Scientific Judgement and Toxic Torts - A Primer in Toxicology for Judges and Lawyers

David L. Eaton
I. GENES, ENVIRONMENT AND DISEASE

Remarkable progress has been made in the past decade in understanding the molecular basis of many chronic diseases such as cancer, degenerative neurological diseases (Alzheimer’s, Parkinson’s), heart disease, and asthma. Although the molecular basis for such diseases has become more apparent, the exact “cause” is seldom identified for a disease in general, and especially for a disease in an individual. It is now recognized, however, that most such chronic diseases result from a complex interplay between our genes and our environment. While our parents predetermine our genes, our environment is somewhat controllable, and thus identifying “environmental risk factors” for chronic diseases holds great promise for disease prevention. It should be noted that “environment” in this context represents virtually everything in the world around us that is not “in our genes.” Thus environmental factors include lifestyle choices such as smoking, drug use and alcohol consumption, exposure to infectious agents (viruses, bacteria), as well as diet and nutrition, environmental pollution (air, water), and even behavioral and social factors such as exercise, reproductive choices, sexual activity, etc.

There is currently great scientific effort committed to

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identifying specific genetic characteristics (so-called “genetic polymorphisms”) that make one individual more susceptible to something in the environment than another.¹ This area of research is sometimes referred to as “ecogenetics,” or the study of “gene-environment interactions.”² There are many classic examples of genetic characteristics that make an individual sensitive to something in his environment. For example, the rare genetic disease, phenylketonuria (PKU), makes individuals with this genetic trait very sensitive to a normal component of our diet—phenylalanine. Phenylalanine is a normal building block of proteins, and in most people is an important nutrient in the diet. For a small part of the population with a mutation in the PKU gene, however, regular “doses” of phenylalanine found in the normal diet can lead to serious mental retardation if infants are exposed to phenylalanine. Because of this, genetic testing for PKU is mandatory in most states in the United States and is part of normal newborn screening. Those rare individuals who test positive for PKU can lead normal lives by following special diets and avoiding foods rich in phenylalanine.

Another common example of a “gene-environment” interaction occurs in many people of Asian descent who carry a genetic variant of a gene involved in alcohol metabolism. Normally, alcohol is fairly rapidly detoxified in the liver. But individuals with a variant form of the gene for an enzyme called “aldehyde dehydrogenase” (ALDH2) are less able to eliminate a toxic by-product of alcohol metabolism, acetaldehyde. If a person with the variant ALDH2 gene consumes even modest amounts of alcohol, toxic amounts of acetaldehyde can accumulate in the blood, causing a very uncomfortable reaction (“flushing” of the skin from vasodilatation, nausea, headache). Not surprisingly, alcoholism and alcohol-related diseases such as cirrhosis of the liver occur very


There is currently a great deal of interest in identifying common genetic traits that might combine with factors in our environment to cause disease. It is hoped that, one day, physicians will be able to characterize, or “genotype,” the entire genetic code of a person, and based on the results (kept on a personal microchip medical card), identify whether the patient is at increased risk for certain diseases and potentially identify specific dietary, workplace, or other environmental factors that should be avoided to lower risk. While we are still a decade or more away from having scientifically validated tests for “environmental susceptibility” to most environmental/occupational hazards, similar approaches for identifying how individuals respond to therapeutic drugs is just around the corner (the field of “Pharmacogenomics”). Indeed, there are now several relatively widespread genetic tests that can identify in advance patients who are likely to have adverse reactions to otherwise “normal” therapeutic doses of specific drugs.3 The concept of “designer drugs” is becoming a reality, but so far in a limited way. For example, there is a common genetic variant in a gene called “N-acetyl transferase.” This gene is involved in the detoxification of a variety of therapeutic drugs, and people with the “slow” genetic variant exhibit increased toxicity (but also enhanced therapeutic effects at lower doses) to a variety of common drugs. Knowing this predisposition in advance allows physicians to prescribe the proper dose.

How will such genetic information be used in the courtroom? In the realm of genetic testing for drug sensitivity, there will be medical malpractice claims filed against physicians who fail to order genetic tests before prescribing certain drugs, once such procedures become the standard of care.4 Drug companies will

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attempt to increase drug safety and limit liability by identifying in advance drugs that may elicit adverse responses in small segments of the population because of genetic sensitivity. In the environmental and occupational arena, employers might use genetic tests as a way of identifying and removing “sensitive” individuals from certain workplace exposures. While such practice might conceivably lower the occurrence of chemical-induced occupational diseases, it is obviously also a means of employment discrimination. It is also likely that plaintiff and defense attorneys will utilize genetic susceptibility as an argument for, or against, causation in toxic tort cases. Currently, however, the scientific data supporting the use of genetic susceptibility information in toxic tort litigation is extremely limited. In the vast majority of circumstances, specific and measurable genetic “susceptibility markers” often do little more than shift a person “up” or “down” the dose-response curve. Such differences tend to be modest (less than a two-fold difference in susceptibility), and the impact of the genetic trait is often lost in the “noisy background” of poor exposure assessment. That is, if one can only “guess” the dose, duration, and frequency of exposure to a specific chemical within a factor of 5 or 10 (not uncommon in toxic tort cases), a genetic factor that theoretically doubles or halves the risk from a given dose will not be particularly informative against the high level of uncertainty of the actual “exposure.” Thus, although genetic information will increasingly find its way into toxic tort


litigation, the fundamental concepts of toxicology and epidemiology continue to serve as the foundation for establishing causation in toxic tort claims. The following information is provided as a “primer” in basic toxicology, as it relates to toxic tort litigation. For a more detailed discussion of considerations of how the science of toxicology and epidemiology should be used in the courtroom, the reader is referred to the Federal Judicial Center’s *Reference Manual on Scientific Evidence*. This publication includes chapters devoted to toxicology and epidemiology, as well as medical testimony and use of DNA in the courtroom.

II. BASIC TOXICOLOGY RELEVANT TO TOXIC TORT LITIGATION

Toxic substances may take many forms, including both human-made (synthetic) and natural chemicals. Although the adverse effects of physical agents such as ionizing radiation fall under the broad rubric of toxicology, this discussion will focus on chemical agents. There are many “sub-disciplines” within the field of toxicology, and a variety of approaches and techniques are used to evaluate the toxicological characteristics of chemicals. A detailed review of the basic principles of toxicology is beyond the scope of this article. The following brief review highlights some of the key principles of toxicology that must be considered in any attempt to establish whether a chemical exposure was causally related to a specific adverse effect or disease in an individual.

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A. Types of Adverse Effects Produced by Chemicals

Virtually all substances are capable of inducing some form of toxic effect, and the type and nature of effects will vary depending on the
dose (amount of substance that finds its way into the body)
• route (i.e., oral, inhalation, skin; injection)
• duration (days, weeks, months, years) and
• frequency (how many times per day, week, month, year)
of exposure.

A given chemical does not cause every possible effect, and the ability of a chemical to cause a particular effect depends upon a variety of factors, as discussed below. Typically, a specific chemical elicits a characteristic pattern of toxic (adverse) effects, although the appearance of specific effects will depend on the dose and other characteristics of exposure. Sometimes chemicals of a common type cause a generalized adverse response. For example, nearly all organic solvents derived from petroleum products (including mixtures such as gasoline or kerosene, or individual solvents such as benzene, hexane, or toluene) share some (but not all) symptoms in common: “defatting” of the skin following dermal exposure, and central nervous system depression (inebriation, loss of consciousness) following relatively high levels of inhalation exposure. However, even though different chemicals of the same general type (e.g., solvents) may have some common effects, they may also differ dramatically in other effects. For example, the industrial solvents benzene and toluene are very similar chemically, and share many common toxic effects noted above for solvents, but benzene is toxic to the bone marrow and can increase the risk of leukemia in workers, whereas these serious toxic effects have not been found for toluene. Thus, some chemicals act in very specific ways at the cellular level, and their effects may be largely limited to a characteristic type of response. As an example, the widely-used class of insecticides known as “organophosphates” inhibit a specific enzyme in the nervous system (acetylcholinesterase), and most of the signs and symptoms of toxicity can be attributed to this one mode of action. However, even small differences in chemical structure can sometimes make
very large differences in the type of toxic response that is produced. This is especially true for chemicals that cause birth defects (teratogens) or chemicals that increase the risk of cancer (carcinogens).

B. Concepts of Dose and Exposure

“Dose” refers to the amount of chemical that enters the body. The units of dose are typically expressed as an amount of substance per kg of body weight (mg/kg bw). Thus, if a 132 lb woman (60 kg) absorbed 60 milligrams of a chemical in a glass of contaminated water, she would have a dose of 1 mg/kg bw. Dose is the single most important factor to consider in evaluating whether an alleged exposure caused a specific adverse effect. Indeed, the basic dictum of toxicology was stated by the Sixteenth Century Physician/Philosopher, Paracelsus, considered the “father of toxicology”: “All substances are poisonous—there is none which is not; the dose differentiates a poison from a remedy.”\(^{11}\)

1. Relationship between “Exposure Concentration” and “Dose”

Dose and “exposure” in terms of media (e.g., air, water, soil) concentration are related, but not identical, terms. Exposure may be referred to as the presence of a chemical in a medium (e.g., air, water, food) that allows for direct contact with potential sites of absorption (e.g., gastrointestinal tract, lungs, skin). The units of such exposure are usually expressed as concentrations—e.g., milligrams of chemical per liter of water (mg/L), milligrams of chemical per cubic meter of air (mg/m\(^3\)), milligrams of chemical per kilogram of food (mg/kg). Frequently, such concentrations are expressed as “parts per million” (ppm) or “parts per billion” (ppb). For chemicals dissolved in water, 1 part per million is the same as 1 milligram of chemical dissolved in 1 liter of water. One part per billion (1 ppb) is a thousand times less—1 milligram dissolved in a thousand liters of water, or 1 microgram of chemical dissolved in 1

liter of water. A part per trillion (ppt), is 1,000 times less than a part per billion. To provide some perspective on these units, consider the following:

- 1 ppm = 1 penny in $10,000, or 1 inch in the distance of 15.8 miles
- 1 ppb = 1 penny in $10 million, or 1 inch in 15,800 miles
- 1 ppt = 1 nickel of Bill Gates net worth (assuming $50 billion), or 6 inches in the distance between the earth and the sun.

Analytical tools developed in the past several decades make it possible to measure substances in water, food, soil or air at the ppt and even parts per quadrillion (ppqd; 1,000 times less than a ppt). It is evident from these simple analogies that, when discussing exposure to chemicals in drinking water, air, or soil, it is critically important that the relationship between exposure, as expressed as a concentration of a pollutant in a medium (measured in ppm, ppb, ppt or even ppqd) and the actual dose to a person not be lost. The science of toxicology can help understand whether the dose of a substance achieved following a particular exposure has any relationship to toxicity or disease.

2. Frequency and Duration of Exposure

Frequency and duration of exposure are important elements of “dose.” Effects caused by chemicals may differ depending on whether exposure was short-term (e.g., acute, single dose or a few days) or long-term (chronic, repeated over years). The dose of a chemical required to produce health effects also differs with frequency and duration of exposure. When exposure occurs repeatedly over weeks, months, or years, the dose is usually expressed as a dose rate, with units of mg of chemical per kg of body weight per day. The dose necessary to produce deleterious effects with short-term exposure is higher than the dose that produces toxic effects when repeated over a long time period. The body can usually tolerate or recover from high doses with brief short-term exposure as compared to long-term repeated exposure. For example, one night of moderate drinking may give you no more than a headache the next day, but heavy drinking frequently
for years could lead to liver cirrhosis and possibly liver cancer. However, it is also possible that repeated, low dose exposures—even for many years—will have no consequence at all, since the body is often able to completely detoxify low doses before they do any damage. This concept of “thresholds” will be discussed in more detail later.

Most chemicals that have been identified to have “cancer-causing” potential (carcinogens) do so only following long-term, repeated exposure for many years. Single exposures or even repeated exposures for relatively short periods of time (e.g., weeks or months) generally have little effect on the risk of cancer, unless the exposure was remarkably high and associated with other toxic effects. Relatively infrequent exposure may also have negligible health consequences even if continued over time because of recovery between doses.

3. Pathways and Routes of Exposure

“Pathways” are the means by which an environmental chemical may reach an “exposed” person. Chemicals can enter the body by four fundamental “routes”: (1) via oral exposure (e.g., ingestion of the toxic substance directly, or in food or drinking water), (2) via inhalation (e.g., breathing air or inhaling dust contaminated with the toxic substance), (3) via direct contact with the skin (e.g., spilling of a pesticide mixture on the skin), or (4) by direct injection into the body (e.g., introduction of a drug by intravenous injection). The “bioavailability”—or ability of the chemical to be taken into the blood stream—differs by route of entry. Most drugs and toxic chemicals will be well absorbed from the gut when ingested in a soluble form versus in other media such as soil. Many chemicals, however, are only slightly absorbed, if at all, if applied to the skin. However, fat-soluble chemicals in high concentration may be well absorbed across the skin, and this can lead to an important pathway of exposure for those using concentrated solutions in the workplace. The extent of inhalation absorption of chemical vapor will depend on a variety of factors, including the relative solubility of the chemical in blood versus air, the rate of breathing, and even whether one breathes through the nose or
mouth.

4. Site of Action in the Body

Ultimately, what matters is the actual concentration of a toxic substance at the “site of action” in the body. The concentration of a chemical in any given organ/tissue in the body is determined by complex interactions between the rates of exposure, and rates of absorption, distribution, metabolism, and excretion. Because chemicals differ in their solubility in body fluids/tissues, in how they are metabolized, and in what cellular processes are altered, toxic effects of a chemical may be limited to specific tissues or organs, referred to as its “target tissue.” For example, lead and mercury typically produce toxic effects associated with the brain and kidneys, whereas certain chlorinated solvents such as chloroform and carbon tetrachloride affect predominantly the liver (although in high doses they also affect the brain).

Many factors determine whether a chemical will be toxic to a particular organ. Some organs metabolize (biotransform) chemicals to toxic intermediates, leading to toxicity in that organ. In such instances, the relative ability of an organ or tissue to metabolize the chemical may determine whether the toxic effect is seen in that tissue. Certain tissues may also accumulate a chemical from the bloodstream at higher rates than other tissues, leading to toxicity in just that tissue. This is particularly true for tissue with a function (e.g., liver), but not necessarily for storage tissue, i.e., fat, which accumulates fat-soluble chemicals such as DDT, but is not directly injured. Metabolic pathways and the amount and type of toxic by-products produced or accumulated may also differ depending on the amount of chemical in the bloodstream (which, of course, is directly related to dose). For example, metabolic pathways at low doses that result in chemical detoxification may be overwhelmed at high doses leading to accumulation of toxic intermediates or production of greater amounts of toxic by-products by alternative pathways.
5. **Dose-response Relationship**

As noted above, the relationship between dose and effect (dose-response relationship) is the hallmark of basic toxicology. The “dose-response” in a given individual describes the relationship between the magnitude or severity of the effect(s) and the dose. In many instances, especially for acute toxicity, the slope of the dose-response curve is quite steep. That is, once a sufficient dose has been achieved to induce a toxic response, further increases in the dose may produce large increases in the response. In the individual, the nature of the response may change with increasing dose. For example, ingestion of one or two glasses of wine will result in an apparent “stimulatory” effect on the nervous system, often expressed as slight changes in personality or character. Further consumption of alcohol will lead to loss of coordination and reaction time, slurred speech, etc. Continued consumption of alcohol beyond this level of intoxication may result in loss of consciousness and even death.

Although individuals within a population may respond differently to the same dose of chemical, the reaction of the population as a whole nevertheless follows a “dose-response relationship” such that the number of people in a population that respond to a chemical exposure increases with dose. Inherent in this concept is that, for the vast majority of chemicals and types of responses, there are doses below which no individual will respond (e.g., a “threshold”) and doses above which nearly everyone responds. For example, no one would exhibit any detectable adverse effect of a few drops of wine or beer (e.g., the dose is below the threshold), yet most everyone in a population would show signs of intoxication after ingestion of an entire bottle of wine (over a relatively short period of time). In between these two extremes, there are clearly differences in the level of intoxication between individuals consuming one, two, three, or four glasses of wine. In a similar fashion, there is inherent human variability in response to chronic exposures to chemicals. Dose-response relationships in populations also exist for both acute and chronic exposures to toxic substances.
6. Concept of “Thresholds”

For most types of dose-response relationships following chronic (repeated) exposure, thresholds exist, such that there is some dose below which even repeated, long term exposure would not cause an effect in any individual. Most toxicological responses, including neurological, reproductive, and developmental effects, exhibit thresholds (e.g., there is a dose below which the probability of an individual responding is essentially zero). One key objective in toxicology is to identify doses for a population below which no one will respond. However, in the case of chemical carcinogens, particularly those that increase risk of cancer by causing direct damage to DNA in cells, many regulatory agencies assume that there are no “thresholds,” and that risk is proportionate to dose at all levels of exposure—e.g., as the dose of carcinogen increases, the probability of developing cancer increases in a proportionate, “linear” fashion.

Nonetheless, many scientific and practical reasons indicate that, at very low doses, the significance of such risks, if real, become trivial and are lost in the background of other daily risks. For example, it is well known that cigarette smoking is strongly associated with increased risk for lung and bladder cancer (and other types), and that the probability of developing such smoking-related cancers is related to both the amount (cigarettes per day) and the frequency (years of smoking) of smoking over a lifetime.\(^\text{12}\) It is also recognized that the carcinogenic properties of cigarette smoke are strongly related to the ability of components of cigarette smoke to damage DNA (cause mutations), and thus it might be assumed that the dose-response relationship for smoking would be a “non-threshold” (linear at low doses) response.\(^\text{13}\) However, while a linear, non-threshold response to cigarette smoke may be hypothesized on theoretical grounds, from a practical


perspective one’s level of increased risk from smoking one cigarette over a lifetime, or even one cigarette a month for a lifetime, is not likely to be distinguishable from “background” risk for cancer from all other causes, known and unknown.

Not all chemical carcinogens increase cancer risk by causing mutation. For such “non-genotoxic” carcinogens, it is generally thought that the dose-response relationship follows a typical threshold-type response. Thus, it is often important to distinguish between “genotoxic” (particularly those that act directly on DNA to cause mutations) and “non-genotoxic” carcinogens for regulatory and risk assessment purposes. Practical thresholds may also exist for “genotoxic” carcinogens that damage DNA by indirect mechanisms (e.g., production of sufficient “reactive oxygen species” to cause oxidative damage, or sufficient inhibition of DNA repair mechanisms), because a sufficient amount of the chemical is needed before enough damage to the DNA occurs to lead to cancerous cells.

C. Chemical Exposures and Chronic Diseases

Traditional toxicology tests in laboratory animals are designed to identify toxic responses following various periods of exposure. Acute toxicity studies examine the toxic effects after single, high doses and are useful to understand the specific organ systems affected by the chemical, as well as the general “potency” of its effect (e.g., does it require microgram, milligram, or gram quantities to produce evidence of toxicity?). Additional “sub-chronic” (usually ninety days of daily exposure) and “chronic” (usually lifetime, or two years of continuous daily exposure) studies are often done to further examine whether the chemical is capable of causing other types of toxic effects following repeated exposures. Such studies may demonstrate that repeated exposure to a chemical could cause liver or kidney or brain damage, for example. Special “3-generation” studies may be done in animals to determine if the chemical can cause reproductive effects and/or

14 CASARETT AND DOULL’S TOXICOLOGY: THE BASIC SCIENCE OF POISONS, supra note 11, at chs. 2, 11-34.
birth defects. Today, new chemicals entering commerce, such as a new pesticide, may be subjected to specialized tests to determine if it can cause neurological effects on learning and behavior, or cause toxicity to the immune system. Each of these toxicological endpoints can be the subject of toxic tort litigation. However, regardless of the end-point, the basic concept of “dose-response” remains essential in evaluating a causal connection between an alleged exposure and a particular disease. As noted above, for non-cancer endpoints, it is generally accepted that “thresholds” exist, and that doses below the threshold represent no risk. However, determining the “true” threshold for humans is difficult, if not impossible, and requires consideration of human variability. Thus, regulatory agencies often determine “safe” levels of exposure for non-cancer endpoints by dividing the highest dose that does not cause any evidence of toxicity upon repeated exposure to a group of laboratory animals (the so-called “No Observable Adverse Effect Level,” or NOAEL) by some “uncertainty” factor.\(^\text{15}\) Usually the factor is 100 or 1000, although the choice of what uncertainty factor to use is dictated by the nature of the toxic response, the quality and quantity of the experimental animal data, and the level of understanding of the mechanism of action of the toxic substance.

Determining the causal relationship between a chemical exposure and a particular chronic disease requires careful consideration of a variety of factors, some of which may be unique to the particular end point in question. For example, establishing an association between a particular drug or chemical and a birth defect requires careful consideration of the exact timing of exposure during pregnancy. Thalidomide, responsible for development of thousands of limb malformation in Europe many decades ago, requires that exposure occur during a very specific period—as short as a few days—early in pregnancy.\(^\text{16}\) Exposure to the drug after the critical period during embryonic development

\(^{15}\) *Id.* at ch. 4.

when the limb buds are forming will not produce that particular birth defect, regardless of dose. Likewise, because the drug is relatively rapidly eliminated from the body, exposure very early in pregnancy—but that was stopped several days prior to the period of limb bud development—also would not produce the birth defect.

Although there is great interest in understanding how environmental factors might contribute to chronic neurological diseases such as Alzheimer’s and Parkinson’s disease, there are relatively few examples where environmental exposures have been shown to contribute to these diseases. Perhaps the most notable example is that of a batch of synthetic heroin that was contaminated with a substance known as MPTP, and subsequently sold on the streets of San Francisco. Numerous young men (under the age of 30) presented with “rapid onset” symptoms essentially identical to Parkinson’s disease. On detailed investigation, it was learned that they had all used a synthetic heroin substance shown later to have contained MPTP. This substance is selectively toxic to certain nerve cells in the brain. These same cells, called “dopaminergic neurons,” are lost progressively with age in all people, resulting in Parkinson’s disease in some (those with a somewhat accelerated loss of cells). Thus, the street drug was able to do in weeks what normally takes a lifetime of “normal” aging. There is now great interest to find other environmental factors that might contribute to the enhanced rate of loss of dopaminergic neurons that seems to be the hallmark of Parkinson’s disease. One environmental chemical, an herbicide called “paraquat,” has a strong structural similarity to the active metabolite of MPTP, and thus there has been substantial toxicological and epidemiological inquiry into whether environmental or occupational exposure to paraquat might contribute to Parkinson’s disease. At this point in time, there is limited toxicological and epidemiological data suggestive of a link between paraquat exposure and Parkinson’s disease, but there remains great controversy and uncertainty over whether paraquat or other pesticides represents a substantial risk factor for Parkinson’s disease.

There is also substantial interest in how chemicals might modify the immune system. There are three ways by which chemical interactions with the immune system could be important. In the first, the chemical may cause direct toxicity to cells of the immune system, thereby interfering with normal immune functions. Numerous chemicals, including “dioxin,” have the ability to interfere with normal immune function, and at sufficient doses, may disrupt immune function. This could lead to enhanced susceptibility to infection, or perhaps even increased risk of cancer, since the immune system plays an important role in destroying precancerous and cancerous cells. Establishing whether a particular chemical has induced immune dysfunction in an individual, however, would require application of the same basic principles of toxicology and epidemiology as for any other type of toxic effect, including “dose-response” and the concept of thresholds.

The second way in which a chemical might interact with the immune system is through the development of an “allergic” reaction to the chemical itself. This is illustrated by the common allergies to penicillin. Some chemicals are capable of triggering the immune system to develop antibodies against the chemical (or, more accurately, to a protein in the body that has been modified by the chemical), and subsequent exposures to that chemical can induce an allergic response. This is a major concern for many drugs, as allergic responses can be fatal. Once “sensitization” has occurred (e.g., the individual has developed antibodies to a specific chemical), relatively small doses of the chemical may be sufficient to trigger a response. Thus, people with allergic sensitization to a specific chemical may respond at a dose much lower than the “average” person, and the response will be qualitatively different (e.g., rather than causing liver damage at a high dose seen in most people, the “allergic” individual may have an asthmatic attack, or develop skin rashes or GI disturbances, at much lower doses). One of the most controversial areas in toxicology and environmental medicine is that related to a number of syndromes such as

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“Multiple Chemical Sensitivity” (MCS), Gulf-War Syndrome, Sick Building Syndrome, Chronic Fatigue Syndrome, etc., for which an immunological basis may be involved. As noted by Kipen and Fiedler:

Symptoms, and especially those without clear underlying medical explanations, account for a large percentage of clinical encounters. Many unexplained symptoms have been organized by patients and practitioners into syndromes such as chronic fatigue syndrome, multiple chemical sensitivity, sick building syndrome, Gulf War syndrome, and the like. All these syndromes are defined solely on the basis of symptoms rather than by medical signs. Some of the above-described conditions overlap strongly with explained conditions such as asthma. The relationship of such symptoms and syndromes to environmental exposure is often sharply debated, as is the distinction between the various syndromes.

Litigation in this area often pits toxicologists, epidemiologists, and/or environmental and occupational medicine specialists against another group of physicians identified as “clinical ecologists.” As noted by Goldstein and Henefin:

Clinical ecologists . . . have offered opinions regarding multiple-chemical hypersensitivity and immune-system responses to chemical exposures. These physicians generally have a background in the field of allergy, not toxicology, and their theoretical approach is derived in part from classic concepts of allergic responses and immunology. This theoretical approach has often led clinical ecologists to find cause-and-effect relationships or

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low-dose effects that are not generally accepted by toxicologists.  

A third way that chemical exposures might involve the immune system involves the development of autoimmune diseases such as lupus, rheumatoid arthritis, and scleroderma. These are important and disabling diseases, yet our understanding of why the immune system sometimes goes awry is limited. Autoimmune diseases arise when the immune system begins to recognize normal tissues as “abnormal” and mounts an attack to destroy the tissue (similar to transplant rejection, where the transplanted organ is recognized as foreign by the immune system). Because the etiology of autoimmune disease is largely unknown and unpredictable, there have been many efforts to identify environmental factors that contribute to the development of autoimmune diseases. Probably the most extensively studied disease in this regard is lupus (Systemic Lupus Erythematosus, SLE). About a half a dozen drugs have been definitively linked with lupus, with dozens more implicated. However, the list of non-drug “environmental” chemicals that have been definitively shown to cause lupus (or other autoimmune diseases) is much shorter. Some inorganic substances, in particular silica, gold, cadmium, and mercury, have been shown to induce autoimmunity in animals and humans. There is suggestive data that exposure to organic solvents, certain chlorinated hydrocarbons such as vinyl chloride, trichloroethylene, and hexachlorobenzene, can also induce autoimmunity, although the scientific evidence (both toxicological and epidemiological) for this is marginal. It remains an area of scientific interest and controversy.

D. Environment and Cancer Risk

Claims of cancer, or increased cancer risk, or fear of cancer,
following chemical exposures are often key elements of toxic tort litigation, and thus I will devote a substantial amount of space to this particular form of toxic response. While it is clear that some chemical pollutants (potentially found in air, food, and/or water) have the ability to cause cancer in either or both experimental animals and humans, deciding whether a particular chemical exposure has “more probably than not” been a “substantial contributing factor”—or whatever the relevant burden of proof might be—in a particular person’s cancer (or risk of cancer, or fear of cancer) is a major challenge for scientists, lawyers, judges, and jurors. To facilitate an understanding of the scientific challenges that are faced in such litigation, it is perhaps useful to look at the “big picture” of what scientists know—and don’t know—about the causes of cancer.

1. Major Causes (Risk Factors) of Cancer

Over the last fifty or so years, a tremendous amount of epidemiological data has been collected on the relationship between a variety of “environmental factors” and the incidence of cancer. Studies comparing cancer risks in different populations with various lifestyle, genetic, cultural, dietary, and behavioral characteristics have led to a reasonable understanding of the major “risk factors” for cancer. These data are of course based on the incidence of cancer in large populations, and thus it is difficult to ascribe “individual” risk to a specific person from these data. Based on such analyses, it has been stated that 85-90% of all cancers are “environmentally-related” and thus potentially preventable. It should again be emphasized, however, that the term “environmentally-related” in this context refers to everything other than genetics (including smoking, diet, lifestyle, etc.) and does not equate directly to “environmental pollution.”

As illustrated in Table 1, approximately 35-40% of all cancer deaths are attributable to tobacco products.\(^23\) While much of this is

l lung cancer (the leading cause of cancer-related deaths in both men and women in most developed parts of the world), smoking also increases risk of oral, bladder, kidney, and several other cancers.

**TABLE 1—“BEST ESTIMATES” OF THE MAJOR RISK FACTORS FOR CANCER**

<table>
<thead>
<tr>
<th>Factors</th>
<th>Best Estimate (%)</th>
<th>Range (%)</th>
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<tr>
<td>Tobacco</td>
<td>35</td>
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<tr>
<td>Diet</td>
<td>35</td>
<td>10 – 70</td>
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<td>Cultural and Lifestyle factors</td>
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<td>1 – 13</td>
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<tr>
<td>Infectious agents</td>
<td>10</td>
<td>5 – 20</td>
</tr>
<tr>
<td>Genetics</td>
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<td>2 – 10</td>
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<td>2 – 8</td>
</tr>
<tr>
<td>Alcohol</td>
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<td>2 – 4</td>
</tr>
<tr>
<td>Geophysical factors (e.g., radon)</td>
<td>3</td>
<td>2 – 4</td>
</tr>
<tr>
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The next most important factor—roughly equal in importance to smoking—is “diet.” What it is about diet that is so important remains uncertain. What is clear is that there are many aspects of the diet that can either increase or decrease cancer risk. For example, diets high in fruits and vegetables have consistently been shown to lower the risk of a variety of cancers.24 In some studies, seminal work of Sir Richard Doll. Id. Recognition of infectious agents as a substantial contributor to several types of cancer, especially for cervical and stomach cancer, became evident in the past decade. Id. The American Cancer Society also discusses the major causes of cancer in their book, entitled: CANCER: WHAT CAUSES IT, WHAT DOESN’T (2003), available for purchase at http://www.cancer.org.

diets high in animal fat have been associated with increased risk of some common cancers (e.g., breast), but the relationship is not always seen, and it remains unclear whether the amount and/or type of fat in the diet are important risk factors. There are also some chemical contaminants in the diet that may increase cancer risk, but again for most populations it is not clear how important natural dietary carcinogens (cancer-causing chemicals) are to overall cancer risk. In some parts of the world, however, a common mold contaminant of corn and peanuts—called “aflatoxin”—is certainly an important contributor to the very high incidence of liver cancer. It has been shown that aflatoxin is much more dangerous in populations where hepatitis B viral infections are common.25

The third most important category of risk factors revolves around cultural and lifestyle factors, which includes sexual practices and reproductive factors. Often these cultural factors interact with other environmental factors, such as viruses. For example, it is now recognized that almost all cervical cancer is due to infection with the human papilloma virus (HPV), which is transmitted through sexual activity. For reasons that are unclear, cervical tissue in teenage women seems more susceptible to HPV infection than that in older women. Thus, sexual activity at a young age is a major risk factor for cervical cancer. While this disease is relatively easily diagnosed (via Pap smear) and treated if detected early, large differences in access to medical care and sex education can make a huge difference in the mortality of this disease across populations.

Breast cancer is the second leading cause of cancer-related deaths among women in the United States and many other developed countries, trailing only smoking-related lung cancer. The major risk factor for breast cancer appears to be a constellation of reproductive factors that influence a woman’s “lifetime dose” of unopposed estrogen. Thus, the age of onset of menstruation, the
age of onset of menopause, the number of children, the age of first pregnancy, and the extent of breast-feeding, all influence breast cancer risk. Thus, it is not surprising that breast cancer incidence and mortality is much lower in countries and cultures where women have their first children early in life, have multiple children, breast feed for extended periods, and often have dietary habits that postpone (or, at least don’t accelerate) the onset of menstruation, compared to a typical “suburban U.S.” lifestyle. Recently, there has been much public press coverage of the discovery of several “breast cancer genes” (BRCA1, BRCA2, BRCA3). Although there is little question that women who carry variant forms of the genes are at substantially increased risk of developing breast cancer (especially at a younger age), the overall contribution of these rather rare genetic causes of breast cancer is probably substantially less than 10% of all breast cancers. Thus, the large majority of breast cancers seem not to have major genetic contributors. But it remains uncertain whether there are important “environmental susceptibility” genes that might interact with environmental factor(s) to substantially increase breast cancer risk.

Of the various identifiable “environmental” factors not associated with diet or lifestyle, infectious agents seem to play a more important role than was expected only a decade ago. It is now clear that well over 90% of cervical cancers are due to HPV infections. Many cases of stomach cancer are directly attributable to helicobacter coli. Most cases

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of liver cancer worldwide can be attributable to hepatitis B (and probably C) viral infections, and alcohol consumption. Even the HIV virus responsible for AIDS is associated with substantially increased risk for certain types of cancer. It is possible, although still not proven, that a significant fraction of human blood-related cancers (leukemias and lymphomas) have a viral etiology, as several leukemia viruses have been identified in animals.

What role do “man-made” chemical pollutants, such as heavy metals, pesticides, industrial solvents, asbestos, etc., play in overall cancer risk? As indicated in Table 1, “occupation” is thought to be responsible for 3-5% of all cancers, although there is reasonable hope and expectation that this will decline substantially as the long history of high-level occupational exposures to cancer-causing substances becomes a relic of the past. But the incidence of asbestos-related lung cancer and mesothelioma, derived from occupational exposures that occurred predominantly in the ’40s, ’50s, ’60s and early ’70s has not yet peaked, since latency period (time from first exposure to the development of clinical disease) may be as long as fifty to sixty years in some individuals. Greatly improved awareness and early identification of potential cancer-causing chemicals, coupled with significant improvements in workplace controls, monitoring, and worker education (at least in developed countries) should result in a drastic reduction in the incidence of occupationally related cancers in the future.

Probably the most uncertain and controversial contributor to cancer risk is that associated with environmental pollution.

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31 See supra note 23 & Tbl. 1.

32 See Julia G. Brody & Riuthann A. Rudel, Environmental Pollutants and Breast Cancer, 111 ENVTL. HEALTH PERSP. 1007, 1007-19 (2003); Shuanfang Li, Stephen D. Hursting, Barbara J. Davis, John A. McLachlan & J. Carl Barrett,
Although it is very difficult to make reasonable “estimates” of the contribution of environmental pollution to overall cancer incidence and mortality, most experts place the number at only a few percent, at most. However, even 1% of 500,000 deaths a year is not an insignificant number (5000) of potentially preventable deaths, so efforts to reduce the use and release of chemical carcinogens are not ill-founded. The challenge comes in balancing the potentially real, but very low, risks of cancer in a large population against the societal benefits that come from the industrial and consumer activities that contribute to the pollution. The basic ways that chemicals can increase cancer risk (chemical carcinogenesis) and the process of “carcinogenic risk assessment” for chemical pollutants are discussed in more detail below.

One example in the area of environmental carcinogenesis that has been the subject of substantial tort and regulatory litigation is that of “dioxins.” Dioxins represent a group of industrial by-products produced inadvertently in the chemical manufacture of trichlorophenol (TCP). TCP was widely used in the synthesis of the herbicide, 2,4,5 trichlorophenoxy acetic acid (2,4,5-T), a component of Agent Orange. TCP was also used in the manufacture of the antibacterial soap ingredient, hexachlorophene, so many antibacterial soaps were also contaminated with trace amounts of dioxins. Although there are more than a dozen specific “dioxin” chemicals, the term is generally used to refer to one
highly toxic form, 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD). When tested in experimental animals, TCDD is extremely toxic, causes cancer and birth defects at extraordinarily low doses, and is generally considered the most toxic and carcinogenic man-made chemical ever studied. Dioxins represent an interesting challenge to the courts for several reasons. First, there are very large species differences in susceptibility to the toxic—and presumably carcinogenic—effects of TCDD. For example, the single lethal dose of dioxin in guinea pigs is approximately 0.1 micrograms per kg of body weight, whereas the lethal dose in hamsters is more than 10,000 times greater. Second, although studies in rats and mice provide consistent evidence that “dioxin” is a potent and effective carcinogen, human epidemiology studies are less convincing. Furthermore, dioxin is not appreciably metabolized in the body, nor does not cause mutations, and the “mechanism” by which it causes cancer is uncertain. Because it is very soluble in fat and is not metabolized in the body, it remains in the body for many years following exposure. Because potentially tens of thousands of military personnel were exposed to dioxin during the Vietnam War, and because of the widespread use of certain herbicides containing small amounts of dioxins in agriculture, forest practices, utility and highway right of ways, and even residential property, it has been the subject of extensive toxic tort litigation. Although it is probably one of the most extensively studied chemical carcinogens, there remains substantial scientific uncertainty as to the actual levels of cancer risk to humans exposed to trace levels of dioxins in the environment.

2. General Mechanisms of Chemical Carcinogenesis

Chemicals that cause an increased incidence of cancer in a population (experimental animals or humans) following exposure are referred to as “carcinogens.” The process of chemical carcinogenesis is “multi-stage,” such that several events must

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34 See sources cited supra note 33.
35 Casarett and Doull’s Toxicology: The Basic Science of Poisons, supra note 11, at Ch. 8.
occur before a normal cell is transformed into a malignant (cancer) cell. Typically, the process of carcinogenesis is divided into four general stages: (1) initiation, (2) promotion, (3) progression and (4) metastasis. A chemical carcinogen may increase the incidence of tumors by acting at any or all of these various stages. One of the most important ways that a chemical may act as a carcinogen is by interacting with DNA in somatic cells to cause mutations. (Somatic cells refer to all cells in the body except sperm and egg cells (ova)). Mutations in somatic cells may lead to permanent changes in the DNA that result in critical changes in the way the cell controls its rate of cell division. Such a permanent alteration in DNA of a somatic cell is referred to as “initiation,” and represents the first stage of chemical carcinogenesis. Because initiation results in a permanent change in the DNA of a cell that is subsequently passed on to all “daughter” cells following division of the mutated cells, initiation is generally considered to be an irreversible process, and initiated cells may accumulate in the body throughout life.

By definition, all chemicals that are “initiators” are mutagenic, and thus short-term tests that demonstrate the mutagenic ability of a chemical make it a suspect carcinogen. However, not all chemicals that test positive in mutagenicity assays are carcinogenic for a variety of reasons. Nevertheless, a chemical that consistently tests positive in numerous different short-term mutagenicity assays is more likely to be carcinogenic than a chemical that routinely tests negative. However, as with all toxicological responses, the “dose-response” for mutagenesis is critically important to consider. Thus, when considering the potential health significance of exposure to chemical mutagens that may act as carcinogens, it is important to keep the total or cumulative “dose” in mind, as the critical issue is whether there is a biologically relevant increase in the “background” rate of DNA damage from all other sources over the lifetime of an individual.

Although initiation is an essential first step toward cancer, most initiated cells do not go on to become cancers because they usually require additional genetic changes and other external stimuli to

\[36\] Id.
become true cancer cells. An initiated cell is analogous to a car whose accelerator is stuck part way on. As long as the brakes work, the speed and control of the car can be adequately maintained. However, if a second change occurs, resulting in the loss of the brakes, then the car can no longer be controlled. It should be recognized that the vast majority of mutations in a cell do not have any effect on cell growth and regulation, just as most mechanical malfunctions in a car do not result in a stuck accelerator or loss of brakes.

The second step of carcinogenesis, referred to as “promotion,” occurs when some external stimuli, including exposure to certain chemicals, increases the rate of cell proliferation of initiated cells or otherwise enhances the ability of an initiated cell to become cancerous, but does not directly interact with DNA. A chemical that increases cancer risk by acting as a promoter of carcinogenesis is generally considered to be less of a concern because the promoting stimulus goes away when the exposure stops—e.g., promotion is “reversible.” The process of promotion can be viewed as a relatively early stage in carcinogenesis where an initiated cell is stimulated to divide repeatedly to give rise to a small colony of initiated cells (a “preneoplastic” lesion).

The third stage of carcinogenesis, referred to as progression, represents the long period of time where the small colony of initiated cells acquires additional mutations that further transform the cell from a normal cell to a cancer cell. To return to the car analogy, additional mutations damage the cell’s “brakes,” “steering,” or other critical functions necessary to properly maintain control.

The probability of a single cell acquiring all of the necessary genetic changes to convert it from a normal cell to a cancer cell depends on a variety of factors, including the dose and duration of exposure to mutagenic substances. Exposure to man-made mutagenic chemicals can increase cancer risk. However, it should be recognized that the vast majority of DNA damage that occurs in our cells results from normal metabolic processes and exposure to natural components in the diet or to UV radiation in sunlight. As a cell “burns” sugars to produce energy it generates reactive by-products of oxygen. These by-products, called “free radicals” or
“reactive oxygen species” (ROS) can and do directly damage DNA. In addition, common chemicals found naturally in the diet and/or formed during cooking can also damage DNA (e.g., act as mutagens). Fortunately, the cells of the body have remarkable processes that can reduce the damage to DNA from ROS and mutagenic chemicals (both natural and man-made), as well as repair damaged DNA. Many vitamins and certain chemicals found naturally in the diet (especially in fruits and vegetables) act as “antioxidants” and can help protect cells from the DNA damaging effects of ROS and chemicals (both man-made and natural) that damage DNA. This is one reason why diet is so important in lifetime cancer risk. Many studies have demonstrated that diets high in fruits and vegetables lower the risk of many types of cancer.37

Because the probability of a single somatic cell acquiring all of the necessary genetic changes (mutations) to become a cancer cell is quite small and is a function of the period of time that a cell has to acquire such mutations, cancers that occur because of exposure to a carcinogen are both relatively rare in an exposed population and are usually not seen until many years after the initial period of exposure. For cancers caused by prolonged or repeated exposure to a chemical, the time frame from first exposure to when the disease becomes clinically evident is referred to as the “latency” period. In general, the latency period is somewhat inversely related to the extent of exposure (dose). For most human cancers that are related to chemical carcinogen exposure (e.g., cancers related to cigarette smoking), the latency period is usually twenty to forty years. Certain cancers (mesotheliomas) that arise from occupational exposure to asbestos typically are not seen for thirty or more years after first exposure.38 The shortest latency period possible appears to be at least a couple of years following very high levels of exposure to mutagenic chemicals used to treat cancers, especially leukemias.39 Because latency seems to be inversely related to dose,

37 See sources cited supra note 24.
39 Richard A. Larson, Michelle M. LeBeau, James W. Vardiman & Janet D.
very low levels of exposure to mutagenic chemicals may be of little practical consequence to an individual because (1) the extent of DNA damage is very small, relative to the “background” rate that occurs from all other sources, and (2) the latency period for developing a cancer is very long and may exceed normal life expectancy. Although these are very important practical biological considerations, they are often not considered in quantitative risk assessment of low dose carcinogen exposures by regulatory agencies, who usually assume that risk is directly proportional to dose at low doses (the “linearized dose-response” approach to cancer risk assessment).

Another factor that affects the occurrence of a cancerous cell is the rate of cell turnover, or “cell proliferation.”40 Because all cells have a background incidence of spontaneous mutations, the likelihood of a cell becoming mutated is related to the rate of cell replication, analogous to rolling dice. The more often the dice are rolled the more likely a specific number is to appear. This factor is especially important in considering the use of high doses of chemicals in laboratory animal test for cancer-causing potential. When doses of the test chemical are so high that they cause tissue damage (and thus stimulate cell division to repair the damage)—which usually would not occur at low doses—direct extrapolation of the rate of tumor formation in the animals given high doses to humans exposed to much lower doses that don’t cause tissue damage is of questionable scientific value.41 The particular rodent strains used also often have a high background rate of spontaneous tumors.42 Thus, any chemical that damages cells and causes considerable regeneration (i.e., cell proliferation) may increase the

Rowley, Myeloid Leukemia After Hematotoxins, 104 (Suppl 6) ENVTL. HEALTH PERSP. 1303, 1303-07 (1996).


42 See Gold, supra note 41; see also Zito, supra note 13.
likelihood of a cancerous cell occurring.

E. Use of Toxicological Data to Assess Chemical Risks in Populations and Individuals

It is important to recognize that the procedures commonly used in “risk assessment” for the purposes of establishing public health guidelines that represent “acceptable” exposure levels for large populations are often, in this author’s opinion, of marginal relevance to estimating “causation” in an individual—e.g., whether a particular chemical caused or contributed to a particular disease or illness in a given person. Although toxicological data—and the basic principles of toxicology outlined above—are useful for both (establishing guidelines for protection of public health and establishing “causation”), there are substantial differences in approach. Thus, use of toxicological data for these two distinct purposes will be discussed separately in the following sections.

F. Use of Toxicological Data to Establish “Acceptable” Levels of Exposure for Large Populations (Public Health)

Much of the available dose-response criteria for assessing chemical toxicity and risks to human health are based on protective guidelines developed by federal (e.g., EPA, ATSDR) and sometimes state agencies. The federal government and national organizations using similar approaches also set occupational health guidelines and standards for protection of workers. Guidelines for protection of the general public are usually more stringent than for workers, who are assumed to be part of a healthier and less sensitive population. Public health guidelines, however, should not be interpreted as predicting exact levels at which effects would occur in a given individual. Because a number of protective, often “worst-case” assumptions (e.g., exposure to any dose of a carcinogenic chemical based on animal studies confers a risk of cancer in humans, high daily exposure for a lifetime) are made in estimating allowable exposures for large populations, these criteria and the resulting regulatory levels (e.g., MCLGs, MCLs) generally
overestimate potential toxicity levels for nearly all individuals. Furthermore, because these guidelines are intended to be protective of all individuals in a population, including the very young, the very old, and other potentially “sensitive” individuals, the theoretical risks from exposure at the guideline level is likely to be substantially overestimated for the large majority of individuals in the population. Nevertheless, they can provide useful guidance to public health agencies that have the responsibility of protecting all individuals in large populations.

Public Health criteria developed by the EPA for individual chemicals usually include determination of non-cancer reference doses and “cancer potency” or “slope factors.” Non-cancer reference doses represent the dose below which no adverse health effects are expected, even in sensitive individuals exposed repeatedly at the defined level for many years. Reference doses are usually derived from “No Observable Adverse Effect Levels” (NOAELs) or “Lowest Observable Adverse Effect Levels” (LOAELs) in the toxicological literature. NOAELs and LOAELs are usually determined from experimental animal studies, rather than human exposures. The term “Reference Dose” is frequently used to refer to a dose of a chemical to humans that could be consumed on a daily basis for a lifetime with no chance of anyone exhibiting an adverse response (the specific definition of such “safe” doses varies from agency to agency and regulation to regulation). Reference Doses are obtained by dividing the “NOAEL” dose determined in animal studies by an Uncertainty Factor. Uncertainty Factors usually range from 100 to 1000, depending on the amount of uncertainty in, for example, extrapolating from animals to humans, short-term to long-term effects, average to sensitive members of the population. Generally, the more uncertainty factors required, the more likely it is that the Reference Dose will be lower than what would actually be necessary for protection of humans because each uncertainty factor errs on the side of overprotection. Thus, although health authorities can confidently expect that exposures below reference dose levels will not result in adverse effects, the converse is not

43 See sources cited supra notes 13 & 14.
true. Exposures in a given individual that exceed a reference dose level do not signify that effects are likely to occur because of the margin of “safety” built into these Reference Doses which are intended to provide guidance for protecting even sensitive members of the population. Thus, such regulatory levels are of substantial value to public health agencies charged with ensuring the protection of the public health, but are of limited value in judging whether a particular exposure was a substantial contributing factor to a particular individual’s disease or illness.

G. Determining Regulatory Guidelines for Chemical Carcinogens for Protection of Public Health

For carcinogens, most regulatory agencies have used “default” assumptions about the dose-response relationship such that it is assumed that the risk of developing cancer is proportionate to dose at all doses (e.g., there is no “threshold” dose). Thus, to establish socially acceptable levels of exposure to carcinogens commonly found in food, air and water, the EPA, FDA, and other regulatory agencies have established guidelines for conducting risk assessments. Using such procedures, “acceptable,” “tolerable,” “permissible,” or “safe” levels of exposure to a specific chemical are often established based on regulatory policy decisions on allowable risk, or tradeoffs between risk reduction and cost. For contaminants in drinking water, such levels are referred to as “Maximum Contaminant Levels,” or MCLs. The EPA has a long-standing policy that dictates that the desired level of cancer risk for a contaminant in drinking water is zero. Thus, for carcinogens the EPA has established MCLGs (Maximum Contaminant Level Goals) of zero. However, because zero levels are generally not
achievable by modern technology, the actual drinking water standards, MCLs, are usually based on other considerations such as technical feasibility, cost-benefit analysis, and background levels. However, even if the standard is based primarily on a technology or cost, MCLs for most such chemicals in drinking water must still be within an acceptable range of health risk. For cancer risk from chemicals in drinking water, the EPA has stated this range to be an excess lifetime risk of cancer over background of one in 1,000,000 to one in 10,000. However, there are some exceptions where MCLs have been established that yield theoretical excess lifetime cancer risks much greater than one in 10,000. The recent adoption of an MCL for arsenic in drinking water of 10 ppb is such an example.48

Because the lifetime probability of dying from cancer for someone living in the United States is about 1 in 4 (25%, or 0.25 lifetime probability), a theoretical increase in lifetime cancer risk (mortality) of 1 in 100,000 would provide a potential increase in overall lifetime probability of dying from cancer from approximately 0.25 to approximately 0.25001. Thus, when citizens are confronted with evidence that their drinking water is contaminated with a “cancer causing chemical” at levels that exceed federal regulatory limits, it becomes important to ensure that the public understands how such standards are derived and the significance of the potential increase in risk, relative to other common risks encountered daily.

EPA cancer “slope factors” represent the slope of the dose-response relationship statistically extrapolated from studies of high dose exposure and cancer in laboratory animals or human populations. The EPA default assumption in these slope factors is that no dose of a carcinogenic chemical is without some risk of cancer and that one can extrapolate high dose exposures and the risk of cancer to low doses. The use of a slope factor in this

manner ignores the ample evidence that for many carcinogenic chemicals, practical thresholds may exist for significant cancer risk because of detoxification mechanisms at low doses (e.g., difference in metabolism of the chemicals at high dose versus lower doses) or because of the mechanisms of action. Therefore, these slope factors cannot be expected to accurately predict a risk of cancer, if any, in a given individual at low doses. Although they may be somewhat useful to make crude estimates of individual risk, many assumptions go into the determination of the cancer slope factor, and it is important to consider the relevance of the particular animal study from which the slope factor was determined when attempting to use such values in individual risk calculations.

\[H. \text{ Use of Toxicological Data in Assessment of Individual Causation}\]

When assessing whether a particular potentially toxic substance is a substantial contributing factor to an individual's disease or illness, the “regulatory approach” is of little value, although much of the same toxicological and epidemiological data may be used in evaluating causation. The key scientific criteria used to establish causation between an alleged chemical exposure and a particular disease or illness includes the following basic concepts:

1. The toxic substance in question must have been demonstrated to cause the type of illness or disease in question. This addresses the issue of general causation as well as specific causation, and may be demonstrated either in humans following known exposures (usually from accidents, occupational exposures, or intentional exposures), or, in the absence of human data, in experimental animals intentionally exposed to the agent in question. Because most chemicals that are widely encountered in the environment, such as pesticides, metals, and industrial solvents, are manufactured, workplace exposure to humans may occur. Occupational health and safety regulations require workplace monitoring, and thus there is frequently a substantial amount of toxicologically relevant data from workplace monitoring that can
be used to assess whether a particular chemical is capable of causing a particular disease or illness. Indeed, virtually all synthetic chemicals identified by EPA or International Agency for Research on Cancer as “known human carcinogens” have been identified as such through studies of workers exposed to the chemical in the workplace. Workplace exposures are typically hundreds to thousands of times greater than incidental environmental exposures that might occur from contamination of drinking water, or off-site migration of chemicals via the air.

2. The individual must have been exposed to a sufficient amount of the substance in question to elicit the health effect in question. As noted above, the main tenant of toxicology is the “dose-response” relationship. If criterion (1) above has been established for a given chemical, then it must be established that the individual’s dose over a defined period of time was sufficient to cause the alleged health effect. It is not adequate to simply establish that “some” exposure occurred. Because most chemically induced adverse health effects clearly demonstrate “thresholds,” there must be reasonable evidence that the exposure was of sufficient magnitude to exceed the threshold before a likelihood of “causation” can be inferred. For carcinogenic chemicals that act via mutagenic action, a threshold may not be evident. Thus, although any level of exposure will theoretically increase the probability of developing the disease, the risk follows a dose-response relationship, and the dose must be sufficient to “significantly” elevate the risk above the background. What represents a “significant” increase in cancer risk is of course subjective and influenced by many factors. However, as noted above, because the process of chemical carcinogenesis is always associated with a “latency,” and the latency period is generally inversely related to dose, at very low doses of even “direct acting,” mutagenic carcinogens, the latency period might exceed life expectancy, thereby imparting a “practical” threshold.

3. The chronological relationship between exposure and effect must be biologically plausible. If a disease or illness in an individual preceded the established period of exposure, then it cannot be concluded that the chemical caused the disease, although it may be possible to establish that the chemical aggravated a pre-
existing condition or disease. For cancer cases, diagnosis of the cancer in a time frame close to the beginning period of exposure (i.e., within a few years) argues strongly against a causal relationship, since, as noted above, chemically-induced cancers have latency periods that are nearly always in excess of five years, and are somewhat inversely related to dose.

4. The likelihood that the chemical caused the disease or illness in an individual should be considered in the context of other known causes. Although this consideration may not be essential to establish general causation, it is a critical consideration in the quantitative assessment of whether the substance was “more likely than not” a cause or substantial contributing factor to the disease or illness in a specific person. This is especially important in cancer causation, because cancer is by its very nature a multi-factorial disease. As discussed above, chemicals that are mutagenic have the theoretical potential to increase cancer risk even at very low doses, although there is a point at which the theoretical risk is trivial, relative to all other causes, known and unknown. As there are literally hundreds, if not thousands, of mutagenic naturally-occurring chemicals present at low levels in our diet and thus also present theoretical cancer risks, it becomes important to put such theoretical, “low dose” risks in perspective.49

J. Multiple Exposures/Mixtures

Another area of relevance to human risk assessment for environmental pollutants is the fact that, unlike experimental animals, humans may be exposed to multiple different chemicals, diets, and lifestyle factors that affect the dose-response relationship for a given chemical. For chemical carcinogens, it is often assumed that the risks are additive even though they may not act through similar mechanistic pathways. That is, the risk for each chemical in a mixture is calculated separately, and the total risk from exposure to the mixture is simply the sum of the risks for each individual chemical. While there are examples of non-additive responses (both “synergistic,” where the response of two chemicals is greater

49 See Kipen & Fiedler, supra note 20.
than predicted from adding the individual response alone, and “antagonistic,” where one chemical appears to reduce the risk of the second), unless there is compelling evidence to the contrary, additivity of risks for all carcinogens is generally assumed. However, for carcinogens that act via different modes (e.g., genotoxic vs. non-genotoxic carcinogens), additivity is less certain, and mechanistic data may warrant consideration of non-additive models for interaction.

EPA risk assessment guidelines also consider non-cancer effects of chemicals to be additive, particularly if they effect the same endpoint at lower doses. If the chemicals act by the same mechanism, then their action could be additive even when exposure to each is below a dose that would cause effects.

III. FUTURE SCIENTIFIC OPPORTUNITIES AND CHALLENGES IN TOXIC TORT LITIGATION

The past decade has seen a tremendous advance in DNA-based technologies that offer exciting challenges and opportunities to the field of toxicology. The growing area of “toxicogenomics”—the application of new molecular technologies to understand how chemicals cause adverse responses in cells, tissues, and organisms—will eventually play an important role in toxic tort litigation. Rather than examining the effect of a chemical on one or a few biochemical pathways, the tools of toxicogenomics provide a means to examine the global response of a cell to a chemical stimulus, resulting potentially in a “fingerprint” alteration in expression of thousands of different genes (transcriptomics), proteins (proteomics), or cellular metabolites (metabonomics). The potential exists for such tools to provide convincing proof that a particular disease was related to a specific chemical exposure, through unique changes that potentially can be measured years after the exposure occurred. As noted by Marchant, however, “many obstacles and uncertainties remain to be resolved before toxicogenomics data should be used outside the research context for practical, regulatory or legal applications.”50

50 See Marchant, supra note 6; see also John C. Childs, Toxicogenomics:
these new scientific approaches to linking specific diseases or illnesses to specific exposures can be proven reliable, judges, lawyers, and jurors must rely upon the basic scientific principles of toxicology and epidemiology to establish causation in toxic torts.