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Toxic Environmental Chemicals: The Role of Reproductive Health Professionals In Preventing Harmful Exposures

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Abstract

Every pregnant woman in the U.S. is exposed to many and varied environmental chemicals. Rapidly accumulating scientific evidence documents that widespread exposure to environmental chemicals at levels encountered in daily life can adversely impact reproductive and developmental health. Preconception and prenatal exposure to environmental chemicals are of particular import because they may have a profound and lasting impact on health across the life course. Thus, preventing developmental exposures to environmental chemicals would benefit greatly from the active participation of reproductive health professionals in clinical and policy arenas.

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Condensation

Steps by reproductive health professionals to prevent developmental exposure to toxic environmental chemicals can contribute to a lifetime of health benefits for their patients.

Keywords

critical windows of development; developmental origins of health and disease; environmental chemicals; toxic chemicals; reproductive environmental health

Environmental Chemicals and Reproductive and Developmental Health

Among the U.S. population, current indicators of reproductive adversity include a decline in the age of onset of puberty;¹ declines in fertility and fecundity;^{2, 3} increased rates of poor birth outcomes such as babies born prematurely,^{4, 5} small for gestational age,⁶ and with certain birth defects;⁷ increased rates of childhood diseases such as autism,⁸ certain types of cancer,⁹ and obesity;¹⁰ and declines in life expectancy with some communities having life expectancies already well behind those of the best-performing nations.¹¹ Because these and other barometers of reproductive health and capacity have changed at a relatively rapid pace, they are unlikely to be explained by changes in genetic makeup.¹² Thus, we need turn our attention to other factors, including the environment, as possibly contributing to these trends.

The environmental contributors to reproductive health begin *in utero* and include the social, physical and nutritional environment, and physical and chemical agents. Each of these factors interacts with the others and with intrinsic biological factors, such as age, gender and genes, to influence individual and population health outcomes (Figure 1).^{13, 14} For example, environmental pollution interacts with stress to the detriment of long-term health,¹⁵⁻¹⁷ the effects of exposure to toxic chemicals can be exacerbated or mitigated by nutritional status,¹⁸⁻²⁰ and exposure to toxic chemicals and good nutrition is influenced by social and other environmental factors such as injustice, poverty, neighborhood, and housing.¹⁸⁻²⁵

Disparities in these environmental contributors are of major health consequence.²⁶⁻²⁸ Many communities with the highest exposures also lack access to medical care, good educational opportunities, good nutrition, employment, and other factors that may help to mitigate related impacts. Thus, the effect of a low dose exposure to an environmental chemical may be quite different depending on the populations degree of exposure to other environmental contaminants and underlying health status (Figure 2).²⁹

Within the field of obstetrics and gynecology, preconception and prenatal exposure to environmental chemicals (which is defined in this paper as including synthetic chemicals and metals) is a key area of inquiry because: (a) exposure to many and varied toxic chemicals among pregnant women in the U.S. is now the norm (Figure 3);³⁰ (b) developmental exposure to certain environmental chemicals is linked to a myriad of health consequences that can manifest across the lifetime of individuals and potentially be transmitted to the next generation (Table 1);³¹ and (c) preconception and prenatal exposure to environmental chemicals can be mitigated and prevented. This paper provides a brief overview of this new science relevant to the practicing obstetrician, gynecologist and other reproductive health professionals and outlines opportunities for preventing harm and associated costs in clinical and policy venues.

Exposure to Environmental Chemicals Among Pregnant Women

In the past 70 years, there has been a dramatic increase in human exposure to both natural and synthetic chemicals. Over this period, U.S. chemical production and use has increased over 16-fold.³² Today, more than 80,000 chemical substances are listed by the U.S. Environmental Protection Agency (EPA) as manufactured or processed in the United States, or imported into the country,^{33, 34} but this is probably an overestimate of the number of chemicals currently in commercial use. About 3,000 to 4,000 chemicals are identified as high volume chemicals, meaning that more than a million pounds of each of them are manufactured or imported annually.³⁴ Moreover, approximately 700 new industrial chemicals are introduced each year.³⁵

Health care professionals and the public cannot assume, as they do with pharmaceuticals, that adequate in vitro and in vivo testing of environmental chemicals has been undertaken and considered by regulatory agencies before widespread human exposure occurs (Figure 4). The vast majority of chemicals in commerce have entered the marketplace without comprehensive testing and standardized information on their reproductive or other chronic toxicities.^{36, 37} For example, in 1976 the U. S. Environmental Protection Agency (EPA) was given the authority to regulate chemicals in commerce under the Toxic Substances and Control Act (TSCA). EPA has used its authorities under the TSCA to require testing of fewer than 200 of the 62,000 chemicals in commerce when TSCA became law.³⁸

The inadequacy of our current regulatory framework for chemicals in commerce is recognized by physicians and organizations of health professionals such as the American Medical Association and the American Academy of Pediatrics,³⁹⁻⁴¹ governmental⁴² and non-governmental organizations,⁴³ and industry.⁴⁴

Toxic chemicals are currently widely distributed throughout homes, workplaces and communities, and contaminate food, water, air and consumer products. A 2011 study using population-based data from the National Health and Nutrition Examination Survey documented ubiquitous exposure among pregnant women in the U.S. to multiple chemicals.³⁰ The study found virtually all pregnant women have measured levels of all of the following chemicals that can be harmful to human reproduction and/or development in their bodies: lead, mercury, toluene, perchlorate, bisphenol A (BPA), and some phthalates, pesticides, perfluorochemicals (PFCs), polychlorinated biphenyls (PCBs) and polybrominated diphenol ethers (PBDEs) (Table 1).³⁰

Several of these environmental chemicals in pregnant women, including phthalates, mercury and PBDEs, are at levels associated with adverse health outcomes in human studies.³⁰ We have incomplete knowledge of what these exposures mean because the reproductive and other potential health impacts of daily exposure to this complex mixture of environmental chemicals have not been studied. This shortcoming is recognized by the National Academy of Sciences (NAS) to be a gap in current scientific methodologies that inform public policy that permits human exposure.²⁹ The NAS has also concluded that in the absence of data one cannot assume (as policy makers and regulators currently do) that there is a threshold or safe

limit of exposure for chemicals that adversely impact reproductive or developmental health outcomes.^{45,46}

Many chemicals in pregnant women can cross the placenta, and in some cases, such as methyl mercury, fetal exposure has been documented to be higher than maternal exposure.⁴⁷⁻⁴⁹ In 2010, the National Cancer Institute's President's Report on Cancer observed that “to a disturbing extent babies are born “pre-polluted.””⁵⁰ Postnatally, maternal exposure to environmental chemicals may continue to expose a newborn through breast-feeding.⁵¹⁻⁵³

Developmental Vulnerability to Environmental Chemicals

Assumptions about the benign nature of “low-level” environmental exposures have been upended by the new science.^{29, 54} We now know that the human reproductive system is particularly vulnerable to biological perturbations caused by ambient levels of environmental chemicals when these exposures occur during critical or sensitive periods of development i.e., *in utero*, and during infancy, childhood and adolescence.⁵⁵⁻⁵⁷ This vulnerability is in part because these are times of extensive developmental changes, such as cellular proliferation and rapidly changing and/or undeveloped metabolic, hormonal and immunologic capabilities.⁵⁸

For example, critical stages of central nervous system development occur from embryogenesis through adolescence. The periods of neuronal proliferation, migration, differentiation, and synaptogenesis are especially sensitive to disruption and permanent damage.^{59, 60} Since these processes are unidirectional, interference at an early stage may result in disruption throughout the further cascade of reactions and interactions which propagate human development.^{59, 60}

The range of potential adverse impacts from *in utero* exposure to exogenous chemicals is already well understood by clinicians familiar with thalidomide's congenital limb and gastrointestinal malformations,⁶¹⁻⁶³ and diethylstilbestrol's (DES's) delayed effects of benign and malignant reproductive tract abnormalities and increased risk of female breast cancer.⁶⁴⁻⁶⁶ DES remains one of the most scientifically robust illustrations of the linkage between developmental exposure to a hormonally active exogenous chemical and adult disease.⁵⁸

Of growing importance for patient health is that exposure of pregnant women to “endocrine disrupting chemicals” (EDCs) beyond DES has proliferated, such that simultaneous exposure to many EDCs is ubiquitous among pregnant women in the U.S. today.³⁰ The EPA defines EDCs as compounds that “interfere with the synthesis, secretion, transport, binding, action, or elimination of natural hormones in the body that are responsible for the maintenance of homeostasis (normal cell metabolism), reproduction, development, and/or behavior.”⁶⁷ Examples of EDCs commonly found in food, water, air, house dust, and/or personal care products include phthalates, BPA, PBDEs, perchlorate and some pesticides.⁶⁸ Because hormonal regulation is critical to human reproduction, chemicals that perturb the system may cause permanent effects.⁶⁹⁻⁷⁴

For example, PCBs and PBDEs can disrupt maternal thyroid function which is crucial for normal fetal development and in utero exposure to these chemicals has been associated with neurological deficits in human and/or animal studies.⁷⁵⁻⁷⁷ Phthalates can interfere with testosterone and studies in animals and humans indicate that exposure to certain phthalates during critical times of development can increase the risk of adverse male reproductive development – in rats, undescended testicles and cryptorchidism, and in humans, there is a relationship with subtle measures of feminization in boys for women who have higher phthalate exposures during pregnancy.⁷⁸

The mechanisms of action related to developmental exposure to environmental toxicants are many and complex and can change depending on when in the pregnancy or other developmental stage the exposure or related insult occurs.^{79, 80} For example, environmental chemicals can interfere with the development of normal fetal lung structure and function by perturbing a variety of transcription factors and morpho-regulatory molecules during critical developmental stages.⁷¹

Normal cell signaling can also be perturbed by EDCs, heavy metals and other environmental chemicals through epigenetic mechanisms, which, while not changing DNA, disrupt gene expression integral to orchestrating healthy human development.¹³ The relationship between the human genome and the environment has been analogized as genes acting to “load the gun” or create the potential for adverse health outcomes, and the environment acting as the “trigger” which activates the physiological or pathological network of biological reactions or events responsible for human health and disease.¹³ Environmental modifications of gene expression can affect embryonic imprinting, cellular differentiation, and phenotypic expression.⁸¹ Beckwith-Wiedemann, Prader-Willi and Angelman are three syndromes that exemplify the significance of epigenetics in real life.⁸²⁻⁸⁴

Human research has begun to expand mechanistic data from animal studies on the effect of environmental chemicals on the epigenome and human health.^{17, 85} However, as with pre-clinical testing of pharmaceuticals, non-human systems of evidence is the preferred method for documenting and developing prevention strategies related to the health impacts of developmental exposure to environmental chemicals because these studies can be undertaken prior to human exposure.⁸⁶ Environmental contaminants are not intended for human use, and it is unethical to knowingly expose humans to these chemicals under experimental conditions to assess for harmful effects.

Table 1 presents examples of the reproductive and/or developmental health effects from human studies of *in utero* exposure to environmental chemicals common in pregnant women today. Exemplary of these data, in 2009, the Endocrine Society reviewed the evidence of health impacts from endocrine disrupting chemicals and concluded that “the evidence for adverse reproductive outcomes (infertility, cancers, malformations) from exposure to endocrine disrupting chemicals is strong, and there is mounting evidence for effects on other endocrine systems, including thyroid, neuroendocrine, obesity and metabolism, and insulin and glucose homeostasis.”⁵¹

New scientific discoveries, i.e., epigenetics, cell signaling and developmental programming, document the vulnerability of the developing human to contemporary levels of environmental chemicals. Environmental exposures during fetal development may lead to changes in organ structure, function, and/or metabolism that are permanent and impact lifetime health risk. For the practicing clinician, the new science means that an important outcome of pregnancy is not just a healthy newborn but a human being optimally programmed for health from infancy through old age.

Implications of the New Science for Reproductive Health Clinicians

The nature and extent of the relationship between reproductive health and environmental chemicals is rapidly unfolding. The current strength of the evidence linking ubiquitous exposure to environmental chemicals to adverse reproductive and developmental health outcomes is sufficiently robust that leading scientists and reproductive health and other clinical practitioners have called for timely action to prevent harm.^{31, 50, 51, 56} Among physicians, obstetricians and gynecologists are uniquely poised to intervene in critical stages of human development (i.e., preconception and during pregnancy) to prevent harm.

Taking Action to Prevent Harm in Clinical Settings

Obstetricians and gynecologists can serve as a science-based source of guidance on how to avoid potentially adverse exposures.^{87, 88} As in other areas of clinical practice, communicating the science and areas of uncertainties about environmental chemicals can provide patients with the information they need to make informed choices based on the evidence, their values and preferences. Studies related to communicating the results of environmental chemicals in breast milk and other biomarkers lend empirical support to this approach.^{89, 90}

Pediatricians have long been attuned to the opportunity that clinical practice offers to identify, evaluate and counsel patients about preventing harm from hazardous environmental exposures. The American Academy of Pediatrics has had an environmental health committee for over half a century and publishes a clinicians' handbook for the prevention of childhood diseases linked to environmental exposures.⁹¹ In light of the importance of preconception and prenatal environmental exposures to the health of the pregnancy, and the child and adult that she or he will become, these pediatric approaches to incorporating environmental health into clinical care are equally applicable to reproductive health professionals. Based on our experience in clinical practice and through our engagement with health professionals, scientists and the public, many patients who are pregnant or thinking about becoming pregnant are intensely and justifiably interested in their environmental exposures; at the same time, other women of childbearing age are unaware of the risk of their exposures. Clinicians should intervene as early as possible to prevent exposures during pregnancy by alerting patients to potential hazards and providing guidance on how to avoid toxic exposures. By the first prenatal care visit, disruptions of organogenesis may have already occurred.

Taking an exposure history is a key first step. Clinicians should always ask women of childbearing age about occupational exposures; the workplace may be an important source

of toxic exposures among pregnant women and legal exposure limits for most workplace chemicals are not designed to protect against harm to a pregnancy or the developing fetus. A variety of examples of how to take an exposure history exist,⁹²⁻⁹⁵ and can be found at: <http://prhe.ucsf.edu/prhe/clinical/index.html#eh>

Clinicians should provide anticipatory guidance to all patients with information about how to avoid toxic exposures at home, in the community and at work. Information and resources about environmental hazards can be successfully incorporated into childbirth class course curriculum to help women and men make optimal choices for themselves and their children.⁹⁶ Patient-centered brochures with tips for preventing toxic exposures and links to many additional resources can be found at: <http://prhe.ucsf.edu/prhe/toxicmatters.html>

Patient-centered actions can reduce body burdens of toxic chemicals. Research documents that when children's diets change from conventional to organic food, the levels of pesticides in their bodies decline.⁹⁷ Likewise, recent studies found that avoiding canned food and other dietary sources of BPA can reduce measured levels of the chemical in children and adult family members,⁹⁸ and that short-term changes in dietary behavior may significantly decrease exposure to phthalates.⁹⁹ It is important to recognize, however, that decisions on the individual level about avoiding toxic exposures are complex and often affected by external factors that limit making healthier choices.¹⁰⁰

Patient purchasing patterns can also send a signal to the marketplace that can help drive society-wide change. This was demonstrated by the burgeoning market in organic food,¹⁰¹ the explosion of the market for alternatives to BPA in food contact uses such as baby bottles,¹⁰² and in Walmart's recent banning of a flame retardant found in hundreds of consumer goods from its supply chain.¹⁰³

In addition, while reproductive health professionals can be certain that the environment influences patient health, the idea of adding yet another topic to a clinician's "to-do" list is likely to seem daunting. The reality of severely constrained patient-contact time and lack of a reimbursement mechanism is compounded by the fact that medical education for obstetricians and gynecologists has thus far been largely devoid of training in reproductive environmental health beyond the dangers of alcohol, tobacco, and recreational drugs. However, reproductive health professionals do not need to be experts in environmental health to provide useful information to patients and make referrals when hazardous exposures are identified. Existing clinical experience and expertise in communicating risks of treatment are also largely transferable to environmental health.

Many useful resources exist to support clinicians in communicating about environmental risks.¹⁰⁴ The Pediatric Environmental Health Specialty Units (PEHSUs) are a network of investigators across the U.S. who support clinical capacity related to environmental health.¹⁰⁵ The PEHSUs respond to requests for information throughout North America on prevention, diagnosis, management, and treatment of environmentally-related health effects in children and as such, are poised to serve as a valuable resource for obstetricians and gynecologists in recognition of the inextricable relationship between reproductive and pediatric health.

Recent case examples in our (MM/TW/NS) experience include a woman who had a high blood lead and was 16 weeks pregnant. She had an evaluation by public health including a home visit without identifying a source and was referred to the PEHSU by her physician. We identified her use of an aruveydic medicine with a history of contamination with lead. We counseled her in general regarding possible health consequences for her baby and made her physician aware of the protocol for management of elevated blood lead in pregnancy. Another example was a mother and newborn identified as having elevated blood mercury. The PEHSU helped determine it was inorganic mercury and made the referral to the EPA region emergency response who identified the source of mercury as face cream.

Taking Action to Prevent Harm in Policy Settings

The role of clinicians in preventing exposure to environmental toxicants extends beyond the clinic or office setting.^{106, 107} Society-wide policy actions are essential for reducing toxic exposures to pregnant women and other vulnerable populations because many exposures are not controllable on an individual level, i.e., from air and water. In addition, environmental justice issues related to exposures to toxic substances cannot be sufficiently redressed by individual action. For example, women and men exposed to pesticides at work and in agricultural communities incur substantively higher exposures than the U.S. population overall.^{108, 109}

There are many examples that demonstrate that clinicians are in an excellent position to take action in policy settings. For example, our industrialized food system is associated with many and varied threats to reproductive and developmental health, including exposure to pesticides, chemical fertilizers, hormones in beef cattle, antimicrobials in beef cattle, swine and poultry, fossil fuel consumption and climate change, toxic chemicals in food packaging and cookware, and the production and promotion of food that is unhealthy for pregnant women.¹¹⁰ Policy interventions by the health care sector and physician' patient engagement offer mutually reinforcing opportunities to advance a healthy food system as a strategy to prevent adverse reproductive health impacts.¹¹⁰

To this end, physician leaders have been instrumental in spurring efforts by healthcare institutions to support the development of urban agriculture programs, farmer's markets and local food sourcing outlets to increase accessibility to healthier foods, and healthcare institutions have undertaken procurement policies to create a sustainable and healthy food service model. Nearly 350 hospitals have taken the Healthy Food in Healthcare Pledge in support of these efforts.¹¹¹ Because the food system purchasing power of the healthcare system is so large---about \$12 billion annually--- clinicians becoming engaged in changing their hospital food system procurement patterns can help leverage food system change more broadly. Other examples of institutional policy arenas for clinical action include the reduction of toxic chemicals in healthcare purchasing coupled to bringing policy gaps that impede less toxic procurement patterns to the attention of decision-makers. (See: <http://www.saferchemicals.org/resources/business/kaiser-permanente.html>) Clinicians have also been engaged in reducing the use of pesticides in institutional pest-control polices. (See: <http://www.mdpestnet.org/projects/ipmHealthcare.html>)

Clinicians can also work for towards policy change in their professional organizations. For example, professional organizations of physicians including obstetricians and gynecologists have been active in calling for regulatory and other efforts to address exposure to toxic chemicals and many other environmental threats to human health. A compilation can be found at: <http://www.prhe.ucsf.edu/prhe/pdfs/ProfessionalStatementsDatabase.pdf>.

In 2009, the Endocrine Society issued a position paper calling for improved public policy to identify and regulate EDCs, and finding that “[u]ntil such time as conclusive scientific evidence exists to either prove or disprove harmful effects of substances, a precautionary approach should be taken in the formulation of EDC policy.”⁷⁴ The application of the precautionary principle in environmental health dates to the 1980s and today precaution is an underlying principle of environmental health policy in the European Union, particularly in the realm of risk management.¹¹² The precautionary principle is defined, “When an activity raises threats of harm to human health or the environment, precautionary measures should be taken even if some cause and effect relationships are not fully established scientifically.”¹¹³ Reversing the burden of proof so that chemical exposures are not presumed safe in the absence of scientific evidence⁴⁶ would exemplify a precautionary approach to environmental chemicals.

Future Directions

Just as the thalidomide tragedy led to strengthened regulatory oversight of the safety and efficacy of all prescription drugs,⁶¹ recent advances in toxicity testing,^{79, 114-119} risk assessment,^{29, 46, 54, 120, 121} and in efforts to address shortcomings in regulatory policy related to chemicals in commerce,^{42-44, 122} are likely to create important change in the amount, type and availability of chemical toxicity data and related health impacts. These anticipated improvements underscore the need for a methodology to ensure timely application of these data to prevention. To this end, a methodology has been developed to evaluate the quality of evidence and strength of recommendations about the relationship between the environment and reproductive health in uniform, simple, and transparent summaries that integrate best practices of evaluation in environmental and clinical health sciences.¹²³ The generation of clinical guidelines needs to proceed with the development and dissemination of validated methods to screen and counsel patients about their exposures and safer alternatives that will prevent exposure for all patients.

It is also expected that electronic medical records will revolutionize medical research by facilitating instant, comprehensive, data that go back years into history and extend longitudinally into the future.¹²⁴ Harnessing these changes could greatly accelerate the creation of knowledge about the impact of the environment on our reproductive health and capacity. Obstetricians, gynecologists and other reproductive health professionals can play a groundbreaking role by intervening in critical stages of human development to translate the new science into healthier pregnancies, healthier children and healthy future generations.

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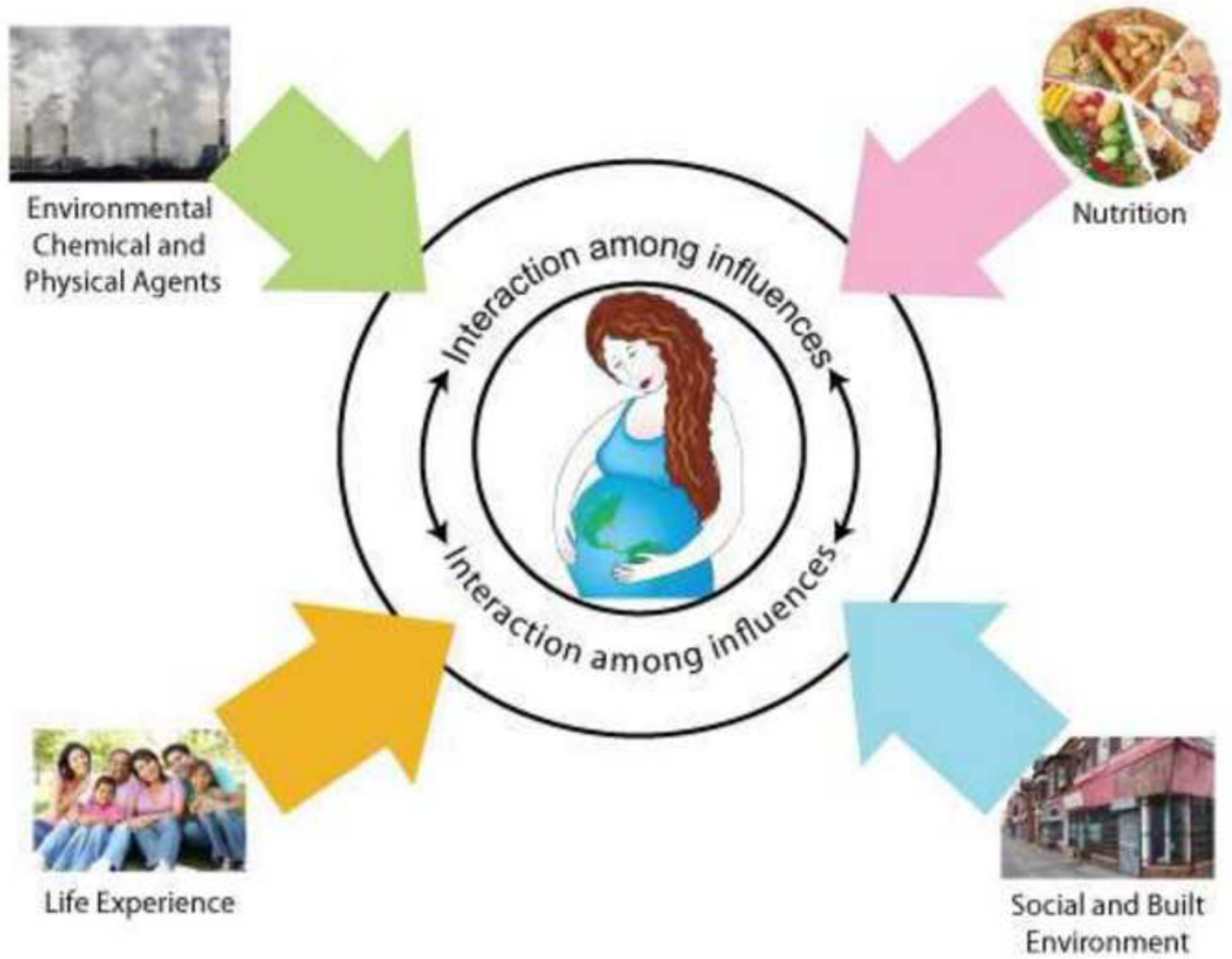


Figure 1. Environmental Influences on Reproductive Health Source: ¹¹⁰ Permission Needed

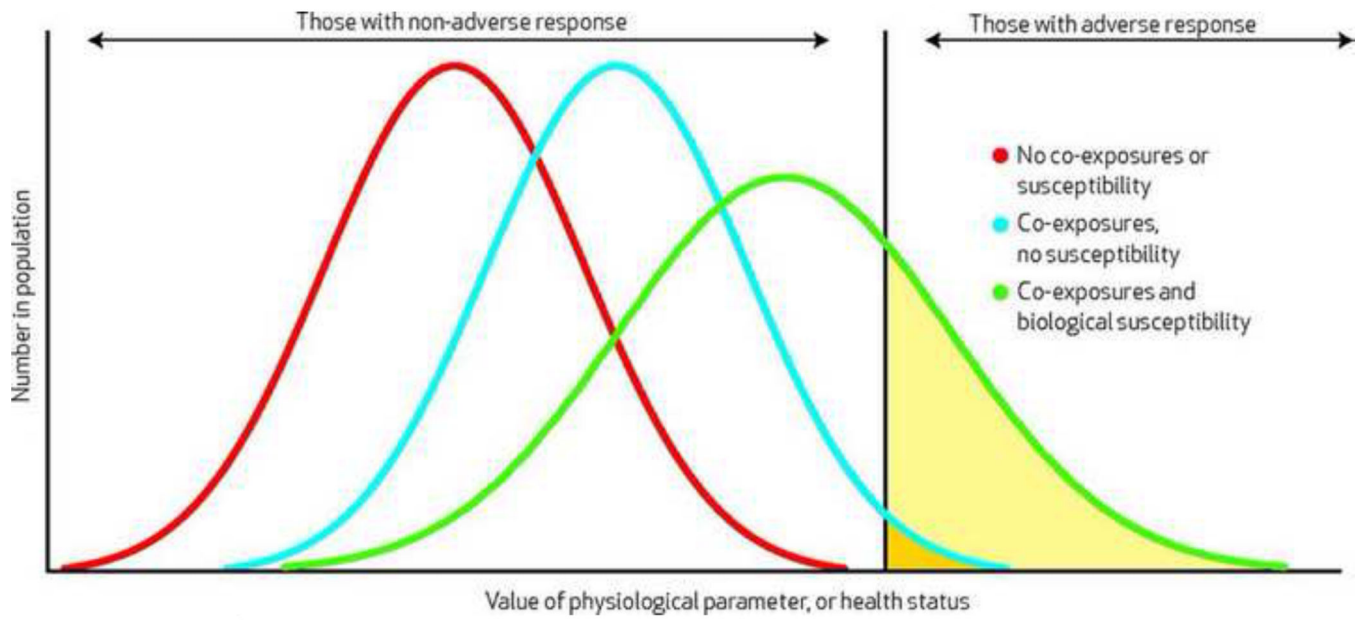


Figure 2.
The Effect of Biological Susceptibility and Co-exposure to Other Chemicals on the Relationship Between Individual Chemical Exposure and Adverse Health Outcomes
Source: ⁴⁶ Permission needed

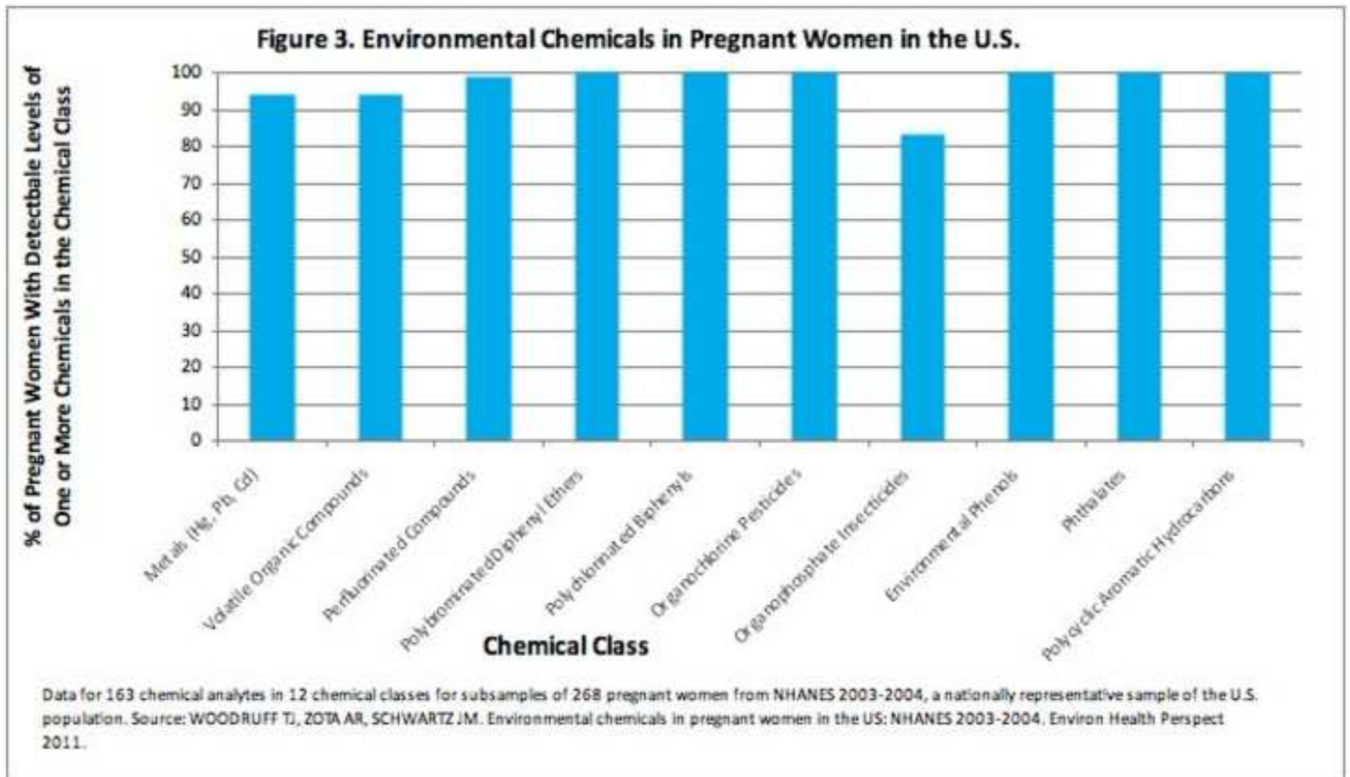


Figure 3. Environmental Chemicals in Pregnant Women in the U.S. Source: Adapted

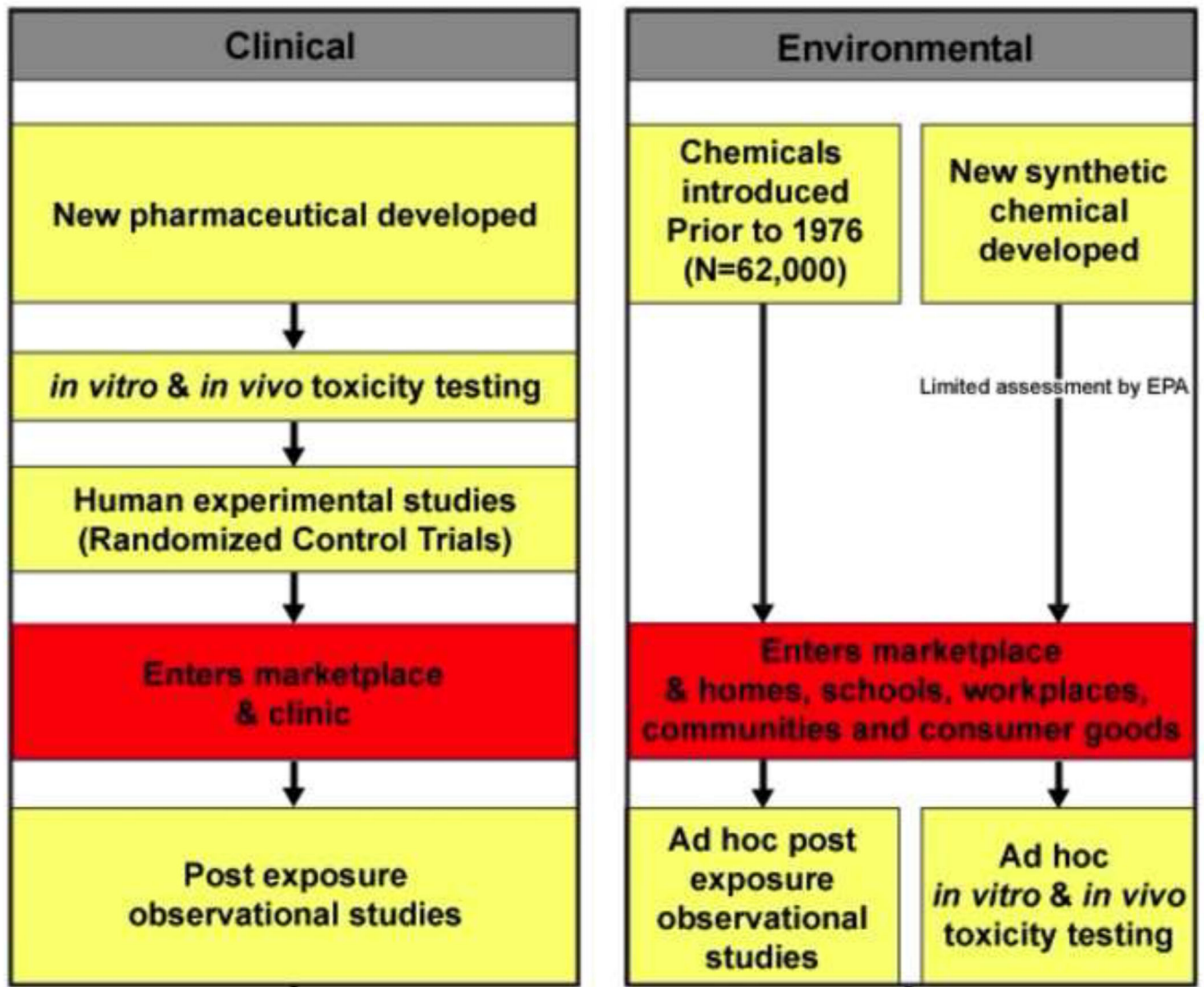


Figure 4.
 Comparison of the evidence streams needed in clinical and environmental health sciences for an exogenous chemical to enter the marketplace
 Source: Adopted from ¹²³ Permission needed

Table 1

Examples of Reproductive Health Impacts of Prenatal Exposure to Environmental Contaminants

Chemical	Exposure Sources and Pathways	Reproductive/Developmental Health Impact
Bisphenol A (BPA)	Chemical intermediate for polycarbonate plastic and resins. Found in consumer products and packaging. Exposure through inhalation, ingestion, and dermal absorption.	Recurrent miscarriage ⁸⁷
		Aggression and hyperactivity in female children ⁸⁸
Lead	Occupational exposure occurs in battery manufacturing/recycling, smelting, car repair, welding, soldering, firearm cleaning/shooting, stained glass ornament/jewelry making; non-occupational exposure occurs in older homes where lead-based paints were used, in or on some toys/children's jewelry, water pipes, imported ceramics/pottery, herbal remedies, traditional cosmetics, hair dyes, contaminated soil, toys, costume jewelry.	Alterations in genomic methylation ⁸⁵
		Increased likelihood of allergies ⁸⁹
Mercury	Mercury from coal-fired power plants is largest source in the U.S. Primary human exposure by consumption of contaminated seafood.	Reduced cognitive performance ^{90, 91}
		Impaired neurodevelopment ^{92, 93}
Polybrominated diphenylethers (PBDEs)	Flame retardants that persist and bioaccumulate in the environment. Found in furniture, textiles, carpeting, electronics and plastics which are mixed into, but not bound to foam or plastic.	Impaired neurodevelopment ⁹⁴
		Premature delivery, low birth weight and stillbirth ⁹⁵
Polychlorinated Biphenyls (PCBs)	Used as industrial insulators and lubricants. Banned in the 1970s, but persistent in the aquatic and terrestrial food chains resulting in exposure by ingestion.	Development of ADHD associated behavior ⁹⁶
		Increased BMI ⁹⁷
		Reduced IQ ⁹⁸
Perfluorochemicals (PFCs)	Widely used man-made organofluorine compounds with many diverse industrial and consumer product applications. Examples are perfluorooctane sulfonate (PFOS) and perfluorooctanate (PFOA), which are used in the manufacture of non-stick Teflon® and other trademark cookware products and in food-contact packaging to provide grease, oil and water resistance to plates, food containers, bags, and wraps that come into contact with food. They persist in the environment. Occupational exposure to workers and general population exposure by inhalation, ingestion, and dermal contact	Reduced birth weight ⁹⁹
Perchlorate	Used to produce rocket fuel, fireworks, flares and explosives and can also be present in bleach and in some fertilizers. Primary pathway for exposure is through drinking water due to contaminated runoff.	Altered thyroid function ¹⁰⁰
Pesticides	Applied in large quantities in agricultural, community and household settings. In 2001, over 1.2 billion pounds of pesticide active ingredients were used in the US. Pesticides can be ingested, inhaled and absorbed by the skin. The pathways of pesticide exposure include food, water, air, dust, and soil.	Impaired cognitive development
		Impaired neurodevelopment ^{101, 102}
		Impaired fetal growth ¹⁰³
		Increased susceptibility to testicular cancer ¹⁰⁴
Phthalates	Synthetically derived, phthalates are used in a variety of consumer goods such as medical devices, cleaning and building materials, personal care products, cosmetics, pharmaceuticals, food processing, and toys.	Childhood cancers ¹⁰⁵
		Reduced masculine play in boys ¹⁰⁶
		Reduced anogenital distance ¹⁰⁷
		Shortened gestational age ¹⁰⁸

Chemical	Exposure Sources and Pathways	Reproductive/Developmental Health Impact
	Exposure occurs through ingestion, inhalation, and dermal absorption.	Impaired neurodevelopment in girls ¹⁰⁹
Toluene	Exposure occurs from breathing contaminated workplace air, in automobile exhaust, some consumer products paints, paint thinners, fingernail polish, lacquers, and adhesives.	Decreased fetal and birth weight ¹¹⁰
		Congenital Malformations ^{111, 112}

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