.000	
1901028000	19A	Basil, fresh leaves	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1901028001	19A	Basil, fresh leaves-babyfood	0.200000	1.000	1.000	
1901029000	19A	Basil, dried leaves	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1901029001	19A	Basil, dried leaves-babyfood	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1901102500	19A	Chive, dried leaves	0.200000	1.000	1.000	
1901118000	19A	Cilantro, leaves	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1901118001	19A	Cilantro, leaves-babyfood	0.200000	1.000	1.000	
1901144000	19A	Dillweed	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1901184000	19A	Herbs, other	0.200000	1.000	1.000	
1901184001	19A	Herbs, other-babyfood	0.200000	1.000	1.000	
1901202000	19A	Lemongrass	0.200000	1.000	1.000	
1901220000	19A	Marjoram	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1901220001	19A	Marjoram-babyfood	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1901249000	19A	Parsley, dried leaves	0.200000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1901249001	19A	Parsley, dried leaves-babyfood	0.200000	1.000	1.000	P 8E21
		Full comment: P 8E2122				
1901334000	19A	Savory	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1902105000	19B	Cinnamon	7.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1902105001	19B	Cinnamon-babyfood	7.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1902119000	19B	Coriander, seed	7.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				

1902119001	19B	Coriander, seed-babyfood	7.000000	1.000	1.000	
1902143000	19B	Dill, seed	7.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1902274000	19B	Pepper, black and white	7.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1902274001	19B	Pepper, black and white-babyfood	7.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
1902354000	19B	Spices, other	7.000000	1.000	1.000	
1902354001	19B	Spices, other-babyfood	7.000000	1.000	1.000	
2001163000	20A	Flax seed, oil	40.000000	1.000	1.000	00ND00
		Full comment: 00ND0025 (S18)				
2001319000	20A	Rapeseed, oil	20.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
2001319001	20A	Rapeseed, oil-babyfood	20.000000	1.000	1.000	P 2E41
		Full comment: P 2E4118				
2001336000	20A	Sesame, seed	40.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
2001336001	20A	Sesame, seed-babyfood	40.000000	1.000	1.000	
2001337000	20A	Sesame, oil	40.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
2001337001	20A	Sesame, oil-babyfood	40.000000	1.000	1.000	
2002330000	20B	Safflower, oil	40.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
2002330001	20B	Safflower, oil-babyfood	40.000000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
2002364000	20B	Sunflower, seed	40.000000	1.000	1.000	P 6F34
		Full comment: P 6F3408				
2002365000	20B	Sunflower, oil	40.000000	1.000	1.000	P 6F34
		Full comment: P 6F3408				
2002365001	20B	Sunflower, oil-babyfood	40.000000	1.000	1.000	P 6F34
		Full comment: P 6F3408				
2003114001	20C	Coconut, oil-babyfood	0.100000	1.000	1.000	P 2F26
		Full comment: P 2F2680				
2003128000	20C	Cottonseed, oil	40.000000	1.000	1.000	
2003128001	20C	Cottonseed, oil-babyfood	40.000000	1.000	1.000	
3100046000	31	Beef, meat byproducts	5.000000	1.000	1.000	
3100046001	31	Beef, meat byproducts-babyfood	5.000000	1.000	1.000	
3100048000	31	Beef, kidney	5.000000	1.000	1.000	P 4F43
		Full comment: P 4F4312				
3100049000	31	Beef, liver	5.000000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
3100049001	31	Beef, liver-babyfood	5.000000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
3200170000	32	Goat, meat byproducts	5.000000	1.000	1.000	
3200172000	32	Goat, kidney	5.000000	1.000	1.000	
3200173000	32	Goat, liver	5.000000	1.000	1.000	
3400291000	34	Pork, skin	5.000000	1.000	1.000	
3400292000	34	Pork, meat byproducts	5.000000	1.000	1.000	
3400292001	34	Pork, meat byproducts-babyfood	5.000000	1.000	1.000	
3400294000	34	Pork, kidney	5.000000	1.000	1.000	
3400295000	34	Pork, liver	5.000000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
3500340000	35	Sheep, meat byproducts	5.000000	1.000	1.000	
3500342000	35	Sheep, kidney	5.000000	1.000	1.000	
3500343000	35	Sheep, liver	5.000000	1.000	1.000	
4000093000	40	Chicken, meat	0.100000	1.000	1.000	P 9F50
		Full comment: P 9F5096				
4000093001	40	Chicken, meat-babyfood	0.100000	1.000	1.000	P 9F50
		Full comment: P 9F5096				
4000094000	40	Chicken, liver	1.000000	1.000	1.000	P 9F50
		Full comment: P 9F5096				
4000095000	40	Chicken, meat byproducts	1.000000	1.000	1.000	P 9F50
		Full comment: P 9F5096				

4000095001	40	Chicken, meat byproducts-babyfoo	1.000000	1.000	1.000	P 9F50
		Full comment: P 9F5096				
4000097000	40	Chicken, skin	1.000000	1.000	1.000	
4000097001	40	Chicken, skin-babyfood	1.000000	1.000	1.000	
5000382000	50	Turkey, meat	0.100000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
5000382001	50	Turkey, meat-babyfood	0.100000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
5000383000	50	Turkey, liver	1.000000	1.000	1.000	
5000383001	50	Turkey, liver-babyfood	1.000000	1.000	1.000	
5000384000	50	Turkey, meat byproducts	1.000000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
5000384001	50	Turkey, meat byproducts-babyfood	1.000000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
5000386000	50	Turkey, skin	1.000000	1.000	1.000	
5000386001	50	Turkey, skin-babyfood	1.000000	1.000	1.000	
6000301000	60	Poultry, other, meat	0.100000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
6000302000	60	Poultry, other, liver	1.000000	1.000	1.000	
6000303000	60	Poultry, other, meat byproducts	1.000000	1.000	1.000	
6000305000	60	Poultry, other, skin	1.000000	1.000	1.000	
7000145000	70	Egg, whole	0.050000	1.000	1.000	P 9F5
		Full comment: P 9F5096				
7000145001	70	Egg, whole-babyfood	0.050000	1.000	1.000	P 9F5
		Full comment: P 9F5096				
7000146000	70	Egg, white	0.050000	1.000	1.000	P 9F5
		Full comment: P 9F5096				
7000146001	70	Egg, white (solids)-babyfood	0.050000	1.000	1.000	
7000147000	70	Egg, yolk	0.050000	1.000	1.000	P 9F5
		Full comment: P 9F5096				
7000147001	70	Egg, yolk-babyfood	0.050000	1.000	1.000	P 9F5
		Full comment: P 9F5096				
8000157000	80	Fish-freshwater finfish	0.250000	1.000	1.000	P 9F21
		Full comment: P 9F2163				
8000158000	80	Fish-freshwater finfish, farm ra	0.250000	1.000	1.000	P 9F21
		Full comment: P 9F2163				
8000159000	80	Fish-saltwater finfish, tuna	0.250000	1.000	1.000	P 9F21
		Full comment: P 9F2163				
8000160000	80	Fish-saltwater finfish, other	0.250000	1.000	1.000	P 9F21
		Full comment: P 9F2163				
8000161000	80	Fish-shellfish, crustacean	3.000000	1.000	1.000	P 3F29
		Full comment: P 3F2956				
8000162000	80	Fish-shellfish, mollusc	3.000000	1.000	1.000	P 3F29
		Full comment: P 3F2956				
8601000000	86A	Water, direct, all sources	0.159000	1.000	1.000	
8602000000	86B	Water, indirect, all sources	0.159000	1.000	1.000	
9500000500	0	Acai berry	0.200000	1.000	1.000	
9500001000	0	Acerola	0.200000	1.000	1.000	
9500001500	0	Agave	0.500000	1.000	1.000	
9500016000	0	Artichoke, globe	0.200000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
9500019000	0	Asparagus	0.500000	1.000	1.000	P 8E36
		Full comment: P 8E3648				
9500019500	0	Atemoya	0.200000	1.000	1.000	
9500020000	0	Avocado	0.200000	1.000	1.000	P 8F20
		Full comment: P 8F2021				
9500022000	0	Bamboo, shoots	0.500000	1.000	1.000	P 9E60
		Full comment: P 9E6003				
9500023000	0	Banana	0.200000	1.000	1.000	P 9F22
		Full comment: P 9F2223				
9500023001	0	Banana-babyfood	0.200000	1.000	1.000	P 9F22
		Full comment: P 9F2223				
9500024000	0	Banana, dried	0.200000	3.900	1.000	P 9F22

	Full comment: P 9F2223				
9500024001	0 Banana, dried-babyfood	0.200000	3.900	1.000	P 9F22
	Full comment: P 9F2223				
9500060000	0 Breadfruit	0.200000	1.000	1.000	P 9E37
	Full comment: P 9E3754				
9500073000	0 Cactus	0.500000	1.000	1.000	
9500074000	0 Canistel	0.200000	1.000	1.000	
9500077000	0 Carob	0.200000	1.000	1.000	
9500089000	0 Cherimoya	0.200000	1.000	1.000	
9500109000	0 Cocoa bean, chocolate	0.200000	1.000	1.000	P 0E38
	Full comment: P 0E3857				
9500110000	0 Cocoa bean, powder	0.200000	1.000	1.000	P 0E38
	Full comment: P 0E3857				
9500111000	0 Coconut, meat	0.100000	1.000	1.000	P 2F26
	Full comment: P 2F2680				
9500111001	0 Coconut, meat-babyfood	0.100000	1.000	1.000	P 2F26
	Full comment: P 2F2680				
9500112000	0 Coconut, dried	0.100000	2.100	1.000	P 2F26
	Full comment: P 2F2680				
9500113000	0 Coconut, milk	0.100000	1.000	1.000	P 2F26
	Full comment: P 2F2680				
9500114000	0 Coconut, oil	0.100000	1.000	1.000	P 2F26
	Full comment: P 2F2680				
9500115000	0 Coffee, roasted bean	1.000000	1.000	1.000	P 6E18
	Full comment: P 6E1809				
9500116000	0 Coffee, instant	1.000000	1.000	1.000	P 6E18
	Full comment: P 6E1809				
9500141000	0 Date	0.200000	1.000	1.000	P 9E37
	Full comment: P 9E3754				
9500151000	0 Feijoa	0.200000	1.000	1.000	
9500153000	0 Fig	0.200000	1.000	1.000	P 3E29
	Full comment: P 3E2929				
9500154000	0 Fig, dried	0.200000	1.000	1.000	P 3E29
	Full comment: P 3E2929				
9500177000	0 Grape, leaves	0.200000	1.000	1.000	
9500178000	0 Grape, raisin	0.200000	4.300	1.000	P 5F15
	Full comment: P 5F1560				
9500183000	0 Guava	0.200000	1.000	1.000	P 1E24
	Full comment: P 1E2443				
9500183001	0 Guava-babyfood	0.200000	1.000	1.000	
9500188000	0 Hop	7.000000	1.000	1.000	
9500193000	0 Jackfruit	0.200000	1.000	1.000	
9500209000	0 Longan	0.200000	1.000	1.000	
9500211000	0 Lychee	0.200000	1.000	1.000	
9500212000	0 Lychee, dried	0.200000	1.850	1.000	
9500214000	0 Mamey apple	0.200000	1.000	1.000	
9500215000	0 Mango	0.200000	1.000	1.000	P 1E24
	Full comment: P 1E2490				
9500215001	0 Mango-babyfood	0.200000	1.000	1.000	P 1E24
	Full comment: P 1E2490				
9500216000	0 Mango, dried	0.200000	1.000	1.000	P 1E24
	Full comment: P 1E2490				
9500217000	0 Mango, juice	0.200000	1.000	1.000	P 1E24
	Full comment: P 1E2490				
9500217001	0 Mango, juice-babyfood	0.200000	1.000	1.000	P 1E24
	Full comment: P 1E2490				
9500235000	0 Olive	0.200000	1.000	1.000	P 3E29
	Full comment: P 3E2929				
9500236000	0 Olive, oil	0.200000	1.000	1.000	P 3E29
	Full comment: P 3E2929				
9500243000	0 Palm heart, leaves	0.500000	1.000	1.000	P 9E60
	Full comment: P 9E6003				
9500244000	0 Palm, oil	0.100000	1.000	1.000	P 6H51

		Full comment: P 6H5115				
9500244001	0	Palm, oil-babyfood	0.100000	1.000	1.000	P 6H51
		Full comment: P 6H5115				
9500245000	0	Papaya	0.200000	1.000	1.000	P 1E24
		Full comment: P 1E2443				
9500245001	0	Papaya-babyfood	0.200000	1.000	1.000	
9500246000	0	Papaya, dried	0.200000	1.800	1.000	P 1E24
		Full comment: P 1E2443				
9500247000	0	Papaya, juice	0.200000	1.500	1.000	P 1E24
		Full comment: P 1E2443				
9500252000	0	Passionfruit	0.200000	1.000	1.000	P 9E37
		Full comment: P 9E3715				
9500252001	0	Passionfruit-babyfood	0.200000	1.000	1.000	
9500253000	0	Passionfruit, juice	0.200000	1.000	1.000	P 9E37
		Full comment: P 9E3715				
9500253001	0	Passionfruit, juice-babyfood	0.200000	1.000	1.000	
9500254000	0	Pawpaw	0.200000	1.000	1.000	
9500263000	0	Peanut	0.100000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
9500264000	0	Peanut, butter	0.100000	1.890	1.000	
9500265000	0	Peanut, oil	0.100000	1.000	1.000	P 0F23
		Full comment: P 0F2329				
9500275000	0	Peppermint	200.000000	1.000	1.000	
9500276000	0	Peppermint, oil	200.000000	1.000	1.000	
9500277000	0	Persimmon	0.200000	1.000	1.000	P 9E37
		Full comment: P 9E3754				
9500279000	0	Pineapple	0.200000	1.000	1.000	P 2F26
		Full comment: P 2F2634				
9500279001	0	Pineapple-babyfood	0.200000	1.000	1.000	P 2F26
		Full comment: P 2F2634				
9500280000	0	Pineapple, dried	0.200000	5.000	1.000	P 2F26
		Full comment: P 2F2634				
9500281000	0	Pineapple, juice	0.200000	1.700	1.000	P 2F26
		Full comment: P 2F2634				
9500281001	0	Pineapple, juice-babyfood	0.200000	1.700	1.000	P 2F26
		Full comment: P 2F2634				
9500283000	0	Plantain	0.200000	1.000	1.000	P 9F22
		Full comment: P 9F2223				
9500284000	0	Plantain, dried	0.200000	3.900	1.000	P 9F22
		Full comment: P 9F2223				
9500289000	0	Pomegranate	0.200000	1.000	1.000	P 1E39
		Full comment: P 1E3978				
9500311000	0	Quinoa, grain	5.000000	1.000	1.000	
9500333000	0	Sapote, Mamey	0.200000	1.000	1.000	
9500346000	0	Soursop	0.200000	1.000	1.000	
9500351000	0	Spanish lime	0.200000	1.000	1.000	
9500352000	0	Spearmint	200.000000	1.000	1.000	
9500353000	0	Spearmint, oil	200.000000	1.000	1.000	
9500358000	0	Starfruit	0.200000	1.000	1.000	P 6E34
		Full comment: P 6E3424				
9500361000	0	Sugar apple	0.200000	1.000	1.000	
9500362000	0	Sugarcane, sugar	2.000000	1.000	1.000	
9500362001	0	Sugarcane, sugar-babyfood	2.000000	1.000	1.000	
9500363000	0	Sugarcane, molasses	30.000000	1.000	1.000	P 9H51
		Full comment: P 9H5196				
9500363001	0	Sugarcane, molasses-babyfood	30.000000	1.000	1.000	P 9H51
		Full comment: P 9H5196				
9500368000	0	Tamarind	0.200000	1.000	1.000	
9500372000	0	Tea, dried	1.000000	1.000	1.000	P 1H53
		Full comment: P 1H5310 & 8H5568				
9500373000	0	Tea, instant	7.000000	1.000	1.000	P 1H53
		Full comment: P 1H5310 & 8H5568				
9500373500	0	Teff, flour	5.000000	1.000	1.000	

9500398000 0 Watercress
Full comment: P 9E6003

0.200000 1.000 1.000 P 9E60

Attachment 2: DEEM-FCID™ Chronic Exposure Estimates.

US EPA Ver. 3.16, 03-08-d
DEEM-FCID Chronic analysis for GLYPHOSATE NHANES 2003-2008 2-day
Residue file name: C:\Users\tbloem\Documents\work\glyphosate\registration
review\417300C.R08

Adjustment factor #2 NOT used.

Analysis Date 06-09-2016/10:40:23

Residue file dated: 06-09-2016/10:37:44

COMMENT 1: THIS R98 FILE WAS GENERATED USING THE CONVERT TO R98 UTILITY VERSION 1.1.2.

=====

Total exposure by population subgroup

Population Subgroup	Total Exposure mg/kg body wt/day
Total US Population	0.091530
Hispanic	0.094838
Non-Hisp-White	0.091452
Non-Hisp-Black	0.086606
Non-Hisp-Other	0.095659
Nursing Infants	0.072309
Non-Nursing Infants	0.174388
Female 13+ PREG	0.076716
Children 1-6	0.218895
Children 7-12	0.139417
Male 13-19	0.097324
Female 13-19/NP	0.082295
Male 20+	0.077524
Female 20+/NP	0.064402
Seniors 55+	0.061294
All Infants	0.142873
Female 13-50	0.070729
Children 1-2	0.230916
Children 3-5	0.214174
Children 6-12	0.149290
Youth 13-19	0.089645
Adults 20-49	0.076405
Adults 50-99	0.062993
Female 13-49	0.071066

Note: The reference dose (RfD) and percent of RfD have been removed from this file because these are based on non-cancer endpoints and non-cancer endpoints are not the focus of this SAP.

Appendix F

Genotoxicity Studies with Glyphosate Based Formulations

While the focus of this analysis to determine the genotoxic potential of glyphosate, the agency has identified numerous studies conducted with glyphosate-based formulations that contain various concentrations of the glyphosate as well as other components of the end use products and are presented in Tables F.1-F.5.

Table F.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Formulations.						
Test/Endpoint	Test System	Concentrations	Test Material/ Concentration	Results	Reference	Comments
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA1537, TA1538, TA98 and TA100; <i>E. coli</i> WP2 <i>uvrA</i> pKM101 $\pm$ S9	1.6-5000 μ g/plate $\pm$ S9 (plate incorporation)	ICIA 0224 57.6% in water	Negative $\pm$ S9	Callander (1988)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537; <i>E. coli</i> WP2P and <i>uvrA</i> $\pm$ S9	100-5000 μ g/plate $\pm$ S9 plate incorporation & pre-incubation protocols	TMSC (tri-methyl-sulfonium chloride) 95% purity	Negative $\pm$ S9	Callander (1993)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 $\pm$ S9	26, 43, 72, 120, 200 μ g/plate	Glyphosate liquid formulation (480 g/L isopropylamine salt)	Negative $\pm$ S9	Camolesi (2009) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 $\pm$ S9	26, 43, 72, 120, 200 μ g/plate	MON 77280 equivalent of glyphosate acid: 495 g/L	Negative $\pm$ S9	Camolesi (2010)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 $\pm$ S9	0.2-2000 μ g/plate	MON 76190 53.2% glyphosate	Negative $\pm$ S9	Catoyra (2009) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA97a, TA98, TA100 and TA102 $\pm$ S9	2 μ g/plate (toxic)	Perzocyd 10 SL formulation	Negative $\pm$ S9	Chruscielska <i>et al.</i> (2000)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 $\pm$ S9	0.03-3.0 μ L/plate	MON 8080 (87.6%)	Negative $\pm$ S9	Flowers (1981)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 $\pm$ S9	3.16-1000 μ g/plate	TROP M (Glyphosate 480); 35.84% purity based on acid, 48.46% pure based on IPA salt	Negative $\pm$ S9	Flügge (2010a) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 $\pm$ S9	0.316-100	Glyphosate 757 g/kg granular formulation (76.1%)	Negative $\pm$ S9	Flügge (2010d) ¹	

Table F.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Formulations.						
Test/Endpoint	Test System	Concentrations	Test Material/ Concentration	Results	Reference	Comments
			monoammonium glyphosate salt)			
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA97a, TA98, TA100, and TA1535 ± S9	1-5000 µg/plate	Roundup WG 784 g/kg ammonium salt equivalent	Negative ± S9	Gava (1998)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537± S9	50-5000 µg/plate	Rodeo® (containing IPA salt and water only); 40% glyphosate (acid equivalent)	Negative ± S9	Kier <i>et al.</i> , (1992)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 ± S9	5-500 µg/plate (-S9)/ 15-1500 µg/plate (+S9)	MON 2139 (Roundup®) 31% Glyphosate (acid equivalent)	Negative ± S9	Kier <i>et al.</i> , (1992)	Cytotoxic at top concentrations
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 ± S9	5-500 µg/plate (-S9)/ 15-1500 µg/plate (+S9)	MON 14445 (Direct®); 75% Glyphosate (acid equivalent)	Negative ± S9	Kier <i>et al.</i> , (1992)	Cytotoxic at the top concentrations, occasionally at lower concentrations
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 ± S9	0.2-2000 µg/plate	MON 79672 (Roundup Ultramax); 74.7% monoammonium glyphosate salt; 68.2% glyphosate	Negative ± S9	Lope (2008) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA1535, TA1537, TA1538, TA98 and TA100 ± S9	0.617-50 µL/plate ± S9	SC-0224, 19.2% purity	Negative ± S9	Majeska (1982)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	TA strains: 10 - 5000 µg/plate (+S9); 3.33-3330 µg/plate (-S9); <i>E. coli</i> : 33.3-5000 µg/plate (+/- S9)	MON 78239 36.6% glyphosate (44.9% potassium salt of glyphosate)	Negative	Mecchi (2003a)	Increase in revertants seen in TA98 and TA1535 -S9 on first trial, not conc-dep; however no increase in revertants seen in repeat in those strains; overall negative.

Table F.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Formulations.						
Test/Endpoint	Test System	Concentrations	Test Material/ Concentration	Results	Reference	Comments
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	TA strains: 3.33-3330 µg/plate (+S9); 1.0-1000 µg/plate (-S9); <i>E. coli</i> : 33.3-5000 µg/plate (+/- S9)	MON 78634 65.2% w/w glyphosate (71.8% w/w as monoammonium salt of glyphosate)	Negative	Mecchi (2003b)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA 98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	10 - 5000 µg/plate (+/-S9)	MON 79864 38.7% glyphosate acid (wt %)	Negative	Mecchi (2008a)	Inhibited growth seen at ≥2000 -S9
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA 98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	33.3-5000 µg/plate	MON 76313 30.9% glyphosate acid	Negative	Mecchi (2008b)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA 98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	10-5000 µg/plate (+/-S9)	MON 76171 31.1% glyphosate	Negative	Mecchi (2008c) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA 98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	10-5000 µg/plate (+/-S9)	MON 79991 71.6% glyphosate acid	Negative	Mecchi (2009a)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA 98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	10-5000 µg/plate (+/-S9)	MON 76138 38.5% glyphosate	Negative	Mecchi (2009b) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA97a, TA98, TA100, and TA1535 ± S9	1-5000 µg/plate	MON 77280 646.4 g/L salt equivalent	Negative	Perina (1998)	

Table F.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Formulations.						
Test/Endpoint	Test System	Concentrations	Test Material/ Concentration	Results	Reference	Comments
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98 and TA100 ± S9	0-1440 µg/plate (calculated as glyphosate IPA salt)	Roundup, 480 g/L glyphosate isopropylamine salt	Negative – S9, Equivocal +S9	Rank <i>et al.</i> (1993)	Stat significant increase at 360 µg/plate for TA98 (-S9) and 720 µg/plate for TA100 (+S9). Not significant at higher concentrations and were not replicated. Effects occurred at close to toxic levels.
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 ± S9	500-5000 µg/plate;	495 g/L glyphosate isopropylamine salt; 371.0 g/L (equivalent of glyphosate acid)	Negative ± S9	Silvino (2011)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 ± S9	1.5-5000 µg/plate	MON 8709 495 g/L glyphosate isopropylamine salt; 371.0 g/L (equivalent of glyphosate acid)	Negative ± S9	Silvino (2011)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, and TA1537 ± S9	15-5000 µg/plate	MON 76313 468 g/L glyphosate isopropylamine salt (351 g/L glyphosate acid equivalent)	Negative ± S9	Silvino (2012)	Cytotoxic at 5000 µg/plate for some strains
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA97a, TA98, TA100 and TA1535 ± S9	1-5000 µg/plate	Glifos formulation (glyphosate isopropylammonium salt, Berol 907 and water)	Negative ± S9	Vargas (1996)	Cytotoxic at the two upper concentrations

Table F.1. <i>In vitro</i> Test for Gene Mutations in Bacteria: Glyphosate Formulations.						
Test/Endpoint	Test System	Concentrations	Test Material/ Concentration	Results	Reference	Comments
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA102, TA1535, TA1537± S9	3.16-316 µg/plate	FSG 3090-H1 360 g/L	Negative ± S9	Uhde (2004) ¹	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100 ± S9	0.01-100 µg/plate	64% (glyphosate Isopropylammonium salt)	Negative ± S9	Wang <i>et al.</i> (1993)	
Bacterial Reverse Mutation	<i>S. typhimurium</i> TA98, TA100, TA1535, TA1537 and <i>E. coli</i> WP2 <i>uvrA</i> ± S9	All strains: 33.3-5000 µg/plate (+S9); 10-3330 µg/plate (-S9)	MON 78910 30.3% glyphosate acid	Negative ± S9	Xu (2006)	Cytotoxic ≥1000 µg/plate (-S9)

¹ Study was cited in Kier and Kirkland (2013). Supplementary information about the study was provided online including test guideline, test material purity, control chemicals and summary data tables.

Table F.2. <i>In Vitro</i> Tests for Chromosome Damage in Mammalian Cells- Glyphosate Formulations						
Test/Endpoint	Test System	Concentrations	Test Material/ Concentration	Results	Reference	Comments
<i>In vitro</i> Chromosomal Aberration using fluorescent in situ hybridization (FISH)	Bovine lymphocytes (from two 6-8 month old calves) -whole chromosome (1) painting probe	28-1120 µM 24 h exposure	62% Isopropylamine salt of glyphosate (38% inert ingredients)	Negative.	Holeckova (2006)	Small but significant increase in polyploidy seen at 56µM No positive control reported.
<i>In vitro</i> Cytokinesis Block Micronucleus Assay (with FISH analysis)	TR146 cells (human- derived buccal epithelial cell line)	0, 10, 15 and 20 mg/L; 20 minute exposure.	Roundup Ultra Max (450 g/l glyphosate acid)	Positive Increase in MN at all test concentratio ns	Koller <i>et al.</i> (2012)	No apoptosis observed at any conc. Necrosis reported at 20 mg/L. Increase in NB and NPB seen at all concentrations

MI= mitotic index. FISH= fluorescent in situ hybridization, MN= micronuclei; NB= nuclear buds; NPB= nucleoplasmic bridges.

Table F.3. <i>In Vivo</i> Tests for Chromosomal Aberrations in Mammals- Glyphosate Formulations.							
Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Chromosomal Aberration	Swiss albino mice (males only) Vehicle: DMSO	Intraperitoneal injection; sampling 24, 48 and 72 h	0, 25 and 50 mg/kg (5/dose)	Roundup (>41% isopropylamine glyphosate)	Positive Increase in MN at all time points at both doses	Prasad <i>et al.</i> (2009)	Significant decrease in mitotic index seen at all doses and time points
Bone Marrow Chromosomal Aberration	C57BL mice (males only) Vehicle: water	Oral administration; sampling 6, 24, 48, 72, 96 and 120 h	0.05, 0.01, 0.5 and 1.0% (8/dose)	Roundup	Negative	Dimitrov <i>et al.</i> (2006)	
Bone Marrow Chromosomal Aberration	New Zealand white rabbits (males only) Vehicle:	Drinking water for 60 days	0, 750 ppm (5/dose)	Roundup	Positive	Helal and Moussa (2005)	

BM= bone marrow, SC= spermatocyte.

Table F.4. <i>In Vivo</i> Tests for Micronuclei Induction in Mammals- Glyphosate Formulations.							
Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Micronucleus Test	Swiss CD1 mice (males only)	Intraperitoneal injection; 2 injections of half the dosage of 135 mg/kg 24 h apart; sampling at 6 and 24 h	0, 450 mg/kg roundup, equiv. to 135 kg glyphosate (3/dose)	Roundup, 30.4% glyphosate	Positive	Bolognesi <i>et al.</i> (1997)	Stat significant increase in MN at 6 and 24 h
Bone Marrow Micronucleus Test	C3H mice (males only) Vehicle: water	Intraperitoneal Injection (single treatment); sampling after 24, 48 and 72 h	0, 90 mg/kg	Not reported	Negative	Chruscielska <i>et al.</i> (2000)	
Bone Marrow Micronucleus Test	Swiss mice (males and females) Vehicle: water	Intraperitoneal Injection (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 50, 100 and 200 mg/kg	480g/L Isopropyla mine salt of glyphosate	Negative	Grisolia (2002)	
Bone Marrow Micronucleus Test	CD-1 mice (males and females)	Intraperitoneal injection; sampling 24, 48 and 72 h	0, 140, 280, and 555 mg/kg	Roundup (31% glyphosate salt)	Negative	Kier (1992)	Some deaths observed at high dose (HD), ↓PCE/NCE ratio at HD at 48 h in males.
Bone Marrow Micronucleus Test	Swiss albino mice (males and females)	Intraperitoneal Injection (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 212.5, 425 and 637.5 mg/kg	MON 77280 646.4 g/L glyphosate salt equivalent	Negative	Monma (1998)	Doses tested corresponded to 25%, 50% and 75% LD50

Table F.4. <i>In Vivo</i> Tests for Micronuclei Induction in Mammals- Glyphosate Formulations.							
Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Micronucleus Test	NMRI-Bom mice	Intraperitoneal Injection (single treatment); sampling after 24 h	0, 133 and 200 mg/kg (4/sex/dose)	Roundup, 480 g glyphosate isopropylamine salt per liter	Negative	Rank <i>et al.</i> (1993)	BM toxicity indicated by %PCE decreased at 200 mg/kg
Bone Marrow Micronucleus Test	Swiss albino mice (males only ²) Vehicle: water	Oral gavage (two treatments, 24 h apart); sampled at 18 and 24 h after last dose	0, 2000 mg/kg	MON 8709494.7 g/L salt of isopropylamine (371.0 glyphosate acid)	Negative	Claro (2011)	OECD 474 Guideline No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	C57BL mice (males only) Vehicle: water	Oral administration; sampling 6, 24, 48, 72, 96 and 120 h	0.05, 0.01, 0.5 and 1.0% (1%=1080 mg/kg) (8/dose)	Roundup	Negative	Dimitrov <i>et al.</i> (2006)	Toxicity seen in 1.0% dose group
Bone Marrow Micronucleus Test	CrI:CD-1(ICR) BR mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 78239 (36.6% glyphosate)	Negative	Erexson (2003a)	EPA Guideline (84-2) No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	CrI:CD-1(ICR) BR mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 78634 (65.2% glyphosate)	Negative	Erexson (2003b)	EPA Guideline (84-2) No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	CrI:CD-1(ICR) BR mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 78910 (30.3% glyphosate)	Negative	Erexson (2006)	EPA Guideline (84-2) No significant signs of toxicity observed in main study.

Table F.4. <i>In Vivo</i> Tests for Micronuclei Induction in Mammals- Glyphosate Formulations.							
Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Micronucleus Test	NMRI mice (males and females) Vehicle: 0.8% hydroxypropylmethyl cellulose	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 mg/kg (5/sex/dose)	TROP M (Glyphosate 480); 358.4 g/L glyphosate acid; 483.6 g/L IPA salt	Negative	Flügge (2010c) ¹	OECD Guideline 474 No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	Swiss mice (males only ²) Vehicle: water	Oral gavage (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 2000 mg/kg (6/dose)	A17035A 289.7 g/L glyphosate	Negative	Negro Silva (2009) ¹	OECD Guideline 474 No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	Swiss mice (males only ²) Vehicle: water	Oral gavage (2 treatments, 24 h apart); sampling after 24 h (last treatment)	0, 2000 mg/kg (6/dose)	Glyphosate SL (499.35 g/L glyphosate)	Negative	Negro Silva (2011) ¹	OECD Guideline 474 No significant signs of toxicity observed in main study
Bone Marrow Micronucleus Test	Hsd:CD-1(ICR) mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 79864 (38.7% glyphosate)	Negative #	Xu (2008a)	EPA Guideline (84-2) /OECD 474 No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	CD-1(ICR)BR mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 76171 (31.1% glyphosate)	Negative	Xu (2008b)	EPA Guideline (84-2) /OECD 474 No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	CD-1(ICR)BR mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 79991 (71.6% glyphosate)	Negative	Xu (2009a)	EPA Guideline (84-2) /OECD 474 No significant signs of toxicity observed in main study.

Table F.4. <i>In Vivo</i> Tests for Micronuclei Induction in Mammals- Glyphosate Formulations.							
Test/Endpoint	Test System	Route of Administration	Doses	Test Material Purity	Results	Reference	Comments
Bone Marrow Micronucleus Test	CD-1(ICR)BR mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 76138 (38.5% glyphosate)	Negative	Xu (2009b) ¹	EPA Guideline (84-2) /OECD 474 No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	Hsd:CD-1(ICR)BR mice (males only ²) Vehicle: water	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 (mg/kg) (5/dose)	MON 76313 (30.9% glyphosate)	Negative	Xu (2009c) ¹	EPA Guideline (84-2) /OECD 474 No significant signs of toxicity observed in main study.
Bone Marrow Micronucleus Test	CD rats (males and females) Vehicle: 0.8% hydroxypropylmethyl cellulose	Oral gavage (single treatment); sampling after 24 and 48 h (high dose only)	0, 500, 1000, and 2000 mg/kg (5/sex/dose)	757 g/kg granular formulation (69.1% glyphosate acid)	Negative	Flügge (2010e) ¹	OECD Guideline 474 No significant signs of toxicity observed in main study

¹ Study was cited in Kier and Kirkland (2013). Supplementary information about the study was provided online including test guideline, test material purity, control chemicals and summary data tables.

² Only males tested; report indicated that there were no difference between sexes seen in range finding study.

BM= bone marrow, CA= chromosomal aberrations, MN= micronucleated erythrocytes, NCE= normochromatic erythrocytes, PCE=polychromatic erythrocytes.

Table F.5. Other Assays for Detecting DNA Damage- Glyphosate Formulations.							
Test/Endpoint	Test System	Route of Administration	Doses/ Concentrations	Test Material/ Concentration	Results	Reference	Comments
Bacterial SOS Chromotest	<i>Escherichia coli</i> PQ37 strain	NA (<i>in vitro</i>)	0.25µg/sample	Roundup BIO formulation;	Positive	Raipulis <i>et al.</i> (2009)	
DNA Adducts ³²P-postlabeling	Swiss CD1 mice (males and females) Liver and kidney evaluated	Intraperitoneal injection	0, 400, 500 and 600 mg/kg, corresponding to 122, 152 and 182 mg/kg glyphosate salt	Roundup (30.4% isopropylammonium salt of glyphosate)	Positive (liver and kidney)	Peluso <i>et al.</i> (1998)	
DNA oxidative damage: 8-OHdG formation	Swiss CD-1 mice (males) liver and kidney evaluated	Intraperitoneal injection (single dose); sampling 4 and 24 h after injection	900 mg/kg corresponding to 270 mg/kg glyphosate (3/dose)	900 mg/kg corresponding to 270 mg/kg glyphosate	Kidney: positive at 8 and 24 h Liver: negative	Bolognesi <i>et al.</i> (1997)	
Single-cell gel electrophoresis (SCGE) assays-COMET assay	TR146 cells (human-derived buccal epithelial cell line). Alkaline conditions	NA (<i>in vitro</i>)		Roundup Ultra Max (450 g/l glyphosate acid)	Induced DNA migration at >20 mg/L	Koller <i>et al.</i> (2012)	Also measured multiple cellular integrity parameters to assess cytotoxicity. Formulation was more toxic than technical. Significant increase in LDHe at all concentrations tested. Cytotoxic ≥ 60 mg/L
Sister Chromatid Exchange (SCE)	Bovine lymphocytes	NA (<i>in vitro</i>)	28 - 1112 µM;; ±S9; sampling at 24 and 48 h	62% Isopropylamine salt of glyphosate	Positive	Sivikova & Dianovsky (2006)	

Table F.5. Other Assays for Detecting DNA Damage- Glyphosate Formulations.							
Test/Endpoint	Test System	Route of Administration	Doses/ Concentrations	Test Material/ Concentration	Results	Reference	Comments
Sister Chromatid Exchange (SCE)	Human lymphocytes (2 donors)	NA (<i>in vitro</i>)	250, 2500 and 25000 µg/mL	Roundup; Isopropylamine salt of glyphosate (purity not stated)	Stat. significant increase (p<0.001) at 250 µg/mL in both donors, and in one donor at 2500 µg/mL	Vigfusson and Vyse (1980)	No growth seen at highest concentration (25 mg/mL)
Sister Chromatid Exchange (SCE)	Human lymphocytes	NA (<i>in vitro</i>)	-S9: 0, 0.1 and 0.33 mg/mL; 72 h exposure	Roundup, 30.4% glyphosate	Positive	Bolognesi <i>et al.</i> (1997)	Stat significant increase in SCE/cell at ≥ 0.1 mg/mL
Alkaline elution assay- DNA single strand breaks	Swiss CD-1 mice (males) liver and kidney evaluated	Intraperitoneal injection (single dose); sampling 4 and 24 h after injection	900 mg/kg corresponding to 270 mg/kg glyphosate (3/dose)	900 mg/kg corresponding to 270 mg/kg glyphosate	Positive (Increased elution rate) at 4 hours in liver and kidney At 24 h, returned to control levels	Bolognesi <i>et al.</i> (1997)	Return to control values at 24 h may indicate DNA repair or reflect rapid elimination of compound

h= hour, NA= not applicable, SCE= sister chromatid exchange, LDHe= extracellular lactate dehydrogenase

Appendix G

The following studies were considered during the systematic review, but were excluded from the analysis.

Amer S.M. et al (2006). In vitro and in vivo evaluation of the genotoxicity of the herbicide glyphosate in mice. Bulletin of the National Research Centre (Cairo) 31 (5): 427-446.

Aboukila, R.S. et al. (2014). Cytogenetic Study on the Effect of Bentazon and Glyphosate Herbicide on Mice. Alexandria Journal of Veterinary Sciences, 41: 95-101.

Majeska (1982d) MRID 00126616

Majeska (1982e) MRID 00126614

Majeska (1982f) MRID 00126615