Shining a Light on Glyphosate-Based Herbicide Hazard, Exposures and Risk: Role of Non-Hodgkin Lymphoma Litigation in the USA

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Roundup, and other glyphosate-based herbicides, are the most heavily used pesticides in the history of the USA and globally. In March 2015, the International Agency for Research on Cancer (IARC) classified glyphosate as a “probable human carcinogen”. A portion of the 695,000 Americans then living in 2015 with non-Hodgkin lymphoma (NHL) became aware of IARC’s decision. Several thousand Roundup–NHL lawsuits had been filed by the end of 2017, rising to 18,400 by July 2019 and 42,000 by November 2019. Three cases have gone to trial, each won by the plaintiffs. The author has served as an expert witness for the plaintiffs in this litigation and has been compensated for his time spent. The impact of the litigation on the independent assessment of the science useful in determining whether glyphosate and glyphosate-based herbicide exposures are linked to NHL is reviewed, as is why the US Environmental Protection Agency (EPA) and IARC reached such different judgements regarding glyphosate human cancer hazard and risk. Two important “lessons learned” regarding the EPA versus IARC assessment of glyphosate cancer hazard and risk are highlighted. The first arises from differences in the magnitude of applicator risks from mostly dermal exposures to formulated glyphosate-based herbicides compared to just dietary exposures to technical glyphosate. The second relates to missed opportunities to markedly lower applicator exposures and risks with little or no impact on sales via reformulation, added warnings and worker safety provisions, company-driven stewardship programmes and greater determination by the EPA in the 1980s to compel Monsanto to add common-sense worker protection provisions onto Roundup labels (eg “wear gloves when applying this product”). Policy reforms designed to alleviate systemic problems with how pesticide hazards, exposures and risks are analysed, regulated and mitigated are described.

* Project Coordinator, The Heartland Study and Benbrook Consulting Services, Port Orchard, WA, USA; email: charlesbenbrook@gmail.com. Thanks to Yogi Hendin and Alessandra Arcuri for organising the 6 June 2019 symposium in Rotterdam “The Science and Politics of Glyphosate” (<www.eur.nl/en/research/erasmus-initiatives/dynamics-inclusive-prosperity/events/science-and-politics-glyphosate>). This paper expands upon my remarks delivered during the symposium. I appreciate the help of Rachel Benbrook with the figures and references cited herein. I have worked as an expert witness for plaintiffs in US litigation involving Roundup and non-Hodgkin lymphoma. This assignment provided me with access to a substantial number of Monsanto emails, studies and other documents not otherwise available, as well as the funding required to carry out detailed analyses of the basis of International Agency for Research on Cancer’s and the US Environmental Protection Agency’s classifications of glyphosate oncogenicity. The lawyers I work for had no role in the decision to write this paper, nor have they reviewed this paper. No external funding or payment was received in support of the writing of this paper. Rachel Benbrook’s contributions were supported by Benbrook Consulting Services.

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I. THE ROOTS OF ROUNDUP–NON-HODGKIN LYMPHOMA LITIGATION IN THE USA

The International Agency for Research on Cancer (IARC) issued its surprising classification of glyphosate-based herbicide (GBH) and glyphosate as a “probable human carcinogen” in March 2015,1 via a press release2 and short paper in The Lancet Oncology.3 Substantial media coverage followed the IARC announcement in the USA. Most stories made reference to controversies arising from the ongoing US Environmental Protection Agency (EPA) glyphosate re-registration process and/or recent developments in the State of California in the wake of the State’s listing of GBH consumer products as possible carcinogens under Proposition 65.4

The IARC classification decision led, within months, to the filing of lawsuits alleging that past use of Roundup had contributed to the plaintiffs’ cases of non-Hodgkin lymphoma (NHL). In the ongoing litigation over Roundup and NHL, the core issues in dispute arise at the nexus of cancer classification schemes, pesticide hazard and risk assessment policy, exposure assessment and risk mitigation requirements, legal standards, levels and burden of proof and evaluation of case-specific causality. Much of the expert witness testimony and arguments by attorneys presented to the juries during the three completed trials has focused on whether the EPA’s or IARC’s assessment of glyphosate oncogenicity is more relevant to deciding the merits of each plaintiff’s case.

After a multiyear review, the EPA concluded that glyphosate is not likely to pose an oncogenic risk.5 The agency’s evaluation focused primarily on risks arising from exposure to pure glyphosate (not formulated Roundup) via the diet across the general population. Given the agency’s focus on dietary exposures and the risks of glyphosate, the lack of weight placed on the toxicology studies done on formulated products is defensible.6 In addition, three factors led to, and sustained the EPA’s lack of concern over glyphosate dietary exposures:

- The generally infrequent and low levels of glyphosate residues in food;
- Glyphosate’s very high chronic reference dose;
- The generally high levels of glyphosate fed to animals in two-year rodent cancer bioassays.

1 The classification of chemical substances by the IARC relative to potential to cause cancer is hazard based and does not take into account the likelihood that a given level of exposure will trigger disease. The IARC takes three areas of data into account: results of animal bioassays, genotoxicity and other mechanistic studies and epidemiology.
2 IARC, “IARC Monographs Volume 112: Evaluation of five organophosphate insecticides and herbicides”. Press release: 20 March 2015: <www.iarc.fr/wp-content/uploads/2018/07/MonographVolume112-1.pdf>.
3 KZ Guyton et al; “Carcinogenicity of tetrachlorvinphos, parathion, malathion, diazinon, and glyphosate” (2015) 16 The Lancet Oncology 490.
4 See <https://oehha.ca.gov/proposition-65>.
5 The fullest and final justification of the EPA’s classification decision is set forth in the agency’s “Revised Glyphosate Issue Paper: Evaluation of Cancer Potential”, accessible at <https://cfpub.epa.gov/si/si_public_record_report.cfm? Lab=OPP&dirEntryID=337935>. The response of the EPA to public comments following the publication of glyphosate’s proposed re-registration and the “Interim Registration Review Decision” were posted in January 2020 and are accessible at <www.epa.gov/ingredients-used-pesticide-products/interim-registration-review-decision-and-responses-public>.
6 By the time a consumer ingests food with glyphosate residues, there is very little if any exposure to the surfactants incorporated in glyphosate-based herbicides.
However, a different set of factors come into play when the EPA, or anyone, assesses the possible oncogenic risks from exposures to formulated GBHs during mixing and loading operations and application episodes. Mixer-loaders and applicators are exposed via dermal absorption to formulated products, which are a mixture of glyphosate and various surfactants and adjuvants. Such mixtures can markedly enhance the toxicity of GBH exposure. This is because the surfactants in most GBHs facilitate the movement of glyphosate through the human epidermis and, once inside the body, through cell membranes. This accelerated movement of glyphosate increases the amount reaching the inside of cells, where it comes into contact with DNA and can trigger direct damage to DNA or oxidative stress, two genotoxicity mechanisms that are known to be associated with the formation of cancerous cell growths.

Despite these differences, the EPA did not conduct an applicator exposure risk assessment, nor render a judgement on the often much higher exposures and risks incurred by applicators, especially in the case of people spraying a GBH with handheld equipment. It chose not to do so because: “Dermal and inhalation endpoints were not selected since there was no toxicity observed in route-specific toxicity studies and there was no concern for increased quantitative susceptibility to offspring”. No dermal exposure endpoints were selected because none was observed. This is not surprising, given that the EPA had not required registrants to conduct any “route-specific toxicity studies”, nor had it carried out such studies itself. The EPA has used a 3% dermal absorption “default value” in conducting glyphosate worker-exposure assessments since the early 1980s, a value weakly supported by studies involving topically applied, pure glyphosate in mouse and primate studies. Evidence presented at trial showed clearly that, by the early 2000s, Monsanto scientists were aware that the surfactants in Roundup-brand herbicides were likely increasing glyphosate’s dermal absorption rates above 3%, but they did not share these data and reasons supporting this conclusion with the EPA, nor other regulators.

The EPA relied mostly on registrant-commissioned, unpublished animal bioassays and genotoxicity studies done on pure glyphosate, and it did not conduct a thorough review of, nor place much weight on genotoxicity studies done with formulated GBHs, including Roundup.

The IARC, on the other hand, classified glyphosate and GBHs as “probable human carcinogens”. The IARC Working Group considered a greater diversity of routes of exposure, including several published studies reflecting applicator exposures to formulated GBHs. The IARC’s evaluation took account of, and placed considerable weight on in vivo studies from people exposed directly to GBHs.

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7 E Wozniak et al, “The Mechanism of DNA Damage Induced by Roundup 360 PLUS, Glyphosate and AMPA in Human Peripheral Blood Mononuclear Cells – Genotoxic Risk Assessment” (2018) 120 Food and Chemical Toxicology 510; R Mesnage, C Benbrook and MN Antoniou, “Insight into the Confusion over Surfactant Co-Formulants in Glyphosate-Based Herbicides” (2019) 128 Food and Chemical Toxicology 137.

8 GA Dedeke, FO Owagboriaye, KO Ademola, OO Olujimi and AA Aladesida, “Comparative Assessment on Mechanism Underlying Renal Toxicity of Commercial Formulation of Roundup Herbicide and Glyphosate Alone in Male Albino Rat” (2018) 37 International Journal of Toxicology 285; N Defarge et al, “Co-Formulants in Glyphosate-Based Herbicides Disrupt Aromatase Activity in Human Cells below Toxic Levels” (2016) 13 International Journal of Environmental Research and Public Health 264.

9 EPA, Draft Human Health and Ecological Risk Assessments for Glyphosate: <www.epa.gov/ingredients-used-pesticide-products/draft-human-health-and-ecological-risk-assessments-glyphosate>.
As a result of these differences, the EPA’s assessment of glyphosate oncogenicity is primarily relevant to one route of exposure (dietary) and generally low rates of exposure, while the IARC’s evaluation spans all routes of exposure and the sometimes high levels of applicator exposure. In this paper, I explain the differences in the following: (1) the questions that the EPA and IARC set out to answer; and (2) the data that the EPA and IARC relied on in reaching their divergent classification decisions. I argue that a full accounting of these differences goes a long way towards explaining why and how the EPA and IARC reached diametrically opposed conclusions.

Throughout the first three trials, attempts to convince jurors that exposure to Roundup likely did or did not contribute to a plaintiff’s NHL were the primary focus of both expert testimony and the presentations of evidence by attorneys. In this body of litigation, two key pieces of the puzzle were clear: the plaintiffs had used Roundup frequently and had been exposed to it many times; and the plaintiffs had suffered and continued to suffer from NHL. The matter of dispute was whether and to what degree exposure to Roundup contributed to a given plaintiff’s NHL, taking into account other possible causes of a given plaintiff’s NHL.

Defence attorney presentations of evidence began, ended and never veered far from the EPA’s “not likely to pose oncogenic risk” classification. Plaintiff attorneys focused instead on the IARC’s synthesis of relevant data and its “probable carcinogenic” classification. They explained why the IARC’s assessment was more relevant to the exposures incurred by the plaintiffs. Plaintiff attorneys and experts did not directly dispute nor strive to question the EPA’s judgement that current, projected dietary exposures to glyphosate posed little if any cancer risk. However, they did explain in detail why the EPA’s analysis did not address nor encompass the much higher levels of exposure experienced by applicators, nor the fact that applicators are exposed to a mixture of chemicals, and not just pure technical glyphosate.

A factor unique to this sort of litigation warrants attention: the applicable level and burden of proof under US law and judicial precedent. To prevail on the question of liability and compensatory damages, plaintiff attorneys and experts must convince a jury that a preponderance of the evidence supports a conclusion that it is “more likely than not” that the plaintiff’s use of and exposure to Roundup contributed to his or her NHL. Thus, it is not necessary to prove that a plaintiff’s NHL was caused solely, or even primarily, by exposures to Roundup. A jury can find in favour of a plaintiff if they feel that a preponderance of the evidence supports the conclusion that Roundup exposures accelerated the progression of a plaintiff’s NHL or made it more difficult to treat and control.

The legal standard to support an award of punitive damages is stricter and the nature of the evidence differs. To support a punitive damage award, there must be “clear and convincing” evidence that a defendant has acted with malice, oppression and/or a blatant disregard for public safety or the rights of others. Both of these levels and burdens of proof are not nearly as onerous as the “beyond a reasonable doubt”

10 <www.law.cornell.edu/wex/preponderance>.
11 “Clear and convincing means that the evidence is highly and substantially more likely to be true than untrue; the fact finder must be convinced that the contention is highly probable”. Quoted from <www.law.cornell.edu/wex/clear_and_convincing_evidence>.
threshold applicable in criminal cases. These evidentiary burdens are well-known and accepted by lawyers and judges engaged in mass tort litigation, but they are not necessarily understood, or accepted, by non-lawyers. Some critics of the litigation argue that plaintiff attorneys and experts have not proven that a given plaintiff’s case of NHL was caused by his or her use of Roundup. In doing so, they impose on the proceedings an evidentiary and scientific threshold beyond that required by law. It is surely fair to question whether the level and burden of proof applicable in ongoing Roundup–NHL litigation is proper, but that debate should be pursued in a forum other than a courtroom in which a given Roundup–NHL case is tried.

In the spring of 2015, approximately 695,000 Americans were living with NHL, and an estimated 74,200 will be newly diagnosed with the disease in 2019. A substantial number of the individuals with NHL no doubt read or were told about the IARC’s classification of GBHs and glyphosate as “probable human carcinogens”. Some of these people likely recalled applying Roundup in the years prior to their diagnosis and consulted an attorney to explore whether the specifics of their case warranted joining the litigation. Two key factors taken into account in initial case reviews were the extent of a plaintiff’s use of Roundup and likely exposure levels, and the presence/absence of other known NHL risk factors.

Several law firms with experience in toxic chemical, mass tort litigation recognised the significance of the IARC’s classification decision and began assessing whether there was a plausible evidentiary and legal basis to pursue litigation against Monsanto, the sole manufacturer of GBHs under the Roundup brand from the mid-1970s through the early 2000s. Monsanto remained the major GBH manufacturer after the entrance of several basic and me-too manufacturers into the GBH market in the 2000s. Bayer acquired Monsanto in June 2018, and today Bayer/Monsanto is still by far the dominant player in the GBH market in the USA and globally.

The first Roundup–NHL case was filed by Enrique Rubio on 22 September 2015 in a US District Court in California. Three other early plaintiffs, Joselin Barrera, Elisa de la Garza and Judi Fitzgerald, filed their cases in Delaware in October 2015. Since few firms had the resources to take on litigation of this scope and expense on their own, a group of law firms began the process in 2016 of coordinating the key tasks required by the litigation. This allowed the firms to share the technical burden and costs of pursuing litigation against then-Monsanto on behalf of individuals with NHL.

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12 <www.law.cornell.edu/wex/burden_of_proof>.  
13 Data from the National Cancer Institute for 2016: <https://seer.cancer.gov/statfacts/html/nhl.html>.  
14 <https://seer.cancer.gov/statfacts/html/nhl.html>.  
15 A “basic” manufacturer of GBHs is a company with its own chemical production facilities that seeks and obtains its own registrations for technical and end-use products from the EPA, without relying on the data of any other registrant. “Me-too” manufacturers of pesticides purchase technical, pure active ingredient from a basic manufacturer and generally pay that company what amounts to a licensing fee to cite EPA-required testing data previously generated and submitted to the agency by the basic manufacturer. These agreements also typically include the right to adopt and use the provisions of already-approved EPA labels on the labels of their so-called “me-too” products.  
16 When Bayer acquired Monsanto in 2018, the company’s stated plan was to retire the Monsanto name. Subsequently, Bayer decided to retain the Monsanto name. Accordingly, in this paper, the corporate entity manufacturing and selling Roundup-brand herbicides since 2018 is referred to as Bayer/Monsanto.  
17 Access the case filings at <https://casetext.com/case/rubio-v-monsanto-co-2> and the Barrera et al case filings at <https://www.courtlistener.com/opinion/4256060/barrera-v-monsanto>.
1. Outcomes of the first three trials

a. Johnson v. Monsanto

Three trials in this area have been completed to date; each is briefly discussed herein. The case that would be the first to come to trial was filed on 28 January 2016 by the Miller Firm in California State Court on behalf of Dewayne Johnson, a school district groundskeeper. The Johnson case trial started on 9 July 2018 and was argued by Brent Wisner, an attorney working for the Baum Hedlund law firm based in Los Angeles, California. I was among the expert witnesses testifying on behalf of Mr Johnson. On 10 August 2018, the jury found in favour of Mr Johnson and awarded US$39.2 million in compensatory damages and US$250 million punitive damages (see the trial transcript and official jury form for the questions placed to the jury by the judge and the jury’s answers).

b. Hardeman v. Monsanto

Concomitant with the progression of the Johnson case through pre-trial motions, trial scheduling, jury selection and the trial itself, two other cases were moving through similar steps towards trial in other California courts. The Edwin Hardeman case was the first scheduled for trial as part of federal-district court multiple-district litigation (MDL) overseen by Judge Vincent Chhabria in San Francisco. On 17 November 2016, Judge Chhabria named Weitz & Luxenberg, the Miller Firm and Andrus Wagstaff as the plaintiff’s co-lead counsel for the MDL. These co-lead counsel firms were joined by three additional firms on an MDL case management executive committee. The role of the lead counsel and executive committees was to oversee and coordinate the work of the plaintiff law firms and interactions with Judge Chhabria’s court as the cases within the MDL moved forwards.

Judge Chhabria decided to split the Hardeman trial into two phases: Phase 1 focused just on the plausibility of Roundup–NHL causality. Due to the limited focus of Phase 1, plaintiff attorneys were not allowed to introduce any documents or present any expert testimony on Monsanto’s behaviour, conduct and actions relative to studies presented to the EPA, or Monsanto-supported studies and reviews published in peer-reviewed journals. The judge’s stated goal was to ensure that the jury made its decision regarding Roundup–NHL causality based only on scientific evidence and without prejudice from information or testimony regarding Monsanto’s behaviour and efforts to shape regulatory decisions or policy.

The Hardeman Phase 1 trial began on 25 February 2019 and went to the jury on 14 March 2019. The Phase 1 verdict in favour of Mr Hardeman came on 19 March. On the next day (20 March), Phase 2 commenced, focusing on Monsanto’s behaviour and compensatory and punitive damages. After just a week, the second phase of the
trial was completed and the case went to the jury. Their verdict and award were announced by Judge Chhabria on 27 March 2018.

The jury awarded Mr Hardeman US$5.1 million in compensatory damages and US$75 million in punitive damages. In response to a post-verdict motion by Bayer/Monsanto attorneys, Judge Chhabria modestly altered the compensatory damages award and reduced the punitive damages to US$20 million, for a total award to Mr Hardeman of US$25.3 million. In allowing US$20 million in punitive damages to stand, Judge Chhabria noted Monsanto’s efforts to shield the EPA from data raising new questions about Roundup’s safety and the company’s aggressive efforts to influence scientific discourse and the glyphosate-focused content of peer-reviewed journals.

Post-Hardeman, potential plaintiffs and law firms became more confident of the scientific merits of the litigation. As a result of the Johnson and Hardeman verdicts and awards, the number of cases filed rose steadily in 2018 and 2019 from a few thousand in 2017, to around 4500 at the time of the Johnson verdict in mid-2018, to over 42,000 in November 2019, as shown in Figure 1.

c. Pilliod v. Monsanto

Just weeks after the conclusion of the Hardeman federal MDL trial, a third case in California State Court began. Alva and Alberta Pilliod had both been diagnosed with the same form of NHL. This married couple had sprayed Roundup several days per year on their rural California properties for over two decades. They often walked together, each with a handheld sprayer, spot-spraying berry bushes and other weeds and vegetation choking the trails they had created and maintained. On some occasions, they wore shorts and flip-flops as they sprayed Roundup on encroaching weeds along trails.

Figure 1: Non-Hodgkin lymphoma lawsuits, milestones and trends.

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21 Access all Pilliod case and trial documents at <www.baumhedinlaw.com/pilliod-v-monsanto-trial>.
Judge Winifred Smith presided over the Pilliod trial. The plaintiff attorneys presented their case in about one month, beginning on 28 March 2019. The defence attorneys presented their case in about one week. Jury deliberations began on 5 May 2018, and the verdict and monetary awards were announced on 13 May. The Pilliods received a stunning punitive damage award of US$2 billion dollars. In response to post-trial motions, Judge Smith reduced the combined (Alva and Alberta) punitive damage award to US$69 million, citing Supreme Court guidance that classifies any punitive damage award greater than seven times the corresponding compensatory damages award as excessive and likely in need of judicial review and adjustment.

2. Why such large awards?

In her post-trial order, Judge Smith set forth what she regarded as part of the justification for a still significant US$69 million punitive damage award:

In this case there was clear and convincing evidence that Monsanto made efforts to impede, discourage, or distort scientific inquiry and the resulting science.22

Comparable sentiments had been voiced earlier by Judge Curtis Karnow of the Superior Court of San Francisco County, California. He presided over several pre-trial motions setting forth the ground rules governing the Johnson trial. In his 17 May 2018 order in response to multiple pre-trial motions, Judge Karnow wrote:

The internal correspondence noted by Johnson could support a jury finding that Monsanto had long been aware of the risk that its glyphosate-based herbicides are carcinogenic, and more dangerous than glyphosate in isolation, but has continuously sought to influence the scientific literature to prevent its internal concerns from reaching the public sphere and to bolster its defenses in products liability actions.23

Table 1 provides an overview of the compensatory and punitive damage awards in the above three trials as initially awarded by the jury, and later adjusted by the trial judges. The bottom two lines in Table 1 provide average compensatory, punitive and total damage awards per plaintiff, highlighting the sizable reduction in the punitive damage awards from an average of US$581 million per plaintiff by juries to US$32.2 million after adjustments by the three judges.

On average, across the first four plaintiffs whose cases have gone to trial, the total awards have been US$47.6 million per person. After reductions by the three judges, the average ratio of punitive-to-compensatory damage awards is about 2:1, well below the threshold of 7:1 set by the Supreme Court for review of punitive damage awards.

22 Judge Winifred Smith, Assistant Presiding Judge for the Superior Court of Alameda County, California, 25 July 2019 Post-trial Order, Pilliod v. Monsanto Company: <www.baumhedlundlaw.com/pdf/monsanto-documents/Pilliod/Pilliod-order-denying-Monsanto-Motion-7-25-19.pdf>.

23 <https://usrtk.org/wp-content/uploads/2016/09/Judges-order-in-Johnson-Case-ahead-of-trial.pdf>.
From the mid-1970s through early 2015, few scientists, doctors, pesticide users, regulators and environmentalists were concerned about Roundup use, exposures and risks to the general public. It was widely stated by industry experts that Roundup breaks down in the environment naturally to harmless, ubiquitous elements. Despite very limited testing of food for glyphosate residues by the US government, there was no reason to expect the presence of glyphosate in food above very low, rarely detectable levels mostly from drift (except for raw soybeans). This is because the herbicide could only be applied before crop germination – and hence months before the edible portion of a crop is formed and ready for harvest. The frequency and levels of glyphosate residues in certain foods changed with the registration of pre-harvest crop desiccation uses of GBHs in the 1980s and 1990s and genetically engineered Roundup Ready (RR) crops in the second half of the 1990s. Since the mid-2000s, glyphosate dietary exposures have been common, and levels have been rising in the urine of Americans.

In addition, in the early 1980s, the EPA set a very high chronic reference dose (cRfD; also known as an “acceptable daily intake” (ADI) in much of the rest of the world) of

24 In 2011, the US Department of Agriculture’s Pesticide Data Program tested 300 samples of soybeans, the majority of which were genetically engineered to resist post-emergent applications of GBHs. Glyphosate was found in 90.3% of samples at an average level of 1.9 ppm, and glyphosate’s major metabolite aminomethylphosphonic acid (AMPA) was present in 95.7% of the samples at an average level of 2.3 ppm.

25 S Mills et al, “Excretion of the Herbicide Glyphosate in Older Adults between 1993 and 2016” (2017) 318 JAMA 1610.
1.75 mg/kg/day. At the time, only two widely used pesticide active ingredients had higher cRfDs (the imidazolinone herbicides imazapyr and imazethapyr had cRfDs equal to 2.5 mg/kg/day). This means that the EPA then viewed glyphosate as less chronically toxic than all but two currently registered pesticides for which a cRfD had been established.

The IARC determination in March 2015 and the ensuing litigation triggered the first in-depth and independent assessment of Roundup/glyphosate uses and exposures in relation to several important risk factors:

- Toxicity;
- Mixer-loader and applicator dermal absorption and exposures;
- Glyphosate absorption, distribution, metabolism and excretion;
- Impacts of the surfactants in GBH formulations on toxicity and absorption;
- The capacity of GBHs to damage DNA in ways that might lead to cancer, and especially NHL.

By “independent” assessment in the context of this litigation, I mean one that is carried out by an experienced and well-trained scientist who does not currently work for a GBH registrant and has not been significantly dependent on funds from a pesticide manufacturer during prior stages of their career. Both the IARC assessment and the reports and testimony by experts working on behalf of plaintiffs meet these criteria. So, too, do some of the experts who have worked for defendant attorneys in the Roundup–NHL litigation. Experts working for plaintiff attorneys are paid for the time invested in their work, but their careers have generally not been shaped by or been dependent on sustaining a mutually rewarding professional and financial relationship with a company with a major stake in the outcome of litigation.

In addition, the scientific work and insights gained by plaintiff experts were informed and shaped by a unique aspect of this litigation: access via the discovery record to Monsanto’s internal discussions of scientific concerns, risk assessment issues and findings from Monsanto-conducted proprietary research.

The record contains thousands of emails discussing what Monsanto could have done, should have done and eventually did in response to newly published data or their own internal studies that might alter regulatory risk assessments and lead to new restrictions on Roundup use. The issues and concerns regarding GBH human health risks that Monsanto has focused on for decades include elevated tumours in animal bioassays, dozens of positive genotoxicity assays, problems with published epidemiological studies

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26 For a current overview of the history of glyphosate’s cRfD, see <www.epa.gov/pesticides/epa-releases-draft-risk-assessments-glyphosate>.

27 cRfDs are set by the EPA by dividing the lowest relevant “no observable adverse effect level” (NOAEL) in animal studies by a 100-fold standard safety factor. cRfDs can change over time when new mammalian toxicology data become available. Ironically, in light of industry claims in the late 1990s and 2000s that Roundup/GBHs were displacing higher-risk herbicides, Roundup/GBHs also displaced then-market-leading, less toxic imidazolinone herbicides, especially in soybean weed management systems.
reporting an association between Roundup exposures and cancer risk, the impact of GBH surfactants on applicator exposures and risk and the appropriate rate of dermal absorption to use in applicator exposure assessments. The way Monsanto responded to new science and risk concerns helped plaintiff experts identify possibly consequential scientific issues to focus on. Differences between what Monsanto scientists said in internal email chains about possible risk concerns compared to what Monsanto told regulators and/or published in peer-reviewed journals received substantial focus in plaintiff expert reports and trial testimony.

Over time, more discovery documents will be unsealed. When studied in conjunction with the already available and extensive body of both plaintiff and defendant expert reports, deposition transcripts, transcripts of trial testimony and trial exhibits, other scientists and the public will have the same opportunity as plaintiff experts to follow Monsanto’s evolving knowledge about Roundup and GBH risks. Those that do so can then reach their own conclusions regarding what the company did and did not do to steward the use of Roundup and sharpen the accuracy of Roundup risk assessments.

1. Insights abound in the massive discovery record
Several million documents have been turned over to the courts by Monsanto in response to queries from plaintiff attorneys, including millions that no regulator or member of the public has seen. Some documents shed light on what Monsanto knew, and was concerned about, regarding Roundup toxicity and exposures, while others outline and track the impact of the often elaborate efforts undertaken by the company and its surrogates to undermine published science, and scientists, questioning Monsanto’s view of Roundup safety.

Because the majority of these documents remain under seal, it is not possible to cite and discuss their content. However, many of the significant documents have been unsealed and are available from several sources. These documents include:

- Essentially all official communications between Monsanto and the EPA over the life history of Roundup, including Monsanto’s internal speculation regarding the basis for new risk concerns raised by regulators and how to assuage such concerns.

- The results of Monsanto-conducted or commissioned studies on glyphosate or Roundup, including internal studies and preliminary reports never disclosed to regulators (including some drafts of study reports containing more worrisome results than final reports).

- Thousands of documents addressing Monsanto’s marketing strategies, efforts and materials.

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28 From a law firm involved in the litigation: <www.baumhedlundlaw.com/toxic-tort-law/monsanto-roundup-lawsuit/monsanto-secret-documents>; from a non-governmental organisation tracking the litigation: <https://usrtk.org/monsanto-papers>; or from an academic program/document repository: <www.industrydocuments.ucsf.edu/chemical/collections/roundup-litigation-documents>.
Monsanto efforts to influence the flow of information to the scientific community and attitudes and opinions regarding Roundup safety among scientists, occupational health specialists, the media, politicians and farmers.

Stewardship policies, programmes and commitments in terms of product safety and worker health, including Monsanto’s input during the evolution of the FAO Code of Conduct for pesticide manufacturers and the company’s obligations as a signatory.

Internal discussions and debates among Monsanto scientists, the company’s regulatory officials and corporate leaders over what the company should do in response to new, negative information in published studies and/or new concerns raised by regulators (usually European regulators, especially since the early 1990s) or scientific bodies (eg the IARC).

Until the four lead law firms began hiring experts in the science areas central to the Roundup–NHL litigation, few if any independent scientists and physicians in the USA had ever had the funding, time or access to data needed to carry out an in-depth assessment of what was known about Roundup-induced human health risks. During 2016–2018, plaintiff attorneys and experts benefited greatly from access to the discovery record. Individuals working for the lead law firms conducted systematic topical searches, often at the request of plaintiff experts, and then shared relevant documents with the experts working in a given area.

A set of revealing Monsanto documents and internal email exchanges have been identified that discuss the concerns of Monsanto scientists and regulatory officials arising from both the company’s internal studies and published science in peer-reviewed journals. Many have been unsealed by judges, and they are often referred to as the “Monsanto Papers”. These documents include internal Monsanto exchanges that address possible risk-driven problems emerging in ongoing US EPA and European glyphosate re-registration reviews. Some of these discussions date back to the early 1980s and primarily focus on adverse glyphosate and GBH health and safety data and cancer risk.

Some key milestones in the history of GBH risk assessment addressed in the “Monsanto Papers” have been:

- The EPA’s conclusion in 1984 that a Monsanto-commissioned mouse study showed a statistically significant increase in a rare tumour (renal tubular adenomas) in male mice, leading to an Office of Pesticide Programs – Toxicology Branch-recommended “possible human carcinogen” classification of glyphosate that remained in place from 1984 through 1991.

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29 Discovery continues. Tens of thousands of additional documents have been disclosed in 2019.
30 Despite the thousands of hours spent by people working since 2016 to identify relevant documents in the discovery record, relevant documents continue to be identified. Plus, as both plaintiff and defence experts and lawyers deepen their understanding of a given issue or series of events, the importance of seemingly obscure comments and information in certain Monsanto documents becomes apparent upon a second (or third) reading. This progression of understanding of what happened and why is common in sustained mass tort litigation and is why the substance and focus in trials tends to evolve.
31 The Baum Hedlund law firm is the original source of most of the “Monsanto Papers”; access them at <www.baumhedlundlaw.com/toxic-tort-law/monsanto-roundup-lawsuit/monsanto-secret-documents>. 
Publication of multiple peer-reviewed papers beginning in the 1990s reporting glyphosate and/or GBH genotoxicity.33

Epidemiological studies reporting statistically significant linkages between GBH use, exposures and the risk of cancer, especially NHL.34

Each of the above milestones triggered concerted efforts by Monsanto to raise questions about the design, conduct and interpretation of studies reporting a possible new risk concern following exposure to Roundup. Such efforts often lasted for years, even decades. Several have played out multiple times, such as in the back-and-forth following the March 2015 IARC classification, and most recently in the course of each of the three trials. Common examples of Monsanto actions taken to raise doubts about adverse data include citing inappropriate tumour historical control data,35 arguing that a tumour observed in a cancer bioassay was not treatment related for specious reasons in violation of EPA cancer risk assessment guidelines36 and dismissing a positive result in an in vivo genotoxicity assay because of an inappropriate method of administration of the test substance or excessively high dose rates.

But unlike all past venues and cycles in which glyphosate and GBH health impacts have been discussed, the US Roundup–NHL litigation has been the first time that the rules of scientific engagement were set by courts to ensure an equal opportunity by plaintiff and defence lawyers and experts to present their best cases possible. The judges overseeing each of the trials rigorously enforced evidentiary and equal time rules.

These trials marked the first time scientists independent of GBH registrants and regulators had the opportunity to assess such an extensive portion of the full record of glyphosate risk assessment science, including Monsanto concerns in the wake of negative study results or new questions from regulators. Other documents reviewed by plaintiff experts and presented to the juries describe Monsanto plans and actions to undermine confidence in any published finding that raised a new or more serious risk concern over Roundup use and exposures. Significantly, the record convincingly shows that Monsanto’s near-universal response to new adverse data was to attack the messenger and the science, rather than to conduct new and more sophisticated studies capable of dismissing or confirming – and possibly quantifying – potential new areas of risk.

In short, the Roundup–NHL litigation has provided the venue and funding for the first open, rigorous and data-driven assessment of Roundup–NHL risks conducted by

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32 <https://archive.epa.gov/pesticides/chemicalsearch/chemical/foia/web/pdf/103601/103601-171.pdf>.
33 See the in-depth review of published genotoxicity studies in the IARC’s Monograph 112, accessible at <https://publications.iarc.fr/Book-And-Report-Series/Iarc-Monographs-On-The-Identification-Of-Carcinogenic-Hazards-To-Humans/Some-Organophosphate-Insecticides-And-Herbicides-2017>.
34 See the IARC’s review and discussion of the epidemiological literature in Monograph 112, pp 331–50.
35 EPA scientists instructed Monsanto to submit historical control data in the same species of laboratory animal from studies done in the same laboratory within a few years of when a given experiment was conducted. Monsanto often submitted, and asked the EPA to rely on historical control data from other laboratories and outside the timeframe considered by the EPA as relevant.
36 For dozens of examples, see the report and transcript of the 13–16 December 2016 Scientific Advisory Panel report: <www.epa.gov/sites/production/files/2017-03/documents/december_13-16_2016_final_report_03162017.pdf>.
scientists with roughly equal access to and familiarity with the pertinent evidence. Each of the plaintiff experts responsible for addressing a critical scientific or medical area – NHL diagnosis and treatment, toxicology, epidemiology, genotoxicity, animal bioassays and risk assessment – spent hundreds if not thousands of hours in the course of preparing for and participating in the first three trials.

For each expert on both sides of the litigation, the process started with preparation of often near-book-length expert reports answering questions put to them by counsel. Each expert was then deposed under oath by Bayer/Monsanto or plaintiff attorneys, who had the right to review and question the basis of any factual statements or opinions offered in an expert’s report. Such videotaped depositions typically lasted a day, but some spanned up to three days. Over the last three years, most plaintiff experts have been deposed eight or more times, beginning with their pre-Johnson trial deposition. Experts must defend their methods and analytical procedures, and they typically are asked to explain any deviations from the methods preferred and used by Monsanto scientists and/or regulators. Often, plaintiff experts are asked to explain why an alternative analysis or judgement is wrong (e.g., IARC versus EPA on genotoxicity). Whenever possible, attorneys attempt to back experts into a corner in order to generate testimony that can be used at trial to characterise the expert’s methods and analysis as outside of the scientific mainstream and purposefully biased. But importantly, everyone involved in the process knows that if such an attempt is made at trial, the expert whose work is called into question will have an opportunity to respond and defend his or her work.

In my work on this litigation, I concluded early on that the differences in the IARC and EPA analyses of the GBH and glyphosate genotoxicity databases were critical to understanding why the IARC and EPA reached such different cancer classification decisions. My first expert report stated this conclusion and set out why I had reached this opinion. During my first depositions, Monsanto attorneys allotted a significant share of their time to questioning me about this opinion. For example, after putting a proprietary genotoxicity study in front of me, I would be asked something like, "Dr Benbrook, are you familiar with this study?" Often, the answer was "no." The next series of questions might repeat this question-and-answer process on several more internal genotoxicity studies that I had not seen or reviewed. Then I would be asked something like, “Now that you are aware of these x number of studies, do they change your opinion?”

Questions of this sort are clearly legitimate and I should have been able to answer them. For this reason, I studied in greater detail the full body of genotoxicity assays available on glyphosate and GBHS and conducted an in-depth comparison of the EPA’s and IARC’s assessments of the relevant genotoxicity and mechanistic data. The results of my ongoing analytical work in this area were explained in my second and third expert reports, filed respectively as part of the Hardeman and Pilliod cases. My analysis of this component of the glyphosate and GBH cancer classification decision process served as the basis for a paper I wrote on this topic and published in *Environmental Sciences Europe.*

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37 Access most expert reports filed as part of the Johnson, Hardeman and Pilliod cases at <www.baumhedlundlaw.com/toxic-tort-law/monsanto-roundup-lawsuit>.

38 C Benbrook, “How Did the U.S. EPA and IARC Reach Diametrically Opposed Conclusions on the Genotoxicity of Glyphosate-Based Herbicides?” (2019) 31 Environmental Sciences Europe 2.
Some of the experts in the Roundup–NHL litigation had, furthermore, been involved with glyphosate and GBH use, regulation and risk issues for many years, if not decades. For example, in 1982–1983, I served as the Staff Director for a Congressional subcommittee with jurisdiction over federal pesticide law and regulation. Roundup was among the pesticides that received considerable attention by the subcommittee. Glyphosate and Roundup oncogenicity, risk assessment, use and regulation have continued to arise throughout my career. Dr Christopher Portier, the plaintiff expert addressing in detail the results of mouse and rat cancer bioassays, has also been involved with Roundup/GBH oncogenicity over several decades. Other examples could be cited among both defence and plaintiff experts.

Throughout the trials, the first challenge facing both attorneys and experts was explaining to juries the basic concepts, tools and approaches used to assess pesticide risk. Their second task was to explain the relevance and reliability of a given study’s reported findings. Last, they had to knit together multiple study results and lines of evidence into a hopefully coherent and persuasive “weight of evidence” judgement regarding the potential link – or lack thereof – between exposure to Roundup and NHL.

III. THE CASE PRESENTED TO JURIES

One of the core questions addressed throughout all three trials was whether the EPA was right in its judgement that glyphosate is unlikely to pose oncogenic risk, or whether the IARC was correct in reaching its “probable human carcinogen” classification. Over several weeks, juries heard plaintiff and defendant experts explain and critique the results and implications of the extensive animal cancer bioassay database (14 roughly 2-year studies, 6 in mice and 8 in rats), the hundreds of genotoxicity assays conducted on glyphosate and GBHs and the epidemiological database composed of over a dozen cohort studies, case–control studies and meta-analyses.

Monsanto experts presented testimony generally praising the design and conduct of studies that supported their assertion of no credible or persuasive data supporting a link between glyphosate or GBH exposures and NHL. In the case of dozens of studies that produced data supportive of a possible link between GBH use and NHL, they focused their testimony on perceived flaws in study design, inappropriate dosage

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39 See, for example, <www.ncbi.nlm.nih.gov/pubmed/26941213>. Dr Portier did his PhD dissertation on the design of two-year animal feeding studies. His work provided important contributions to the methods incorporated in the cancer feeding studies conducted on glyphosate and all other pesticides. See also the 2020 Portier paper summarising his assessment of the valid animal bioassays on glyphosate in *Environmental Health*, doi: 10.1186/s12940-020-00574-1.

40 The fullest and final justification of the EPA’s classification decision is set forth in the agency’s “Revised Glyphosate Issue Paper: Evaluation of Cancer Potential”, accessible at <https://cfpub.epa.gov/si/si_public_record_report.cfm?Lab=OPP&dirEntryID=337935>.

41 IARC revised Monograph 112: <https://publications.iarc.fr/Book-And-Report-Series/Iarc-Monographs-On-The-Identification-Of-Carcinogenic-Hazards-To-Humans/Some-Organophosphate-Insecticides-And-Herbicides-2017>.

42 Multiple statements and testimonies by Monsanto lawyers and experts actually went further, asserting that there were “no data” or “no evidence” of a link between Roundup use and exposures and any adverse health outcome. In a subset of such cases, the assertion is conditioned on a phrase such as “... at expected levels of exposure”, or “... when the product is applied in accord with the label”. Significantly, given the focus of the Roundup–NHL litigation on applicator and occupational exposures, no such statement by any regulatory agency nor Monsanto has claimed that there is “no evidence” of potential adverse health outcomes among more heavily exposed mixer-loaders and applicators.
levels or routes of administration, inappropriate statistical inferences, sources of bias or other criticisms. Consistently, Bayer/Monsanto attorneys and experts cited the views of regulators on specific studies and findings, but only when such views were aligned with Monsanto’s views.

Plaintiff experts addressed the findings of most of the same studies analysed by the Monsanto team. Often plaintiff experts acknowledged the existence of study flaws pointed out by Monsanto experts, but argued that reliable findings and/or inferences could still be derived from a given study. They also often highlighted the number and diversity of studies in each key area (animal bioassays, genotoxicity and epidemiology) reporting statistically significant results supportive of a Roundup–NHL link. They also pointed out that both Monsanto and the EPA deviated from applicable EPA cancer risk assessment policies and guidelines in advancing various reasons to dismiss dozens of positive results (eg historical controls, too-high doses, inappropriate statistical tests, errant histopathology).

During trial testimony, juries heard about an EPA Scientific Advisory Panel (SAP) convened in December 2016 to review the EPA’s and the IARC’s reviews of the glyphosate and GBH cancer data. The purpose of the SAP meeting was to shed light on why the EPA and IARC reached such different conclusions. After three days of detailed presentations and debate, the SAP members came to consensus on few important conclusions. One unanimous conclusion was that the EPA had not followed its own cancer risk assessment guidelines in its review of glyphosate oncogenicity. Key deviations addressed by the SAP occurred in the EPA’s assessment and use of historical control data, requiring a positive tumour response in both male and female treatment groups to regard a statistically elevated tumour in one sex as biologically relevant, a variety of issues arising around the presence/absence of a maximum tolerated dose and whether dozens of tumour types that were elevated in treatment groups based on linear statistical tests but not under pairwise comparisons were “treatment related”.

Nearly all of the 14 cancer bioassays produced statistically elevated levels of one or more tumours in one or both sexes. Each of the six mouse studies reported evidence of treatment-related lymphatic tumours. Some of the rat studies revealed elevated levels of carcinomas. Most published epidemiology studies reported an elevated risk of NHL associated with more frequent GBH use, as did all of the meta-analyses discussed during the trials. Dozens of published genotoxicity assays were reviewed that reported one or more positive assay, including a small number of important positive studies in exposed human populations.

43 For example, see Dr Portier’s expert report on glyphosate animal bioassay data at <https://usrtk.org/wp-content/uploads/2017/10/Chris-Portier-expert-report.pdf>; and Dr Reitz’s assessment of the GBH epidemiological database at <www.baumhedlundlaw.com/pdf/monsanto-documents/Hardeman/Beate-Ritz-Expert-Report.pdf>.

44 Access the SAP final report dated 16 March 2017 at <www.epa.gov/sites/production/files/2017-03/documents/december_13-16_2016_final_report_03162017.pdf>. The full, >1200-page transcript of the SAP meeting captures in great detail the many questionable aspects of the EPA’s assessment of glyphosate and GBH oncogenicity.

45 Portier’s 2020 published analysis (doi: 10.1186/s12940-020-00574-1) notes 37 significant tumour findings across 13 valid cancer bioassays, including multiple forms of malignant lymphomas relevant in the assessment of Roundup–NHL risk.
Throughout the trials, Bayer/Monsanto lawyers asserted that there was no evidence supporting a link between exposure to Roundup and NHL, despite the fact that experts for both plaintiffs and the defence had spent weeks discussing hundreds of discrete study findings supporting a possible role of Roundup in the initiation and/or progression of NHL. Explaining away such a diversity of experimental results supporting a role of GBH exposures in the plaintiffs’ NHLs was among the hurdles faced by the Bayer/Monsanto trial teams.

1. Differences in the EPA and IARC assessments of GBH exposures and oncogenicity

Beyond the EPA versus IARC interpretations of specific GBH–NHL relevant study findings, there were several other factors and considerations described to the three juries that likely played a role in tipping the weight of evidence:

1. The failure of Monsanto and the EPA to acknowledge or address the known higher risks following exposures to formulated GBHs compared to exposures to pure technical glyphosate.
2. The mostly registrant-generated genotoxicity studies and data relied on by the EPA, in contrast to the IARC’s reliance on genotoxicity studies published in peer-reviewed journals, including a significant number of studies focused on GBHs instead of just technical glyphosate.
3. The significant differences that exist in the levels of applicator GBH exposures among people applying Roundup via a handheld wand (as is the case with the first three sets of plaintiffs), in contrast to the operators of a modern pesticide sprayer with a glass–steel cab and air filtration system.
4. The virtual absence of requirements for personal protective equipment (PPE) on Roundup labels (including, for example, “wear gloves when mixing and loading or applying this product”).
5. The importance of multiple exposure episodes per year, over multiple years, including a certain percentage of high-exposure episodes caused by application equipment problems, leaky hoses and valves, spills, wind, spray patterns and equipment clean-up and repair.

Regarding Factor 1, substantial published data presented at trial support the conclusion that formulated GBHs are more toxic than pure technical glyphosate, especially GBHs containing polyethoxylated tallowamine (POEA)-based surfactants.46 Multiple internal Monsanto emails express the same conclusion, and they are cited and accessible via expert reports and trial testimony and exhibits. Nearly all GBHs sold in the USA by Monsanto since the mid-1970s have contained POEA-based surfactants, which have been phased out in GBHs in Europe due to safety concerns.47

46 For an overview, see C Benbrook “How Did the U.S. EPA and IARC Reach Diametrically Opposed Conclusions on the Genotoxicity of Glyphosate-Based Herbicides?”, supra, note 38.
47 Benbrook expert report in Hardeman, accessible at <https://usrtk.org/wp-content/uploads/2019/01/Benbrook-expert-report-November-2018.pdf>.
The most important reason why the POEA-based surfactants in US Roundup products are more toxic than technical glyphosate is the proclivity of the POEA in Roundup to increase dermal (and weed leaf) absorption rates. Once through the human epidermis, POEA-based GBHs also promote the movement of glyphosate through cell membranes, where it can then damage DNA via oxidative stress or some form of chromosomal or genetic aberration, including possibly epigenetic changes.48

Regarding Factor 2, the EPA’s assessment of the genotoxicity of Roundup and other GBHs was limited predominately to unpublished, registrant-submitted studies. Some 94 of the 95 registrant-submitted glyphosate and GBH genotoxicity assays were negative.49 Nearly half were reverse bacterial mutation studies (ie Ames tests) unlikely to produce positive results because bacteria lack mitochondria, and hence are not subject to oxidative stress caused by alterations in mitochondrial function, as is the case within mammalian cells.50

The IARC, however, focused on a much larger and more diverse body of published genotoxicity assays. Nearly 76% of the 191 assays reviewed by the IARC reported one or more positive genotoxicity response. The dramatic differences in the genotoxicity databases relied on by the EPA and IARC are summarised in Figures 2 and 3, and they explain in large part why the EPA and IARC reached such different conclusions in the case of glyphosate and GBH genotoxicity. Incidentally, over two dozen genotoxicity studies conducted with glyphosate, GBHs or both have been published since the EPA and IARC conducted their reviews; all but one reported one or more positive assay.51

Regarding Factor 3, the absence of high-quality mixer-loader and applicator exposure studies that compare various types of application equipment and varying combinations of PPE is one of the most glaring deficiencies in the overall glyphosate and GBH database. Exposure estimates cited or made by experts during the first three trials relied predominantly on the predictive operator exposure model (POEM)52 and a Monsanto-commissioned UK field study that partially relied on POEM simulations. The UK study estimated exposures to different parts of the body per hour of spraying by application method and type of equipment, by rate of application and conditions during the spray application, and in the presence/absence of PPE. It compared estimated exposures following six hours of spraying relative to levels of exposure regarded by UK regulators as acceptable. Multiple scenarios resulted in applicator exposures well over acceptable limits, especially when applications were made without gloves.

48 D Kubsad et al, “Assessment of Glyphosate Induced Epigenetic Transgenerational Inheritance of Pathologies and Sperm Epimutations: Generational Toxicology” (2019) 9 Scientific Reports 6372.
49 This and other comparisons of the EPA’s and IARC’s assessments of glyphosate and GBH genotoxicity are from C Benbrook “How Did the U.S. EPA and IARC Reach Diametrically Opposed Conclusions on the Genotoxicity of Glyphosate-Based Herbicides?”, supra, note 38.
50 Bailey et al, “Chronic Exposure to a Glyphosate-Containing Pesticide Leads to Mitochondrial Dysfunction and Increased Reactive Oxygen Species Production in Caenorhabditis elegans” (2018) 57 Environmental Toxicology & Pharmacology 46.
51 See table 4 in C Benbrook “How Did the U.S. EPA and IARC Reach Diametrically Opposed Conclusions on the Genotoxicity of Glyphosate-Based Herbicides?”, supra, note 38, for the list of 27 genotoxicity assays published after the release of the EPA and IARC reviews and through late 2018.
52 Access documentation and the POEM at <https://www.hse.gov.uk/pesticides/pesticides-registration/data-requirements-handbook/operator-exposure.htm>.
The exposure assessment branch of the EPA’s Office of Pesticide Programs has developed detailed surrogate reference values for worker exposures based on type of application, methods of application and PPE.53 While it is an imperfect basis for quantifying mixer-loader and applicator exposures, crude approximations of exposures to glyphosate following a workday across several common application scenarios can be calculated. Figure 4 compares simulated occupational exposures based on different application equipment and scenarios, noting as well the EPA’s most recent estimate of adult dietary exposures to glyphosate and acceptable daily exposures (glyphosate’s cRfD).54

Some insights are clear: handheld and backpack sprayers result in far higher exposures than in the case of tractors or spray rigs with glass–steel cabs and air filtration systems. On days when something goes wrong while applying a GBH through a handheld wand (a leaky valve, gusts of wind, pinhole leaks in a hose, excessive application rate) exposures can be vastly higher.

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53 Accessible at <www.epa.gov/pesticide-science-and-assessing-pesticide-risks/exposure-surrogate-reference-table-pesticide-risk>.

54 Dietary exposures across age groups are reported in the EPA’s assessment of glyphosate dietary exposures at <www.regulations.gov/document?D=EPA-HQ-OPP-2009-0361-0071>.
Regarding Factor 4, the EPA’s glyphosate oncogenicity risk assessment does not encompass, nor address high-exposure occupational scenarios. It does not take into account the added risks stemming from repeated use of a GBH during a spray season, nor users who spray for several hours per day. It does not consider the added risks facing people who use – and are exposed to – a GBH for many continuous years.

In addition, current glyphosate and GBH risk assessments do not take into account nor address several factors that can markedly heighten an individual’s GBH-related risks. Individuals with a compromised immune system or battling another form of cancer should exercise added caution when applying a GBH, or they should avoid occupational exposures altogether. Pregnant women should also take added precautions to avoid any exposure to a GBH, and indeed exposures to most pesticides. Such contraindications are common on drug labels, and for good reasons. Should these same reasons apply occupational exposures to pesticides and associated label precautions?

IV. IMPLICATIONS FOR PESTICIDE USE, RISK ASSESSMENT AND REGULATION

Intense global focus on glyphosate and GBH uses, exposures and risks has brought into sharp relief systemic shortcomings in pesticide hazard assessment, how exposures and risks are quantified and how excessive or rising risks are mitigated. The science base supporting the quantification of applicator and occupational exposures, and especially those using handheld wands to direct GBH sprays, is profoundly deficient. The uncertainty embedded in current GBH applicator dermal exposure estimates is a
striking example of how both the public and private sectors have failed to produce credible, real-world exposure and risk estimates.

There is growing agreement among independent pesticide risk assessment scientists, physicians and occupational health experts that pesticide product labels should disclose fully all of the ingredients in pesticide formulations. For example, Mesnage et al (2019)\textsuperscript{55} advanced multiple reasons why it is time to drop “Confidential Business Information” (CBI) protection of co-formulants, arguing that the need for policies and science advancing public and environmental health is more compelling than the need for pesticide formula CBI protection. Other policy tools and interventions can be deployed to reward innovation in pesticide formulation chemistry.\textsuperscript{56}

Pesticide manufacturers and their supporters argue that it is not feasible to put all pesticide formulations through a full battery of toxicological studies. I agree. But it would be feasible and beneficial to subject two or three distinctly different but common formulations of high-volume pesticides to genotoxicity, sub-chronic and, in some cases, chronic toxicity testing. When an initial set of such studies produces no basis for added concern, no further testing of similar formulations would be required. But when different formulations produce significantly different results, steps should be taken by industry and regulators to phase out higher-risk formulations, while continuing to invest in the search for lower-risk formulations.

Today’s human health and environmental impacts stemming from agricultural GBH uses and the emergence and spread of GBH-resistant weeds are inextricably bound to the scope and frequency of GBH uses. Impacts on monarch butterfly habitats, aquatic ecosystems, beneficial insect habitats, pollinators and human health are not linear relative to pounds of active ingredient applied in a given watershed or region. The scale and frequency of use of pesticides must become a variable that is considered by regulators when re-registering pesticides that have gained substantial market share. It is ridiculous to continue basing pesticide regulation on the myth that if one acre of wheat in Kansas can be safely treated with Pesticide X, then all acres of wheat in the state – or the world – can also be treated safely, year in and year out.

Political leaders and regulators should consider the merits of a simple, clear change in policy: pesticide companies and farmers must find ways to use herbicides such that residues in foods as eaten are rare, and, when detected, very low. In the case of many insecticides and fungicides, such a universal “no residues in food” goal is not achievable without significant costs and at least short-term disruption of pest management systems. But it surely is readily attainable in the case of herbicides, most of which are applied very early in the crop production cycle.

Glyphosate residues could be essentially eliminated from human foodstuffs by ending all pre-harvest crop desiccation uses, in conjunction with some additional restrictions on labelled GBH uses on RR crops. Collectively, such dietary exposure-driven limits and risk mitigation measures would likely have a modest impact on the role of GBHs in

\textsuperscript{55} R Mesnage et al, “Insight into the Confusion over Surfactant Co-Formulants in Glyphosate-Based Herbicides”, supra, note 7.

\textsuperscript{56} For some suggested policy alternatives, see C Benbrook, “Why Regulators Lost Track and Control of Pesticide Risks: Lessons From the Case of Glyphosate-Based Herbicides and Genetically Engineered-Crop Technology” (2018) 5 Current Environmental Health Reports 387.
agriculture, including in RR weed management systems, and no impact on applicator exposures and risk among those applying a GBH with handheld or small-scale equipment. Regulation-driven efforts to slow the emergence and spread of glyphosate-resistant weeds may eventually be put in place and, if meaningful, would impact GBH use more significantly than restrictions stemming from the need to get glyphosate out of the human food supply.

Last, the regrettable history of GBH testing and regulation points to the need for a much-expanded role for independent science – and scientists – in conducting studies that are essential to the pesticide risk assessment process. Monsanto seems to have an uncanny ability to design and conduct studies that are almost always negative using the same experimental methods that frequently result in positive studies when done by independent scientists. This highlights the need for greater diversity in who conducts the basic studies supporting the pesticide regulatory decision-making process.