A Study on the Current State of Contaminants of Concern Research, With a Focus on Biosolids and Regulations

Maren Fulton
Portland State University

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A Study on the Current State of Contaminants of Emerging Concern Research, With a Focus on Biosolids and Regulations

by

Maren Mariah Fulton

A research project report submitted in partial fulfillment of the requirement for the degree of

MASTER OF SCIENCE
IN
CIVIL AND ENVIRONMENTAL ENGINEERING

Project Advisor:
Dr. Gwynn Johnson

Portland State University
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ACKNOWLEDGEMENTS

I would like to thank my advisor, Dr. Gwynn Johnson of the Department of Civil and Environmental Engineering at Portland State University. Her door was always open, even after my extended sabbatical in my career prior to returning to Portland State University to complete my degree. Her enthusiasm for research and advising helped steer my research and guide my writing when I needed direction, while consistently allowing the freedom for this paper to be my own work.

Additionally, I would like to thank the Columbia River Toxics Reduction Work Group for allowing me to attend several of the meetings. I would also like to thank the following people associated with the Group for their advice regarding the relevant research topic of CECs in biosolids in the Columbia River Basin: Jennifer Morace, Hydrologist at USGS, Northwest Region; Elena Nilsen, Research Chemist at USGS, Northwest Region; and Mary Lou Soscia, Columbia River Coordinator, EPA Region 10.

Thanks is also due to my employers at Ecology and Environment, Inc., for their support and flexibility that allowed me to complete my research while employed full-time.

I would like to thank my sister, Stephanie Fulton, PhD student at the Department of Crop and Soil Sciences, University of Georgia, Athens, for providing valuable feedback and criticism on this paper, as well as her tireless support. I would also like to acknowledge Jack Young, PhD student at the Department of Mechanical and Aerospace Engineering, University of California, Los Angeles, as the second reader of this paper, and I am gratefully indebted to him for his very valuable comments on this paper.

Finally, I would like to express my gratitude to my partner for providing unwavering support and continual encouragement throughout my years of study and researching and writing to complete this project. I also would like to thank my parents, who have always provided endless support and encouragement. This accomplishment would not have been possible without them. Thank you.

Author,
Maren Fulton
Abstract

A Study on the Current State of Contaminants of Emerging Concern Research, With a Focus on Biosolids and Regulations

Maren Mariah Fulton

Chair of the Supervisory Committee:

Maseeh College of Engineering
Associate Professor Dr. Gwynn Johnson

Wastewater treatment plants are a major source of contaminants of emerging concern, as these facilities are the main receptors of these products via household, commercial, and industrial drains, and transport via stormwater runoff. Biosolids are composed of numerous constituents, with a number of environmentally persistent and potentially toxic contaminants of emerging concern identified as present in large concentrations. Research is needed to understand the transport and fate mechanisms of these compounds in biosolids. Additionally, this research is needed to determine a new priority framework to regulate CECs, both on the national level and regional levels. Wastewater treatment plant influent and effluent treatment regulations and practices may be improved upon as more knowledge accrues regarding contaminants of emerging concern behavior in the environment.

The United States environmental regulatory process is a constant work in progress. Inherited from decades-old public health traditions, environmental regulatory programs arose to address the issues of public health as water and air quality issues surfaced. The Toxic Substances Control Act (TSCA) (15 U.S.C. 2601), enacted in 1976, provided the EPA regulatory authority to protect the environment and consumers’ health against risks posed by chemicals in commerce. TSCA only retroactively addressed chemical hazards, after they had been deemed unsafe and previously unrestricted in the public. A bill to overhaul TSCA, the Frank R. Launtenberg Chemical Safety for the 21st Century Act
(House Amendment to the Senate Amendment to H.R. 2576, TSCA Modernization Act of 2015), was passed by the Senate on June 7, 2016 and sent on for President Obama’s signature. The amended law will give the EPA new authority to evaluate the safety of a new chemical before it enters the marketplace. It will also allow EPA to evaluate the safety of chemicals already known to be risks, including chemicals found to persist in the human body and in the environment.

The Columbia River is presented as a case study as an example of a significant waterbody with research and regulatory gaps concerning CECs. The Columbia River waterbody was chosen as it represents an important natural resource for which data gaps exist on CEC sources and pathways into the waterbody, and which also does not receive adequate protective regulations under the national regulatory framework.
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<tr>
<td>ATSDR</td>
<td>Agency for Toxic Substances and Disease Registry</td>
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<tr>
<td>BPA</td>
<td>bisphenol A</td>
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<tr>
<td>CEC</td>
<td>contaminants of emerging concern</td>
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<tr>
<td>CFR</td>
<td>Code of Federal Regulations</td>
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<tr>
<td>cfs</td>
<td>cubic feet per second</td>
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<tr>
<td>CERCLA</td>
<td>Comprehensive Environmental Response, Compensation and Liability Act</td>
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<tr>
<td>CRTRWG</td>
<td>Columbia River Toxics Reduction Work Group</td>
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<td>CWA</td>
<td>Clean Water Act</td>
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<tr>
<td>DBP</td>
<td>drinking water disinfection byproducts</td>
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<td>DDT</td>
<td>dichlorodiphenyltrichloroethane</td>
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<tr>
<td>ECHA</td>
<td>European Chemicals Agency</td>
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<tr>
<td>EPA</td>
<td>United States Environmental Protection Agency</td>
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<tr>
<td>EU</td>
<td>European Union</td>
</tr>
<tr>
<td>FIFRA</td>
<td>Federal Insecticide, Fungicide, and Rodenticide Act of 1972</td>
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<tr>
<td>FWPCA</td>
<td>Federal Water Pollution Control Act of 1948</td>
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<tr>
<td>FWPCA</td>
<td>Federal Water Pollution Control Act of 1972</td>
</tr>
<tr>
<td>FWQA</td>
<td>Federal Water Quality Administration</td>
</tr>
<tr>
<td>ITC</td>
<td>Interagency Testing Committee</td>
</tr>
<tr>
<td>kg/yr</td>
<td>kilogram per year</td>
</tr>
<tr>
<td>LCREP</td>
<td>Lower Columbia River Estuary Program</td>
</tr>
<tr>
<td>MS4</td>
<td>municipal separate storm and sewer systems</td>
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<tr>
<td>NAS</td>
<td>National Academy of Science</td>
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<tr>
<td>NEP</td>
<td>National Estuary Program</td>
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<tr>
<td>NEPA</td>
<td>National Environmental Policy Act</td>
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<tr>
<td>NOAA</td>
<td>National Oceanic and Atmospheric Administration</td>
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<td>NOM</td>
<td>natural organic matter</td>
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<tr>
<td>NP</td>
<td>nanoparticles</td>
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<tr>
<td>NPDES</td>
<td>National Pollutant Discharge Elimination System</td>
</tr>
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<td>NPL</td>
<td>National Priority List</td>
</tr>
<tr>
<td>NRC</td>
<td>National Research Council</td>
</tr>
<tr>
<td>NSSS</td>
<td>National Sewage Sludge Survey</td>
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<tr>
<td>OPA</td>
<td>Oil Pollution Act of 1990</td>
</tr>
<tr>
<td>OTs</td>
<td>organotins</td>
</tr>
<tr>
<td>PAHs</td>
<td>polycyclic aromatic hydrocarbons</td>
</tr>
<tr>
<td>PBDEs</td>
<td>polybrominated diphenyl ethers</td>
</tr>
<tr>
<td>PCA</td>
<td>polychlorinated alkanes</td>
</tr>
<tr>
<td>PCB</td>
<td>polychlorinated biphenyl</td>
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<tr>
<td>PCDD</td>
<td>polychlorinated dibenzo-p-dioxins</td>
</tr>
<tr>
<td>PCDFs</td>
<td>polychlorinated dibenzo-furans</td>
</tr>
<tr>
<td>PCNs</td>
<td>polychlorinated napthalenes</td>
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<tr>
<td>PFC</td>
<td>perfluorinated compounds</td>
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<tr>
<td>PFOA</td>
<td>perfluorooctanoic acid</td>
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<tr>
<td>PFOS</td>
<td>perfluorooctanesulfonic acid</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>PHS</td>
<td>United States Public Health Service</td>
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<tr>
<td>P.L.</td>
<td>Public Law</td>
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<tr>
<td>POP</td>
<td>Persistent Organic Pollutants</td>
</tr>
<tr>
<td>POTW</td>
<td>publicly owned treatment works</td>
</tr>
<tr>
<td>QAC</td>
<td>quaternary ammonium compounds</td>
</tr>
<tr>
<td>RCRA</td>
<td>Resource Conservation and Recovery Act</td>
</tr>
<tr>
<td>REACH</td>
<td>Registration, Evaluation, Authorisation and Restriction of Chemicals</td>
</tr>
<tr>
<td>TCA</td>
<td>1,1,1-trichloroethane</td>
</tr>
<tr>
<td>TCC</td>
<td>triclocarban</td>
</tr>
<tr>
<td>TCS</td>
<td>triclosan</td>
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<tr>
<td>TSCA</td>
<td>Toxic Substances Control Act</td>
</tr>
<tr>
<td>TMDL</td>
<td>Total Maximum Daily Load</td>
</tr>
<tr>
<td>USCG</td>
<td>United States Coast Guard</td>
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<tr>
<td>USDA</td>
<td>United States Department of Agriculture</td>
</tr>
<tr>
<td>USGS</td>
<td>United States Geological Survey</td>
</tr>
<tr>
<td>USACE</td>
<td>United States Army Corps of Engineers</td>
</tr>
<tr>
<td>UV</td>
<td>ultraviolet</td>
</tr>
<tr>
<td>WDOE</td>
<td>Washington State Department of Ecology</td>
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<tr>
<td>WQA</td>
<td>Water Quality Act</td>
</tr>
<tr>
<td>WRDA</td>
<td>Water Resources Development Act</td>
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<tr>
<td>WWTP</td>
<td>wastewater treatment plant</td>
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Chapter 1

1. Introduction

Over the past few decades, research has documented with increasing frequency the effects and occurrence of a class of environmental pollutants now commonly called contaminants of emerging concern (CECs). CECs include all classes of pollutants for which the environmental risks are previously unknown, unrecognized, or unsuspected. A contaminant of ‘emerging concern’ refers to a compound that is not currently regulated (e.g., not designated as a Priority Pollutant in the United States (U.S.)), is commonly detected at low levels in the environment, and may present a threat to environmental and human health.

The designation, ‘emerging concern,’ represents a shift in perspective of what is traditionally considered to be an environmental contaminant. Despite being found at low concentrations, due to their ubiquitous nature, CECs are now being more widely detected in far-ranging and diverse locations in the environment, from surface and groundwaters (Vulliet, 2011) to household dust (Ali, 2011), and from plants (Divan Jr., 2009) and animals (Sun, 2012) to arctic ice (De Laender, 2011; Hermanson, 2010). Most of these chemicals have not been adequately examined for toxicological or environmental impacts, and screening methods and regulatory restrictions are still under development (EPA, 2016). Research into the occurrence and risk of exposure to these compounds has been rapidly expanding, and legislation to regulate sources and discharges of CECs to the environment has been evolving to keep up with these scientific findings.

Many CECs are industrially produced, yet are dispersed to the environment often via common domestic, commercial, and industrial uses. CECs include many pharmaceuticals, personal care products (such as shampoos, deodorants, and fragrances), commercial and household chemicals, some pesticides, nanomaterials, and hormones. It has been estimated that over 100,000 chemicals are currently in commerce, with up to 1,000 new compounds added to current-use each year. Of these thousands of chemicals,
more than 40,000 organic chemicals have been identified as CECs, and that estimate fails to include or consider the associated degradation by-products of those chemicals in environmental systems (Diamond, 2011). Many of these chemicals make their way to wastewater treatment plants (WWTPs), where they may be released to the environment via treated effluent and land application of treated sewage sludge (biosolids).

The land application of biosolids is considered to be a relatively positive reuse option of WWTP sludge, both nationally and internationally. Researchers have shown that biosolids contribute nutrients and beneficial soil amendments such as nitrogen and phosphorus (National Resource Council (NRC) 2002). A significant volume of research has been conducted on the presence of persistent organic pollutants (POPs) in biosolids such as polychlorinated dibenzo-p-dioxins/dibenzo-furans (PCDD/PCDFs), polychlorinated biphenyls (PCBs) and polycyclic aromatic hydrocarbons (PAHs) (e.g., Wild et al., 1991; Alcock et al., 1996; Stevens et al., 2001), and concentrations of these ‘traditional’ POPs in biosolids have declined due to effective source control (Wild et al., 1990; Clarke et al., 2008, 2010). However, analytical results have indicated the presence of large concentrations of a number of CECs, identified as environmentally persistent and potentially toxic, as being present in large concentrations in biosolids (NRC 2002; Davis et al., 2012; LaGuardia, 2004). Research on the transport, fate, and potential ecological effects of CECs once biosolids are applied to agricultural fields, garden plots, and landscaped plants and shrubs is still emerging (Clarke, 2011).

Finally, the Columbia River Basin is presented in this paper as a study of a significant waterbody with threatened water quality. The Columbia River discharges an average volume of 265,000 cubic feet per second (cfs), and is the fourth largest river by volume in the U.S. following the Mississippi, the St Lawrence, and the Ohio Rivers, respectively. The Columbia River waterbody was chosen as it represents an important natural resource for which data gaps exist on CEC sources and pathways into the waterbody, and which also does not receive adequate protective regulations under the national regulatory framework.
1.1 Objectives and Scope

Research regarding the transport and fate, and potential ecological effects of CECs sourced from land-applied biosolids is ongoing in both the U.S. and internationally. The purpose of this paper is to present an overview of the current state of CEC research, with a focus on CECs in biosolids. This paper also aims to present a review on the current state of legislation governing regulations of monitoring requirements and acceptable use of land-applied biosolids. The literature review included over 25 articles, spanning from 1990 to 2016, and numerous reports and documents that referenced CEC research, from which a subset was selected based on relevance of studies of CECs and water quality, and biased towards recent and current research. A case study on the Columbia River watershed is also presented to provide a topical perspective on the relevance of regulatory and CEC issues.

1.2 Problem Statement and Relevance

Approximately 5.6 million dry tons of biosolids are used or disposed of annually in the United States, of which approximately 60% are land-applied as soil amendments. EPA estimates that biosolids are applied to approximately 0.1% of available agricultural land in the United States on an annual basis (NRC, 2002). Biosolids are composed of numerous constituents, with a number of environmentally persistent and potentially toxic CECs identified as present in large concentrations (NRC, 2002; Davis et al., 2012; LaGuardia, 2004), which may be released to the environment via land application.

The presence of priority pollutants such as heavy metals and polychlorinated biphenyls (PCBs) in land-applied biosolids has been extensively studied, including the effects on human health and the environment, and environmental behavior (NRC, 2002). However, data gaps exist in information regarding risks to human and environmental health, occurrence, persistence, transport and fate, and the transformation of emerging contaminants (Chase et al., 2012; Calderon-Preciado, 2011a). Consumption of
unregulated chemicals is constantly increasing, along with increasing detection rates of significant concentrations of these chemicals in biosolids. Research is needed to understand the transport and fate mechanisms of these unregulated compounds in biosolids. Additionally, this research is needed to determine a new priority framework to regulate CECs, both on the national level and regional levels, including the Columbia River Basin. Advancement in the body of CEC research can also provide guidance on improving regulatory oversight, such as implementing appropriate risk evaluation procedures of new chemicals prior to commercial release, and potentially decrease the frequency of one banned chemical being replaced by a new and unevaluated alternative chemical, such as flame-resistant polybrominated diphenyl ethers (PBDEs) and their replacement congeners (Davis et al., 2012). Understanding the concentrations and environmental behavior of CECs in land-applied biosolids is key to determining the significance and implications of these emerging pollutants on human health and the environment.
Chapter 2

2. Introduction to CECs and Relevant Regulations

This section provides an introduction to CECs and environmental concerns regarding CECs, as well as a brief history of relevant environmental regulations. The summary of environmental regulations includes a description of Federal regulatory agencies and regulations, as well as international environmental regulations.

2.1 Introduction to CECs

CECs include all classes of pollutants for which the environmental risks are previously unknown, unrecognized, or unsuspected. CECs are commonly dispersed to the environment via domestic, commercial, and industrial uses. Sources of CECs include agriculture, landfills, residential household drains and WWTPs, and pesticide use in landscaping applications. CECs include many pharmaceuticals, personal care products, commercial and household chemicals, some pesticides, nanomaterials, and hormones. Classes of CECs identified as problem or priority chemicals include perfluorinated chemicals (e.g., PFOS, PFOA); polychlorinated alkanes (PCAs); polychlorinated naphthalenes (PCNs); organotins (OTs); unregulated congeners of polybrominated diphenyl ethers (PBDEs); triclosan (TCS); triclocarban (TCC); benzothiazoles; antibiotics and pharmaceuticals; synthetic musks; bisphenol A (BPA); quaternary ammonium compounds (QACs), steroids; personal care products; unregulated pesticides; and a wide range of industrial chemicals and nanomaterials.

One of the primary reasons for the increase in awareness of CECs in the environment has been due to recent improvements in the sensitivity of analytical techniques, allowing the detection of previously undetectable low-concentration contaminants. Studies on the risks posed by these pollutants to human and environmental health and safety, and research on the sources, transport, fate, and behavior of CECs are now at the forefront of environmental research. The topic has even gained exposure in media with published
articles concerning these compounds. One reason that CECs attract popular attention is because the major source of these contaminants is from the general population, via everyday use of products that contain the very compounds that pose environmental risks.

The Athens, Georgia, EPA National Exposure Research Laboratory conducted a 2010-2011 biennial review to identify particular CECs as major trends in research or as new emerging contaminants (Richardson, 2011a). The EPA acknowledged that many new environmental CECs were identified in this biennial review due to improved analytical techniques that have allowed detection levels at previously undetectable limits in the low ng/L concentrations. This is significant, as prior analytical techniques did not allow for such low detection levels, resulting in many chemicals occurring undetected in the environment. The CECs identified in this biennial review are summarized below.

Perfluorinated compounds (PFCs) were identified as a major trend of research in the biennial review conducted by the Georgia EPA research laboratory. PFCs have been in production for more than 50 years, and are used in the production of stain repellents, paints, adhesives, waxes, polishes, metals, electronics, fire-fighting foams, caulks, and food packaging such as microwave popcorn bags and French fry boxes. PFCs are persistent and environmentally mobile, due to their unique chemistry of being composed of one of the strongest chemical bonds of carbon-flourine, and their hydrophobic and lipophilic behavior. Between 2000 and 2002, an estimated 5 million kg/yr of PFCs were produced worldwide, with 40% of this in North America alone (Richardson, 2011a).

Brominated flame retardants have been used for many years, and were updated in manufacturing use by polybrominated diphenyl ethers (PBDEs) since the polybrominated biphenyls were banned about 30 years ago. While penta- and octa-PBDEs were banned in various states, replacement fire retardant chemicals have been produced to keep pace with legislative regulations.

Perchlorate is now recognized as a worldwide environmental issue, and has been found in environmental waters across the United States and in other parts of the world. It has been
detected in food products in the US, in Europe, and the Far East. Perchlorate is very water-soluble and environmentally stable, and can accumulate in plants which can lead to exposure in humans and animals. Natural sources of perchlorate include naturally sourced nitrate fertilizer. Ammonium perchlorate is used as an oxidizer in solid propellants for rockets, fireworks, and highway flares. Perchlorate can also be a contaminant from the drinking water treatment chemical sodium hypochlorite, and is not removed by conventional water treatment processes, leading to potential for human exposure to occur through drinking water.

Additional CECs include nanomaterials, which are 1 to 100 nm in size and can have unique properties, including high strength, thermal stability, low permeability, and high conductivity. The chemical structures of nanomaterials are highly varied, including fullerenes, nanotubes, quantum dots, metal oxanes, TiO₂ nanoparticles (NPs), nanosilver, nanogold, and zerovalent iron NPs (Richardson, 2011a).

Pharmaceuticals and hormones have been detected in environmental waters. Concerns regarding pharmaceuticals and hormones include potential threats to drinking water, and possible estrogenic and other effects to wildlife and humans. An additional issue of the release of pharmaceuticals to the environment includes the development of bacterial resistance. These CECs are transported to the environment most commonly via effluent from WWTPs due to incomplete removal in wastewater treatment, and also through livestock farm discharges (Richardson, 2011a). An estimated 3200 different substances are used as pharmaceutical ingredients, including painkillers, antibiotics, antidiabetics, betablockers, contraceptives, lipid regulators, antidepressants, chemotherapy drugs, and impotence drugs, and only a very small subset of these compounds has been investigated in environmental studies.

Drinking Water disinfection byproducts (DBPs) were also identified as CECs, and are formed by the reaction of disinfectants (chlorine, chloramines, ozone, chlorine dioxide, etc.) with natural organic matter (NOM) and bromide or iodide in source waters. Nitrosamines, discovered as DBPs in 2002, are probable human carcinogens. Additional
new CECs include UV filters, used in sunscreens, cosmetics, and other personal care products. UV filters have potential for endocrine disruption and developmental toxicity. Environmental levels of UV filters were detected at levels close to the doses that cause toxic effects in animals (Richardson, 2011a).

1,4-Dioxane has been detected in environmental waters, has also been found in drinking water, and is a probable human carcinogen. Dioxane is a high production chemical used as a solvent stabilizer for the production of textiles including paper and cotton, and also in automotive coolants, cosmetics, and shampoos, as well as a stabilizer in 1,1,1-trichloroethane (TCA). In 2002 alone, an excess of 500 tons of dioxane were produced or imported in the United States.

Reports of benzotriazoles as an environmental contaminant have only been detected since approximately 2004, and studies indicate that they are likely ubiquitous environmental contaminants. Benzotriazoles are complexing agents widely used as anticorrosives. The two common forms, benzotriazole (1H-benzotriazole) and tolyltriazole (a mixture of 4- and 5-methyl-1H-benzotriazole), are water-soluble, resistant to biodegradation, and only partially removed in wastewater treatment.

Siloxanes have become a relatively new area of research of CECs, with concerns about potential toxicity and transport into the environment due to reportable quantities in wastewater, river water, and landfill biogases (Richardson, 2011b). Siloxanes are used in the production of a number of personal care products and common utensils and household products such as cosmetics, deodorants, soaps, hair conditioners, hair dyes, car waxes, baby pacifiers, cookware, cleaners, furniture polishes, and water-repellent windshield coatings.

Synthetic musk compounds are also a growing area of research of CECs, as they have been widely detected in wildlife and humans. Musks are highly lipophilic, and tend to accumulate in sediments, sludges, and biota. Musks are commonly used as fragrance
additives in many consumer products, including perfumes, lotions, sunscreens, deodorants, and laundry detergents.

Finally, microorganisms and algal toxins (mostly cyanobacterial toxins produced from blue-green algae) have also been reported as CECs. An unusual CEC identified in this review was sucralose, identified as a persistent (half-life up to several years) contaminant found in surface water, groundwater, and coastal waters (Richardson, 2011b). The research paper by Soh et al. (2011) stated that sucralose is one of very few contaminants that are highly persistent but do not bioaccumulate, and have little or no reported toxicity at environmentally relevant concentrations. Their paper raised an important question:

“Is persistence reason enough for concern or regulation?” (Soh et al., 2011)

2.1.1 Wastewater Treatment Plants as Sources of CECs

WWTPs are a major source of CECs, as these facilities are the main receptors of these products via disposal down household, commercial, and industrial drains, and transport via stormwater runoff (e.g., from areas treated with land-applied biosolids). CECs subsequently get introduced to the environment via wastewater effluent that drains to major waterbodies, via leachate after disposal of WWTP sludge in landfills, or transported with treated biosolids and applied to agricultural and other lands as a soil amendment. Approximately 5.6 million dry tons of biosolids are used or disposed of annually in the United States, of which approximately 60% is used for land application. EPA estimates that biosolids are applied to approximately 0.1% of available agricultural land in the United States on an annual basis. All biosolids are treated to achieve contaminant concentration limits, as established by the EPA under the 40 CFR Part 503 Standards for the Use or Disposal of Sewage Sludge. Based on the extent of treatment, biosolids may be land-applied under restricted and regulated conditions, as determined by the Part 503 Rule.
2.2 Relevant Environmental Regulatory Agencies and Environmental Regulations

Potential conflict of interests can arise between public health goals and commerce and industry regulations, as is often evident in the evolution of environmental legislation. Regulatory targets are complicated by the myriad factors and stakeholders that hold influence.

This section provides a brief history of relevant environmental regulatory agencies and environmental regulations and an overview of agencies providing regulatory oversight for environmental protection, including the regulation of CECs. Between 2012 and 2016, during which time this literature review was conducted, an effort was made to focus on the most current and relevant papers available at the time on CEC research. At the time of the 2012 research, the Toxic Substances Control Act (TSCA) reform was under consideration, and pending legislation. By the time of the 2016 research period, TSCA reform legislation had been passed by the Senate and signed by President Obama on June 22, 2016. TSCA reform is covered under Section 2.2.2.2. Additionally, a brief summary of international regulations is included for perspective, as well as reference for future direction of regulations.

2.2.1 United States Federal Environmental Agencies

This section provides a summary of the evolution and creation of U.S. Federal agencies providing regulatory oversight for environmental protection. The modern climate of the United States environmental regulatory process was inherited from decades-old public health traditions, from which environmental regulatory programs arose to address issues of public health as threatened by water and air quality issues. In addition, maritime navigation protection regulations arose to protect and promote commerce and industry related to harbors, ports, and otherwise navigable waters.
The U.S. Public Health Service (PHS) originally began in 1798 as the U.S. Marine Hospital Service. Congress changed the name to the “U.S. Public Health and Marine Service” in 1902, expanding its functions to deal with the broad issues of public health, and in 1912 to the PHS. Environmental authorities founded under the PHS included the National Air Pollution Control Administration, originally a research body founded in 1955, and the Federal Water Quality Administration (FWQA), authorized in 1965 by the Water Quality Act (WQA). The FWQA was authorized to issue federal water quality standards for interstate waters, where states failed to do so. The FWQA left the PHS in 1966 to become part of the Department of the Interior, and later was absorbed under the EPA.

The U.S. Environmental Protection Agency (EPA) was established in December 1970, effectively integrating the administration of a variety of federal research, monitoring, standard-setting and enforcement activities, to ensure environmental protection under the umbrella of a single agency. The founding of the EPA was the result of a hybrid of regulations and policy standards originally enacted under agencies such as the PHS and USACE, and many duties were transferred to the EPA from other Federal Agencies. The EPA was assembled from parcels of three federal Departments, three Bureaus, three Administrations, two Councils, one Commission, one Service, and many diverse offices (EPA, 1992).

2.2.2 United States Federal Environmental Regulations

This section provides a chronological history of environmental regulations, with a focus on regulations that pertain to water quality, or potential sources of water quality contamination. This background is referenced from the Congressional Research Service Report RL30798, Environmental Laws: Summaries of Major Statutes Administered by the Environmental Protection Agency (Bearden et al., 2013).

The Rivers and Harbors Appropriation Act of 1899 is the oldest environmental law in the United States, and is administered by the U.S. Army Corps of Engineers (USACE).
Section 9 of the Act, applying to bridges and causeways within navigable waters, was re-delegated to the U.S. Coast Guard (USCG) under the provisions of the Department of Transportation Act of 1966, based on the conflict of interest of both ownership and regulation by USACE of many bridges. Actions regulated under section 10 of the Act include excavation, fill, or alteration of any port, harbor or channel, including damming of navigable streams for the purposes of hydroelectric generation or other navigable purposes. Section 13 of the Act controls discharge of refuse of any kind into navigable waters of the United States without a permit, otherwise known as the Refuse Act. Additional environmental enforcement authority under the USCG includes jurisdiction of certain aspects of the Clean Water Act (CWA), such as enforcement of the Oil Pollution Act of 1990 (OPA).

The Federal Water Pollution Control Act (FWPCA) of 1948 was the first comprehensive federal clean water program, passed under Public Law (P.L.) 80-845 (Act of June 30, 1948), and administered under the PHS. At this time, water pollution was viewed as a state and local problem, and there were no federally required goals or guidelines. Federal enforcement and involvement was limited to issues over interstate waters, and only with the consent of the state from which the pollution originated. The FWPCA of 1948 specifically provided state and local government with technical assistance funding to address water pollution problems, including research grants.

In the latter half of the 1950s into the 1960s, water pollution control programs were modified by four laws amending the 1948 FWPCA statute, extending the federal role and federal jurisdiction to include navigable, as well as interstate waters. The four laws were:

- Water Pollution Control Act of 1956 (P.L. 84-660 (Act of July 9, 1956))
- 1961 Federal Water Pollution Control Act Amendments (P.L. 87-88)
- Water Quality Act of 1965 (P.L. 89-234)
- 1966 Clean Water Restoration Act (P.L 89-753)
Water quality standards became a feature of the law under the 1965 Water Quality Act, and required states to set standards for interstate waters to be used to determine actual pollution levels.

The Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA) was originally passed in 1947 to address shortcomings of the 1910 Federal Insecticide Act, and regulatory authority was assigned to the United States Department of Agriculture (USDA). FIFRA was revised in 1972 by the Federal Environmental Pesticide Control Act (FEPCA), which transferred authority to the EPA. The revised Act emphasized protection of the environment and human health, and shifted the burden of proof of environmental compliance of pesticide products to the chemical manufacturer (EPA, 1996a).

Other notable statutes transferred to the EPA at its inception included certain functions of the National Environmental Policy Act (NEPA) that pertain to ecological systems, and the Federal Water Pollution Control Act (FWPCA) of 1972, better known as the Clean Water Act (CWA).

### 2.2.2.1 Clean Water Act

The Federal Water Pollution Control Act Amendments of 1972 (P.L. 92-500), referred to as the Clean Water Act (CWA), totally revised the original 1948 FWPCA. The CWA is the primary federal law that governs water pollution, and established the basic structure for regulating the discharge of pollutants into the waters of the United States. This represented a fundamental change in Federal policy, as the CWA shifted the regulatory focus from water quality standards to effluent standards as the foundation for the strategy to control pollution from point sources, primarily industrial dischargers and publicly owned treatment works (POTWs), henceforth referred to as wastewater treatment plants (WWTPs) in this paper. Point sources are defined as discharges from a “discrete conveyance” (or outfall) by industrial facilities (including mining, manufacturing, oil and gas extraction, etc.), municipal governments and other government facilities (such as military bases), and some agricultural facilities (such as animal feed lots).
The Clean Water Act of 1972 transferred administration of the FWPCA to the EPA, in coordination with state governments. The 1972 law gave the EPA authority to develop pollution control programs such as setting wastewater and industry effluent standards, thereby establishing effluent limitations for the amounts of specific pollutants that may be discharged by municipal sewage plants and industrial facilities. Congress created a major public works financing program for bringing WWTPs up to treatment standards, authorized and funded under Title II.

The CWA also authorized the setting of water quality standards for all contaminants in surface waters. Title IV of the CWA made it unlawful to discharge any pollutant from a point source into navigable waters, unless a permit was obtained under the newly introduced National Pollutant Discharge Elimination System (NPDES), established under section 402. The NPDES program is the primary mechanism under the permit program for regulating point sources of pollution. Initially, the NPDES program focused on WWTPs and industrial wastewater. Nonpoint source pollution was not specifically addressed until the Water Quality Act (WQA) of 1987. A nonpoint source is defined as a diffuse source of pollution that does not have a point of origin. They include stormwater runoff from industrial sources, municipal storm drains, and agricultural stormwater discharges and irrigation return flows.

Additional major changes to the FWPCA have been introduced via additional amendments, including the Clean Water Act of 1977, the Water Quality Act of 1987, and the 1990 Oil Pollution Act. The WQA of 1987 expanded the NPDES program under CWA section 402. The updated program addressed certain nonpoint sources not subject under the 1972 CWA, in particular stormwater runoff, requiring separation of sewer and stormwater systems. Under the Stormwater Phase II Final Rule MS4 Program, operators of regulated small municipal separate storm and sewer systems are required to capture stormwater and provide stormwater treatment at the WWTP, instead of allowing direct discharge of nonpoint stormwater runoff to surface waterbodies. The permit exemption for agricultural discharges continued, but Congress created a grant program at EPA to
expand research of nonpoint controls and management practices. Additionally, the 1987 WQA created a program for management of biosolids generated by WWTPs.

The CWA was amended again in 1992 to set site-specific allowable pollutant levels for individual water bodies and to create an antidegradation policy to maintain and protect existing uses and high quality waters. Water bodies that do not meet applicable water quality standards are placed on the section 303(d) list, requiring development of a Total Maximum Daily Load (TMDL) of contaminant(s) specific for that water body. A TMDL establishes water quality-based limitations of the maximum amount of a pollutant that a water body can receive and still meet water quality standards. Once a TMDL is issued for a water body, appropriate modification of NPDES permits must be implemented to reflect the TMDL requirements.

### 2.2.2.2 Toxic Substances Control Act

The Toxic Substances Control Act (TSCA) (15 U.S.C. 2601), enacted in 1976, provided the EPA regulatory authority to protect the environment and consumers’ health against risks posed by chemicals in commerce. Chemicals in current-use are subject under TSCA, with the exception of chemicals regulated under other federal laws, such as those laws concerning food, drugs, cosmetics, firearms, ammunition, pesticides, tobacco, or mixtures. EPA is required by Section 8 of TSCA to develop and maintain an inventory of all chemicals, or categories of chemicals, manufactured or processed in the United States. EPA reviews approximately 700 new chemical manufacturing notices annually.

The TSCA inventory in 1979 identified approximately 55,000 chemicals in commerce. While this Act authorized EPA to passively screen new and existing chemicals used in U.S. manufacturing and commerce to identify potentially dangerous products or uses that should be subject to federal regulation, to require chemical manufacturers to conduct reporting and record-keeping, and require testing for chemical products, TSCA did not give EPA authority to independently evaluate the safety of a chemical before it went to marketplace. Based on these evaluations of new and existing chemicals and their
environmental risks, the EPA could only place restrictions relating to chemical substances and/or mixtures.

TSCA only retroactively addressed chemical hazards, after they had been deemed unsafe and previously unrestricted in the public. Title I, enacted in 1976, addressed two chemical substances, PCBs and elemental mercury. Specifically, Section 6(e) originally regulated PCBs and banned most uses, and with 2008 amendments restricting sales of elemental mercury (P.L. 110-414). In addition, five titles have been added to address the following specific chemical concerns:

- asbestos in 1986 (Title II, P.L. 99-519)
- radon in 1988 (Title III, P.L. 100-551)
- lead in 1992 (Title IV, P.L. 102-550)
- environmental issues in schools in 2007 (Title V, P.L. 110-140)
- formaldehyde in 2010 (Title VI, P.L. 111-199)

TSCA was formed with the intention to conduct and report test data by producers (i.e., manufacturers, importers, and processors) of chemicals in commerce, however the law did not give the EPA the ability to control substances on the market. A bill to overhaul TSCA, the Frank R. Launtenberg Chemical Safety for the 21st Century Act (House Amendment to the Senate Amendment to H.R. 2576, TSCA Modernization Act of 2015), was passed by the Senate on June 7, 2016, and sent on for President Obama’s signature. President Obama signed the Act into Law on June 22, 2016.

**TSCA Limitations and TSCA Reform**

Many chemicals, even some in widespread use, are not well characterized in terms of their potential health and environmental effects, and many of these are potential or existing CECs. TSCA is one of the first lines of defense in regulating new and existing CECs, but has been considered ineffective and out of date (Schierow, 2009), and incapable of assessing the safety of all chemicals in use today.
The Frank R. Lautenberg Chemical Safety for the 21st Century Act will amend TSCA to give the EPA new authority to evaluate the safety of a new chemical before it enters the marketplace. It will also allow EPA to evaluate the safety of chemicals already known to be risks, including chemicals found to persist in the human body and in the environment. The bill also limits companies’ ability to claim product information as confidential, allowing regulators, health professionals and the general public access to previously restricted information about the chemical components in chemical products. The TSCA Modernization Act of 2015 is presented in full in Appendix A.

The first version of the TSCA inventory in 1979 identified approximately 55,000 chemicals in commerce. Due to the overwhelming number of chemicals, Section 4(e) of TSCA established the Interagency Testing Committee (ITC) as an independent advisory committee to the Administrator of the EPA. The ITC was created to make recommendations on prioritizing and selecting chemicals for testing or information reporting to meet the data needs among government agencies. These chemicals are then added to the “Priority Testing List”.

Chemicals were assigned a higher priority if known or suspected to cause or contribute to cancer, gene mutations, or birth defects. Section 4(a) of TSCA directed the EPA to require test data to be reported on existing chemicals when certain conditions prevail, including:

- the manufacture, processing, distribution, use, or disposal of the chemical “may present an unreasonable risk;” (Sec. 4(a)(1)(A)(i)
- the chemical is produced in very large volume and there is a potential for a substantial quantity to be released into the environment or for substantial or significant human exposure. (Sec. 4(a)(1)(B)(i)

If either condition existed, EPA shall by rule require that testing be conducted if:

(ii) existing data are insufficient to resolve the question of safety, and
(iii) testing is necessary to develop the data
Section 5 and Section 6 of TSCA Title I directed EPA to require manufacturers and processors to conduct testing for existing chemicals in order to:

- prevent future risks through pre-manufacture screening and regulatory tracking of new chemical products (Section 5);
- control unreasonable risks already known, or as they are discovered for existing chemicals (Section 6).

TSCA also required EPA to be given a short notice of 90 days when there were plans to produce, process, or use an existing chemical in a way that differs from previously permitted uses, if the Administrator determined by rule that new uses of the chemical may produce significant changes in human and environmental exposures and therefore require notification. Although the legislative history of TSCA included a presumption that testing of new products would take place before they were widely used, either as the chemical was developed, or as its markets grew, TSCA forbade promulgation of blanket testing requirements for all new chemicals. This reflected a concern that uniform testing requirements could stifle innovation in the chemical industry. The purpose of the screening procedure was to identify potential hazards, and control them before use of a chemical becomes widespread; however, the ability of EPA to direct regulation of new chemical products was limited by the original TSCA language. Thus, EPA was restricted to determining only which chemicals, or which categories of chemicals, warrant the costs of premarket testing, and was required by TSCA to regulate only “to the extent necessary to protect adequately” against a risk, and to use the “least burdensome” regulatory approach, even in controlling unreasonable risks.

The TSCA Modernization Act of 2015 updates TSCA Section 6 – Prioritization, Risk Evaluation, and Regulation of Chemical Substance and Mixtures, subsection (a), by striking the language “to protect adequately against such risk using the least burdensome requirements,” and inserts, “so that the chemical substance or mixture no longer presents such risk.” Additionally, the TSCA Modernization Act of 2015 requires the Administrator to establish a risk-based screening process, including criteria for
designating substances as high-priority or low-priority for risk evaluations. The Act further clarifies that:

*The process to designate the priority of chemical substances shall include a consideration of the hazard and exposure potential of a chemical substance or a category of chemical substances (including consideration of persistence and bioaccumulation, potentially exposed or susceptible subpopulations and storage near significant sources of drinking water), the conditions of use or significant changes in the conditions of use of the chemical substance, and the volume or significant changes in the volume of the chemical substance manufactured or processed.*

The TSCA Modernization Act of 2015 also gives the EPA greater time periods for evaluating new chemicals, expands the authority of EPA to determine the risk-based screening process required by a manufacturer, and grants the authority to EPA to independently designate a chemical as high-risk. Chemical manufacturers will be required to make a safety finding to get a product on market, which gets away from the passive system as TSCA originally existed. Instead of requiring that EPA had to document that a chemical posed a risk before it could ask the manufacturer to conduct toxicity or exposure tests, EPA will be able to determine that chemicals in furniture, clothing, cleaning products and other common household items are safe before such products are allowed into commerce.

However, the Act still allows for some regulatory leeway in interpretation, as language in the legislation states EPA must consider the “cost-effectiveness” of any proposed rule, which may be considered restating the previously mentioned “least-burdensome requirements” on industry.
2.2.2.3 Additional Environmental Protection Acts

The Safe Drinking Water Act (1974), Title XIV of the Public Health Service Act, protects the public water supplies from harmful contaminants. The law focuses on all waters actually or potentially designed for drinking use, whether from above-ground or underground sources.

The Resource Conservation and Recovery Act (RCRA), passed in 1976, created authority for the EPA to control hazardous waste from “cradle to grave.” This includes the generation, transportation, treatment, storage and disposal of hazardous waste. This law also set a framework for the management of non-hazardous waste.

The Comprehensive Environmental Response, Compensation and Liability Act (CERCLA), enacted in 1980, provides for a federally funded “Superfund” to clean up uncontrolled or abandoned hazardous waste sites as well as accidents, spills and other releases of pollutants into the environment. To prioritize cleanup action, CERCLA directed EPA to create a National Priorities List (NPL) of the most contaminated sites which present the greatest risks to human health and the environment. The NPL includes both non-federal sites and federal facilities deemed to present a sufficient level of risk to warrant listing. The law gives the U.S. EPA the authority to locate the parties responsible for any release and assure their cooperation in the cleanup. Additionally, Section 104(i) of CERCLA established the Agency for Toxic Substances and Disease Registry (ATSDR) mainly to assess potential health risks at NPL sites. The ATSDR assesses individual sites based on the likelihood of human exposure to contamination through the air, soil, surface water, groundwater, and other pathways such as consumption of contaminated food sources. These assessments serve two purposes: to inform the public of potential health hazards at a contaminated site and to aid decision-makers in evaluating what cleanup actions may be warranted to prevent potentially harmful exposure.

The Pollution Prevention Act (1990) focuses industry, government and public attention on reducing pollution through cost-effective changes in production, operation and raw
material use. Pollution prevention also includes other practices such as source reduction, recycling and sustainable agriculture that increase efficiency in the use of energy, water and other natural resources.

2.2.2.4 Wastewater Treatment Plant and Land-Applied Biosolids Regulatory Requirements

Wastewater treatment produces two end products: effluent and sewage sludge. Biosolids are the treated component of sewage sludge. Approximately 75% of the United States population contributes to wastewater directly through a sewerage system to over 16,000 wastewater treatment plants (DHS, 2016).

Wastewater pretreatment regulations were established through 40 CFR Part 403 (as of June 26, 1978). These regulations addressed industrial facilities contributing to the influent stream, and aimed to prevent introduction of pollutants into the WWTPs that would interfere with the operation of a plant, including interference with disposal of municipal biosolids due to contamination. These regulations under Part 403 dramatically reduced concentrations of selected pollutants (e.g., pollutants which create a fire hazard, oil and grease, and corrosive pollutants, among other hazardous wastes) discharged to WWTPs and therefore reduced the concentrations of pollutants in treated biosolids.

Section 402 of the CWA (the NPDES permit program), and 40 CFR Part 503 (Standards for Use or Disposal of Sewage Sludge), allows regulation of land-applied biosolids from the consideration of a point-source discharge to groundwater. The 1993 Code of Federal Regulations, 40 CFR Part 503, under Section 405 of the 1987 WQA, created a program for general requirements, pollutant limits, management practices, and operational standards, for the final use or disposal of sewage sludge produced during treatment of domestic sewage in WWTPs. Contaminants considered under this rule were selected based in part on analytical results from the 1988 National Sewage Sludge Survey (NSSS), which analyzed sewage sludge for 411 possible contaminants from 176 WWTPs within the U.S.
The Part 503 rule established management practices for usage and disposal of biosolids, including land-application of biosolids, concentration limits, and loading rates for chemicals occurring in biosolids, and treatment and use requirements designed to control and reduce pathogens (EPA, 1993). The chemical land-application standards in the Part 503 rule were determined by EPA through risk assessments aimed at identifying the chemical constituents in biosolids judged likely to pose the greatest hazard. Likely exposure scenarios and calculated concentration limits and loading rates were also identified. The regulations to guide the management practices and operational standards are as follows:

1. Identify uses for sewage sludge, including disposal;
2. Specify factors to be taken into account in determining the measures and practices applicable to each such use or disposal (including publication of information on costs); and
3. Identify concentrations of pollutants which interfere with each such use or disposal.

Under Part 503, the first primary regulated contaminants in biosolids were ten inorganic chemicals: arsenic, cadmium, chromium (limits later deleted under 60 FR 54764), copper, lead, mercury, molybdenum, nickel, selenium, and zinc.

Different land application rules apply to different classes of biosolids, and the following quality control regulations were established under the Part 503 regulation:

Two levels of sewage sludge quality with respect to heavy metal concentrations:
- Pollutant ceiling concentrations
- Pollutant concentrations ("high quality" sewage sludge)

Two levels of quality with respect to pathogen densities:
- Class A
- Class B
Two types of approaches for meeting vector attraction reduction:

- Sewage sludge processing
- The use of physical barriers.

Class A biosolids contain no detectible levels of pathogens, must meet strict vector attraction reduction requirements and low levels metals contents, and are permitted to ensure that these standards have been met. Class B biosolids are treated but still contain detectible levels of pathogens, and have buffer requirements to protect water quality, restricted public access, and crop harvesting restrictions for virtually all forms of Class B biosolids.

Under the Part 503 regulation, fewer restrictions are imposed on the use of higher quality sewage sludge. Class A biosolids may be applied in small quantities without restriction by the general public on public-contact sites, including parks, golf courses, lawns, and home gardens. When used in bulk, Class A biosolids are subject to buffers for water quality protection, but not to crop harvesting restrictions. Based on the extent of treatment, Class B biosolids may be applied where little exposure of the general public is expected to occur on the sites, such as on agricultural land, forests, and reclamation sites.

In 1995 EPA revisited the pollutants considered in Rule 503 under the so-called “Round Two” evaluation. However, a second comprehensive analytical survey of contaminants in biosolids was not conducted, and EPA instead focused largely on compounds previously considered during the original NSSS, with an emphasis on chlorinated dioxins, furans and co-planar PCBs (EPA, 1996b). In 2003, EPA concluded that these compounds did not present a significant risk to human health or the environment, and made the decision to not regulate levels of these compounds in biosolids (EPA Headquarters Press Release October 17, 2003).

As public health concerns regarding the use of biosolids increased, EPA in 1999 asked the National Academy of Science (NAS) to conduct an independent evaluation
reassessing the scientific basis of the Part 503 Rule. NAS produced the resulting report *Biosolids Applied to Lands: Advancing Practices and Standards* in 2002. The NAS committee stated in the 2002 report that the Part 503 Rule relied on an outdated biosolid contaminant characterization (NRC, 2002), and that the original NSSS did not address possible adverse changes in biosolids composition due to changes in treatment processes and chemical uses over the last decade. Suggested actions recommended by the committee included that a new national sewage sludge survey be conducted, to ensure that Part 503 Rule risk assessment assumptions are based on sound science (NRC 2002). At the 2003 Biosolids Research Summit, a research agenda was created to address research gaps identified by the NRC report. The top identified research gaps were to create a targeted characterization of pathogens, and to conduct a new national survey of CECs in biosolids. Both were ranked as the second and third highest research priorities, following only the development of a rapid incident response study aimed at examining whether a linkage existed between biosolids land application and reports of human health impacts (WERF, 2004).

### 2.2.3 International Environmental Regulatory Agencies and Environmental Regulations

Europe has passed tough chemical regulations, including one of the world’s most extensive chemical safety regulations, known as REACH (Registration, Evaluation, Authorization and Restriction of Chemicals), passed on June 1, 2007. The European Chemicals Agency (ECHA) is the prime regulatory authority for REACH, and helps companies comply with the legislation, advances the safe use of chemicals, and provides information on chemicals and addresses chemicals of concern (ECHA REACH, 2016). The law requires companies that produce or sell chemicals in the European Union (EU) to register toxicity data on the compounds and to outline any new tests needed to clarify their biological effects and places the burden of proof on companies. To comply with the regulation, companies must identify and manage the risks linked to the substances they manufacture and market in the EU.
The European Union Council Directive 86/278/EEC was adopted over 20 years ago to promote land application of biosolids in agriculture and to regulate its use, to prevent harmful effects on soil, vegetation, animals, and humans. The Directive was initiated June 12, 1986, with the intent of the protection of the environment, and in particular of the soil, when sewage sludge is used in agriculture. The Directive currently sets limit values for seven heavy metals that may be toxic to plants and humans: cadmium, chromium, copper, nickel, lead, zinc, and mercury. Since its adoption, several Member States have enacted and implemented stricter limit values for heavy metals and set requirements for other contaminants. Currently, the European Commission is assessing whether the current Directive should be reviewed, and to what extent (EC, 2016).
3. Case Study on Columbia River

The Columbia River is presented as a case study as an example of a significant waterbody with research and regulatory gaps concerning CECs.

3.1 Introduction/Problem Statement

The Columbia River is important to the entire Pacific Northwest region as a food and water source, for transportation and shipping, for recreation, and also as a cultural resource for the many tribes within the Columbia River watershed. The Columbia spans three states and two nations, which makes for complex oversight authority.

The Columbia River is a receptor of numerous point-sources of pollution throughout its length, including urban stormwater runoff, wastewater treatment plant and industrial manufacturing effluent, as well as non-point source pollution such as overland flow from agricultural and industrial fields and air deposition of contaminants. Currently, the Columbia River is extensively monitored for known contaminants (mercury, PCBs, DDT, PBDEs) while limited resources are dedicated to identifying and monitoring emerging contaminants in the Columbia River. Additionally, the Columbia River Basin is a prime candidate for targeted studies on the effects land-applied biosolids within the watershed.

The purpose of this case study is to present an in-depth compilation of current research on emerging contaminants and to identify potential new monitoring methods and programs. This case study also provides an overview of the current Columbia River Monitoring Plan and describes current actions being undertaken by monitoring agencies and resource management agencies to monitor and control emerging contaminants.
3.2 Relevant Agencies and Programs that Affect the Columbia River

This section provides a summary of regional agencies and watershed protection groups providing regulatory oversight for environmental protection of the Columbia River basin.

**National Estuary Program and LCREP**

Between 1989 and 1995, a six-year study was conducted by the Bi-State Water Quality Program on the lower Columbia River between the Bonneville Dam and the Pacific Ocean. Research studies collected water quality data and toxic contaminant data within the lower Columbia River and estuary, generating a large dataset on the threats to the health of the river and river organisms.

The findings of the six-year study highlighted four problems in the lower Columbia River estuary:

- Toxics in sediment and fish tissue that can affect the health of humans, fish, and wildlife
- Habitat loss/modification that can affect fish and wildlife resources
- Water quality problems that affect beneficial uses in portions of the estuary
- Overall decline in anadromous fish runs that has resulted in threatened and endangered species listings (WDOE, 1997)

Based on the results of this research, the lower Columbia River was nominated by the governors of Washington and Oregon states for the EPA National Estuary Program (NEP) in July 1995. The NEP is a non-regulatory program established by Congress via amendments to the Clean Water Act in 1987, and is a place-based program established to protect and restore “water quality and ecological integrity of estuaries of national significance” (EPA NEP Overview 2016). The NEP provides support for the development of management plans for the Nation’s most significant estuaries that are threatened by degradation caused by human activity. Currently, 28 estuaries are
designated as significant, and the program provides resources to help manage these estuaries of national, regional, and local significance.

The EPA accepted the lower Columbia River into the NEP, and established the Lower Columbia Estuary Partnership (LCREP). LCREP is supported by the US Environmental Protection Agency, the states of Oregon and Washington, and numerous other public and private entities. LCREP advocates for long-term monitoring, to address issues about the sources, distribution, and persistence of toxics in the lower river.

Columbia River Toxics Reduction Work Group

As part of the NEP, a long-term monitoring plan was developed for the lower Columbia River and estuary to address many of the issues that face the lower Columbia River Basin. The Columbia River Toxics Reduction Work Group (CRTRWG) was formed in 2005 as a collaborative workgroup among EPA and federal, state, tribal, local, industry, and nonprofit partners to focus on toxics in the Columbia River, with the goal to reduce toxics in the Columbia River Basin and prevent further contamination. This group was modeled on EPA collaborative efforts underway throughout the U.S. including the Chesapeake Bay and the National Estuary Program (EPA and CRTRWG, 2010). In 2009, the EPA produced *The Columbia River Basin State of the River Report for Toxics* with the support and guidance of CRTRWG.

The *State of the River Report for Toxics* determined that four contaminants were above risk action levels: mercury, dichlorodiphenyltrichloroethane (DDT) and associated breakdown products, polychlorinated biphenyls (PCBs), and polybrominated diphenyl ether (PBDE) flame retardants. Additional contaminants are found in the Basin, including arsenic, dioxins, radionuclides, lead, pesticides, industrial chemicals, and CECs including pharmaceuticals found in wastewater.

EPA and the CRTRWG released the Columbia River Toxics Reduction Action Plan (Action Plan) in September 2010. The Action Plan included five general initiatives and
actions to be accomplished through 2015, to better understand and reduce toxic contamination in the Columbia River Basin:

- Increase public understanding and political commitment to toxics reduction in the Basin
- Increase toxic reduction actions
- Conduct monitoring to identify sources and then work to reduce toxic contamination
- Develop a regional, multi-agency research program
- Develop a data management system that will allow sharing of information on toxics in the Basin

The work with the CRTRWG is currently conducted via coordination and partnerships without any designated funding sources, with the exception of work done in the estuary through LCREP. The CRTRWG states that:

“To a great extent, success in reducing toxics in the Basin will depend on a commitment by all levels of government, in both the United States and Canada, tribal governments, nongovernmental organizations, industry groups, and the public to work together. The problems are too large, widespread, and complex to be solved by only one organization or country.”

3.3 Gaps in Columbia River Research and Restoration Funding

The CRTRWG identified a number of priorities for state and local governments to address to reduce toxics in the Columbia Basin. The workgroup believes that a focus on enhancing programs in these areas will help advance the prevention and reduction of toxics in the Columbia River Basin. The Priority Initiatives of the Columbia River Toxics Reduction Working Group, January 2013, are summarized below:
• **Sustainable Purchasing**: Develop a list of sustainable products and a list of chemicals of concern that could be used by all entities in the Basin for greening their operations

• **Green Chemistry**: Initiate a Regional Green Chemistry Center to advance the discussion on how to develop chemical processes that provide less toxic materials

• **Chemicals of Emerging Concern**: Develop monitoring programs and toxicity information to inform actions to address chemicals of emerging concern that are currently unregulated

• **Pesticide Stewardship Partnerships**: Enhance and expand the successful Pesticide Stewardship Partnership model used by the State of Oregon to encourage voluntary changes in pesticide use and practices that lead to measurable environmental improvements

• **Stormwater**: Expand stormwater technical assistance programs to small and medium businesses, providing much-needed pollution prevention expertise at the local level

Contaminants such as DDTs have been determined to persist in the Columbia River despite being banned decades ago, while CECs, including flame retardants (PBDEs) and personal care products, pose new threats to human health and fish and wildlife. However, the Columbia River Basin is the only major EPA-designated ‘large aquatic ecosystem’ to receive zero funding pursuant to the NEP designation. Additionally, the Columbia River Monitoring Plan does not address CECs in biosolids.

In May 2015, U.S Senator Jeff Merkley (D-OR) and Congressman Earl Blumenauer (D-OR), reintroduced the Columbia River Basin Restoration Act. On September 15, 2016, the U.S. Senate passed the Act as part of the Water Resources Development Act (WRDA) of 2016 (LCREP September 16, 2016). This bill addresses critical fish and wildlife habitat, water quality, and infrastructure needs in a total of 18 states and would authorize Congress to appropriate funds for a voluntary grant program to expand and add to monitoring efforts and provide the resources for sustained action to reduce
contaminants and evaluate them throughout the Columbia River system. If the Act becomes law, which has not yet been passed as of the time of this report, the status of the Columbia River would be elevated to that of other Large Aquatic Ecosystems, and it would authorize Congress to appropriate funds through the EPA to implement its Columbia Basin Toxics Reduction Plan and the Estuary Partnership Management Plan.
Chapter 4

4. Conclusions

With over 100,000 chemicals in current use and new chemicals added each year, the environmental risks and behavior of these chemicals need to be properly evaluated and regulated. Ideally, problematic chemicals should be identified prior to their release into the environment and controlled accordingly. TSCA is one of the first lines of defense in regulating new and existing CECs, and is an important regulatory tool to assess the safety of chemicals in use today. Additionally, WWTPs are a destination for environmental contaminants sourced from residential and industrial sewer and storm drains, and many of these contaminants end up partitioning onto the treated sewage sludge. This sewage sludge that turns into treated biosolids then becomes a vector for environmental contamination as it is dispersed as land-applied biosolids.

4.1 Research Gaps

Data gaps exist for CECs in understanding their occurrence, persistence, transport and fate, their transformation products, the characterization of CECs in biosolids, their risks to human and environmental health, and research on prioritizing CECs for regulatory purposes. Data gaps exist on national and regional scales for impacts from CECs released from biosolids to environments such as the Columbia River Basin. Additional concerns regarding CECs in environment include unknown exposure duration and mixture effects, or unknown synergistic and cumulative effects of CECs in the environment.

Questions raised by this research include:

- How do you decide which pollutants to permit? Toxicologists defer to EPA, and human-health revisions are based on EPA reference doses, etc.
- Toxicity data is needed to reach conclusions on regulating compounds
- Standards are needed for methods to perform monitoring requirements
This literature review also concluded that many new CECs were identified due to improved analytical techniques that have allowed detection levels at previously undetectable limits in the low ng/L concentrations. This is significant, as prior analytical techniques did not allow for such low detection levels, and many chemicals went undetected in the environment, their presence unknown previous to the new low-detection analytical techniques.

4.2 Legislative Gaps

The reformed TSCA law still allows for some regulatory leeway in interpretation and regulatory enforcement. The regulatory language states EPA must consider the “cost-effectiveness” of any proposed rule, which may be considered restating the previously mentioned “least-burdensome requirements” on industry. Strong legislation is needed to predict and prevent chemicals from being unnecessarily released to the environment. Additionally, wastewater pretreatment regulations and wastewater treatment methods can be improved to reduce CECs from both WWTP influent streams and effluent products.

The extent of environmental protection afforded by the current evaluation approaches for risk-based assessment does not consider the cumulative risk of the mixture of all CECs present. A multifaceted approach is needed to address these challenges, including a set of tools to characterize CEC exposure at the suborganism, organism, and population levels, to identify potential or actual effects of CECs on aquatic communities. Possible approaches include incorporating existing approaches used by the various water agencies to assess the risk of individual chemicals. However, addressing the extent of the potential effects of mixtures of the chemicals in combination with more common pollutants and other environmental stressors is complex.

There is a need to characterize and better understand the environmental and biological fate, transport, and transformation of CECs. This knowledge gap becomes more important as water supplies become more limited and water re-use practices change.
More than 40,000 organic chemicals have been identified as CECs, which does not include the associated breakdown products in the environment. However, agencies responsible for monitoring efforts, such as the National Oceanic and Atmospheric Administration (NOAA), the U.S. Geological Survey (USGS), and the U.S. Environmental Protection Agency (U.S. EPA), have widely different definitions as to what a CEC actually is (Diamond, 2011). For example, some researchers consider an already regulated chemical to be a CEC if there are additional, unregulated effects, such as endocrine disruption. Others broadly define a CEC as a chemical that is currently unregulated. With no consensus on the definition of a CEC, each agency monitors its own subjective list of chemicals.

4.3 Summary

Biosolids are composed of numerous constituents, with a number of environmentally persistent and potentially toxic CECs identified as present in large concentrations. The presence of priority pollutants such as heavy metals and polychlorinated biphenyls (PCBs) in land-applied biosolids has been extensively studied, including the effects on human health and the environment, and environmental behavior. However, with the constantly increasing consumption of unregulated chemicals and the detection of significant concentrations of these chemicals in biosolids, research is needed to understand the transport and fate mechanisms of these compounds in biosolids. Additionally, this research is needed to determine a new priority framework to regulate CECs, both on the national level and regional levels. WWTP influent and effluent treatment regulations and practices may be improved upon as more knowledge accrues regarding CEC behavior in the environment. Understanding the concentrations and environmental behavior of CECs in land-applied biosolids is key to determining the significance and implications of these emerging pollutants on human health and the environment.
References


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Appendix A – TSCA Modernization Act of 2015
MAY 20, 2016

RULES COMMITTEE PRINT 114-54

TEXT OF HOUSE AMENDMENT TO THE SENATE AMENDMENT TO H.R. 2576, TSCA MODERNIZATION ACT OF 2015

[Showing the text of the Frank R. Launtenberg Chemical Safety for the 21st Century Act.]

SECTION 1. SHORT TITLE; TABLE OF CONTENTS.

(a) SHORT TITLE.—This Act may be cited as the “Frank R. Launtenberg Chemical Safety for the 21st Century Act”.

(b) TABLE OF CONTENTS.—The table of contents of this Act is as follows:

Sec. 1. Short title; table of contents.

TITLE I—CHEMICAL SAFETY

Sec. 2. Findings, policy, and intent.
Sec. 3. Definitions.
Sec. 4. Testing of chemical substances and mixtures.
Sec. 5. Manufacturing and processing notices.
Sec. 6. Prioritization, risk evaluation, and regulation of chemical substances and mixtures.
Sec. 7. Imminent hazards.
Sec. 8. Reporting and retention of information.
Sec. 9. Relationship to other Federal laws.
Sec. 10. Exports of elemental mercury.
Sec. 11. Confidential information.
Sec. 12. Penalties.
Sec. 15. Citizens’ civil actions.
Sec. 16. Studies.
Sec. 17. Administration of the Act.
Sec. 18. State programs.
Sec. 19. Conforming amendments.
Sec. 20. No retroactivity.
Sec. 21. Trevor’s Law.

TITLE II—RURAL HEALTHCARE CONNECTIVITY

Sec. 201. Short title.
Sec. 202. Telecommunications services for skilled nursing facilities.

1 TITLE I—CHEMICAL SAFETY

2 SEC. 2. FINDINGS, POLICY, AND INTENT.

3 Section 2(c) of the Toxic Substances Control Act (15
4 U.S.C. 2601(c)) is amended by striking “proposes to
5 take” and inserting “proposes as provided”.

6 SEC. 3. DEFINITIONS.

7 Section 3 of the Toxic Substances Control Act (15
8 U.S.C. 2602) is amended—

9 (1) by redesignating paragraphs (4) through
10 (14) as paragraphs (5), (6), (8), (9), (10), (11),
11 (13), (14), (15), (16), and (17), respectively;
12 (2) by inserting after paragraph (3) the fol-
13 lowing:

“(4) The term ‘conditions of use’ means the cir-
14 cumstances, as determined by the Administrator, under
15 which a chemical substance is intended, known, or reason-
16 ably foreseen to be manufactured, processed, distributed
17 in commerce, used, or disposed of.”;
18 (3) by inserting after paragraph (6), as so re-
19 designated, the following:

“(7) The term ‘guidance’ means any significant writ-
20 ten guidance of general applicability prepared by the Ad-
21 ministrator.”; and
(4) by inserting after paragraph (11), as so re-designated, the following:

“(12) The term ‘potentially exposed or susceptible subpopulation’ means a group of individuals within the general population identified by the Administrator who, due to either greater susceptibility or greater exposure, may be at greater risk than the general population of adverse health effects from exposure to a chemical substance or mixture, such as infants, children, pregnant women, workers, or the elderly.”.

SEC. 4. TESTING OF CHEMICAL SUBSTANCES AND MIXTURES.

Section 4 of the Toxic Substances Control Act (15 U.S.C. 2603) is amended—

(1) by striking “standards” each place it appears and inserting “protocols and methodologies”; (2) in subsection (a)—

(A) by striking “If the Administrator finds” and inserting “(1) If the Administrator finds”;

(B) in paragraph (1), as so designated—

(i) by striking “(1)(A)(i)” and inserting “(A)(i)(I)”;

(ii) by striking “(ii)” each place it appears and inserting “(II)”;

(3) by inserting after paragraph (11), as so re-designated, the following:

“(12) The term ‘potentially exposed or susceptible subpopulation’ means a group of individuals within the general population identified by the Administrator who, due to either greater susceptibility or greater exposure, may be at greater risk than the general population of adverse health effects from exposure to a chemical substance or mixture, such as infants, children, pregnant women, workers, or the elderly.”.

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(A) by striking “If the Administrator finds” and inserting “(1) If the Administrator finds”;

(B) in paragraph (1), as so designated—

(i) by striking “(1)(A)(i)” and inserting “(A)(i)(I)”;

(ii) by striking “(ii)” each place it appears and inserting “(II)”;

(3) by inserting after paragraph (11), as so re-designated, the following:

“(12) The term ‘potentially exposed or susceptible subpopulation’ means a group of individuals within the general population identified by the Administrator who, due to either greater susceptibility or greater exposure, may be at greater risk than the general population of adverse health effects from exposure to a chemical substance or mixture, such as infants, children, pregnant women, workers, or the elderly.”.
(iii) by striking “are insufficient data” and inserting “is insufficient information” each place it appears;

(iv) by striking “(iii)” each place it appears and inserting “(III)”;

(v) by striking “such data” and inserting “such information” each place it appears;

(vi) by striking “(B)(i)” and inserting “(ii)(I)”;

(vii) by striking “(I)” and inserting “(aa)”;

(viii) by striking “(II)” and inserting “(bb)”;

(ix) by striking “(2)” and inserting “(B)”; and

(x) in the matter following subparagraph (B), as so redesignated—

(I) by inserting “, or, in the case of a chemical substance or mixture described in subparagraph (A)(i), by rule, order, or consent agreement,” after “rule”;
(II) by striking “data” each place it appears and inserting “information”; and

(III) by striking “and which are relevant” and inserting “and which is relevant”; and

(C) by adding at the end the following:

“(2) ADDITIONAL TESTING AUTHORITY.—In addition to the authority provided under paragraph (1), the Administrator may, by rule, order, or consent agreement—

“(A) require the development of new information relating to a chemical substance or mixture if the Administrator determines that the information is necessary—

“(i) to review a notice under section 5 or to perform a risk evaluation under section 6(b);  

“(ii) to implement a requirement imposed in a rule, order, or consent agreement under subsection (e) or (f) of section 5 or in a rule promulgated under section 6(a);  

“(iii) at the request of a Federal implementing authority under another Fed-
eral law, to meet the regulatory testing

needs of that authority with regard to tox-
icity and exposure; or

“(iv) pursuant to section 12(a)(2);

and

“(B) require the development of new infor-
mation for the purposes of prioritizing a chem-
ical substance under section 6(b) only if the Ad-
ministrator determines that such information is
necessary to establish the priority of the sub-
stance, subject to the limitations that—

“(i) not later than 90 days after the
date of receipt of information regarding a
chemical substance complying with a rule,
order, or consent agreement under this
subparagraph, the Administrator shall des-
ignate the chemical substance as a high-
priority substance or a low-priority sub-
stance; and

“(ii) information required by the Ad-
ministrator under this subparagraph shall
not be required for the purposes of estab-
lishing or implementing a minimum infor-
mation requirement of broader applica-
bility.
“(3) STATEMENT OF NEED.—When requiring the development of new information relating to a chemical substance or mixture under paragraph (2), the Administrator shall identify the need for the new information, describe how information reasonably available to the Administrator was used to inform the decision to require new information, explain the basis for any decision that requires the use of vertebrate animals, and, as applicable, explain why issuance of an order is warranted instead of promulgating a rule or entering into a consent agreement.

“(4) TIERED TESTING.—When requiring the development of new information under this subsection, the Administrator shall employ a tiered screening and testing process, under which the results of screening-level tests or assessments of available information inform the decision as to whether 1 or more additional tests are necessary, unless information available to the Administrator justifies more advanced testing of potential health or environmental effects or potential exposure without first conducting screening-level testing.”;

(3) in subsection (b)—

(A) in paragraph (1)—
(i) in subparagraph (B), by striking “test data” and inserting “information”;

(ii) in subparagraph (C), by striking “data” and inserting “information”; and

(iii) in the matter following subparagraph (C), by striking “data” and inserting “information”;

(B) in paragraph (2)—

(i) in subparagraph (A)—

(I) by striking “test data” and inserting “information”;

(II) by inserting “Protocols and methodologies for the development of information may also be prescribed for the assessment of exposure or exposure potential to humans or the environment.” after the first sentence;

and

(III) by striking “hierarchical tests” and inserting “tiered testing”;

and

(ii) in subparagraph (B), by striking “data” and inserting “information”;

(C) in paragraph (3)—
(i) by striking “data” each place it appears and inserting “information”;  

(ii) in subparagraph (A), by inserting “or (C), as applicable,” after “subpara- 

graph (B)”;

(iii) by striking “(a)(1)(A)(ii) or (a)(1)(B)(ii)” each place it appears in sub-

paragraph (B) and inserting “(a)(1)(A)(i)(II) or (a)(1)(A)(ii)(II)”;

(iv) in subparagraph (B), in the matter before clause (i), by striking “sub-

section (a)” and inserting “subsection (a)(1)”;

(v) by adding at the end the following:

“(C) A rule or order under paragraph (1) or (2) of subsection (a) may require the development of information by any person who manufactures or processes, or intends to manufacture or process, a chemical substance or mixture subject to the rule or order.”;

(D) in paragraph (4)—

(i) by striking “of data” each place it appears and inserting “of information”; and
(ii) by striking “test data” each place it appears and inserting “information”; and

(E) by striking paragraph (5);

(4) in subsection (c)—

(A) in paragraph (1), by striking “data” and inserting “information”;  

(B) in paragraph (2), by striking “data” each place it appears and inserting “information”;  

(C) in paragraph (3)—

(i) by striking “test data” each place it appears and inserting “information”;  

and  

(ii) by striking “such data” each place it appears and inserting “such information”; and

(D) in paragraph (4) by striking “test data” each place it appears and inserting “information”;  

(5) in subsection (d)—

(A) by striking “test data” each place it appears and inserting “information”;  

(B) by striking “such data” each place it appears and inserting “such information”; and
(C) by striking “for which data have” and inserting “for which information has”;

(6) in subsection (e)—

(A) in paragraph (1)—

(i) in subparagraph (A)—

(I) by striking “promulgation of a rule” and inserting “development of information”; and

(II) by striking “data” each place it appears and inserting “information”; and

(ii) in subparagraph (B), by striking “either initiate a rulemaking proceeding under subsection (a) or if such a proceeding is not initiated within such period, publish in the Federal Register the Administrator’s reason for not initiating such a proceeding” and insert “issue an order, enter into a consent agreement, or initiate a rulemaking proceeding under subsection (a), or, if such an order or consent agreement is not issued or such a proceeding is not initiated within such period, publish in the Federal Register the Administrator’s reason for not issuing such an order, en-
entering into such a consent agreement, or
initiating such a proceeding”; and
(B) in paragraph (2)(A)—
(i) by striking “eight members” and
inserting “ten members”; and
(ii) by adding at the end the follow-
ing:
“(ix) One member appointed by the Chairman
of the Consumer Product Safety Commission from
Commissioners or employees of the Commission.
“(x) One member appointed by the Commis-
sioner of Food and Drugs from employees of the
Food and Drug Administration.”;
(7) in subsection (f)—
(A) in paragraph (1), by striking “test
data” and inserting “information”; and
(B) in the matter following paragraph
(2)—
(i) by striking “from cancer, gene
mutations, or birth defects”;
(ii) by striking “data or”;
(iii) by striking “appropriate” and in-
serting “applicable”; and
(iv) by inserting “, made without con-
sideration of costs or other nonrisk fac-
tors,” after “publish in the Federal Register a finding”;

(8) in subsection (g)—

(A) by amending the subsection heading to read as follows: “PETITION FOR PROTOCOLS AND METHODOLOGIES FOR THE DEVELOPMENT OF INFORMATION”;

(B) by striking “test data” each place it appears and inserting “information”; and

(C) by striking “submit data” and inserting “submit information”; and

(9) by adding at the end the following:

“(h) REDUCTION OF TESTING ON VERTEBRATES.—

“(1) IN GENERAL.—The Administrator shall reduce and replace, to the extent practicable, scientifically justified, and consistent with the policies of this title, the use of vertebrate animals in the testing of chemical substances or mixtures under this title by—

“(A) prior to making a request or adopting a requirement for testing using vertebrate animals, and in accordance with subsection (a)(3), taking into consideration, as appropriate and to the extent practicable and scientifically justi-
fied, reasonably available existing information, including—

“(i) toxicity information;

“(ii) computational toxicology and bioinformatics; and

“(iii) high-throughput screening methods and the prediction models of those methods; and

“(B) encouraging and facilitating—

“(i) the use of scientifically valid test methods and strategies that reduce or replace the use of vertebrate animals while providing information of equivalent or better scientific quality and relevance that will support regulatory decisions under this title;

“(ii) the grouping of 2 or more chemical substances into scientifically appropriate categories in cases in which testing of a chemical substance would provide scientifically valid and useful information on other chemical substances in the category; and

“(iii) the formation of industry consortia to jointly conduct testing to avoid
unnecessary duplication of tests, provided
that such consortia make all information
from such testing available to the Adminis-
trator.

“(2) Implementation of alternative testing methods.—To promote the development and
timely incorporation of new scientifically valid test
methods and strategies that are not based on
vertebrate animals, the Administrator shall—

“(A) not later than 2 years after the date
of enactment of the Frank R. Launtenberg
Chemical Safety for the 21st Century Act, de-
velop a strategic plan to promote the develop-
ment and implementation of alternative test
methods and strategies to reduce, refine, or re-
place vertebrate animal testing and provide in-
formation of equivalent or better scientific qual-
ity and relevance for assessing risks of injury to
health or the environment of chemical sub-
stances or mixtures through, for example—

“(i) computational toxicology and
bioinformatics;

“(ii) high-throughput screening meth-
ods;
“(iii) testing of categories of chemical substances;

“(iv) tiered testing methods;

“(v) in vitro studies;

“(vi) systems biology;

“(vii) new or revised methods identified by validation bodies such as the Inter-agency Coordinating Committee on the Validation of Alternative Methods or the Organization for Economic Co-operation and Development; or

“(viii) industry consortia that develop information submitted under this title;

“(B) as practicable, ensure that the strategic plan developed under subparagraph (A) is reflected in the development of requirements for testing under this section;

“(C) include in the strategic plan developed under subparagraph (A) a list, which the Administrator shall update on a regular basis, of particular alternative test methods or strategies the Administrator has identified that do not require new vertebrate animal testing and are scientifically reliable, relevant, and capable of providing information of equivalent or better
scientific reliability and quality to that which
would be obtained from vertebrate animal test-
ing;

“(D) provide an opportunity for public no-
tice and comment on the contents of the plan
developed under subparagraph (A), including
the criteria for considering scientific reliability
and relevance of the test methods and strate-
gies that may be identified pursuant to sub-
paragraph (C);

“(E) beginning on the date that is 5 years
after the date of enactment of the Frank R.
Lautenberg Chemical Safety for the 21st Cen-
tury Act, and every 5 years thereafter, submit
to Congress a report that describes the progress
made in implementing the plan developed under
subparagraph (A) and goals for future alter-
native test methods and strategies implementa-
tion; and

“(F) prioritize and, to the extent con-
sistent with available resources and the Admin-
istrator’s other responsibilities under this title,
carry out performance assessment, validation,
and translational studies to accelerate the devel-
opment of scientifically valid test methods and
strategies that reduce, refine, or replace the use of vertebrate animals, including minimizing duplication, in any testing under this title.

“(3) VOLUNTARY TESTING.—

“(A) IN GENERAL.—Any person developing information for submission under this title on a voluntary basis and not pursuant to any request or requirement by the Administrator shall first attempt to develop the information by means of an alternative test method or strategy identified by the Administrator pursuant to paragraph (2)(C), if the Administrator has identified such a test method or strategy for the development of such information, before conducting new vertebrate animal testing.

“(B) EFFECT OF PARAGRAPH.—Nothing in this paragraph shall, under any circumstance, limit or restrict the submission of any existing information to the Administrator.

“(C) RELATIONSHIP TO OTHER LAW.—A violation of this paragraph shall not be a prohibited act under section 15.

“(D) REVIEW OF MEANS.—This paragraph authorizes, but does not require, the Administrator to review the means by which a person
conducted testing described in subparagraph (A).”.

SEC. 5. MANUFACTURING AND PROCESSING NOTICES.

Section 5 of the Toxic Substances Control Act (15 U.S.C. 2604) is amended—

(1) in subsection (a)—

(A) in paragraph (1)—

(i) by striking “Except as provided in” and inserting “(A) Except as provided in subparagraph (B) of this paragraph and”;

(ii) by redesignating subparagraphs (A) and (B) as clauses (i) and (ii), respectively;

(iii) by striking all that follows “significant new use” and inserting a period; and

(iv) by adding at the end the following:

“(B) A person may take the actions described in subparagraph (A) if—

“(i) such person submits to the Administrator, at least 90 days before such manufacture or processing, a notice, in accordance with subsection (d), of such person’s intention to
manufacture or process such substance and such person complies with any applicable requirement of, or imposed pursuant to, subsection (b), (e), or (f); and

“(ii) the Administrator—

“(I) conducts a review of the notice;

and

“(II) makes a determination under subparagraph (A), (B), (C), or (D) of paragraph (3) and takes the actions required in association with that determination under such subparagraph within the applicable review period.”; and

(B) by adding at the end the following new paragraphs:

“(3) REVIEW AND DETERMINATION.—Within the applicable review period, subject to section 18, the Administrator shall review such notice and determine—

“(A) that the relevant chemical substance or significant new use presents or will present an unreasonable risk of injury to health or the environment, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed or susceptible
subpopulation identified as relevant by the Administrator under the conditions of use, in which case the Administrator shall take the actions required under subsection (f);

“(B) that—

“(i) the information available to the Administrator is insufficient to permit a reasoned evaluation of the health and environmental effects of the relevant chemical substance or significant new use; or

“(ii)(I) in the absence of sufficient information to permit the Administrator to make such an evaluation, the manufacture, processing, distribution in commerce, use, or disposal of such substance, or any combination of such activities, may present an unreasonable risk of injury to health or the environment, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed or susceptible subpopulation identified as relevant by the Administrator; or

“(II) such substance is or will be produced in substantial quantities, and such substance either enters or may reasonably
be anticipated to enter the environment in substantial quantities or there is or may be significant or substantial human exposure to the substance, in which case the Administrator shall take the actions required under subsection (e);

“(C) that the relevant chemical substance or significant new use is likely not to present an unreasonable risk of injury to health or the environment, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed or susceptible subpopulation identified as relevant by the Administrator under the conditions of use, in which case the submitter of the notice may commence manufacture of the chemical substance or manufacture or processing for a significant new use; or

“(D) that the relevant chemical substance is a low-hazard substance, in which case the submitter of the notice may commence manufacture of the chemical substance or manufacture or processing of the chemical substance for a significant new use.

“(4) FAILURE TO RENDER DETERMINATION.—
“(A) FAILURE TO RENDER DETERMINATION.—If the Administrator fails to make a determination on a notice under paragraph (3) by the end of the applicable review period and the notice has not been withdrawn by the submitter, the Administrator shall refund to the submitter all applicable fees charged to the submitter for review of the notice pursuant to section 26(b), and the Administrator shall not be relieved of any requirement to make such determination.

“(B) LIMITATIONS.—(i) A refund of applicable fees under subparagraph (A) shall not be made if the Administrator certifies that the submitter has not provided information required under subsection (b) or has otherwise unduly delayed the process such that the Administrator is unable to render a determination within the applicable review period.

“(ii) A failure of the Administrator to render a decision shall not be deemed to constitute a withdrawal of the notice.

“(iii) Nothing in this paragraph shall be construed as relieving the Administrator or the
submitter of the notice from any requirement of this section.

“(5) ARTICLE CONSIDERATION.—The Administrator may require notification under this section for the import or processing of a chemical substance as part of an article or category of articles under paragraph (1)(A)(ii) if the Administrator makes an affirmative finding in a rule under paragraph (2) that the reasonable potential for exposure to the chemical substance through the article or category of articles subject to the rule justifies notification.”;

(2) in subsection (b)—

(A) in the subsection heading, by striking “TEST DATA” and inserting “INFORMATION”; 

(B) in paragraph (1)—

(i) in subparagraph (A)—

(I) by striking “test data” and inserting “information”; and

(II) by striking “such data” and inserting “such information”; and

(ii) in subparagraph (B)—

(I) by striking “test data” and inserting “information”;
(II) by striking "subsection (a)(1)(A)" and inserting "subsection (a)(1)(A)(i)"; and

(III) by striking "subsection (a)(1)(B)" and inserting "subsection (a)(1)(A)(ii)";

(C) in paragraph (2)—

(i) in subparagraph (A)—

(I) by striking "test data" in clause (ii) and inserting "information";

(II) by striking "shall" and inserting "may"; and

(III) by striking "data prescribed" and inserting "information prescribed"; and

(ii) in subparagraph (B)—

(I) by striking "Data" and inserting "Information";

(II) by striking "data" both places it appears and inserting "information";

(III) by striking "show" and inserting "shows";
(IV) by striking “subsection (a)(1)(A)” in clause (i) and inserting “subsection (a)(1)(A)(i)”; and

(V) by striking “subsection (a)(1)(B)” in clause (ii) and inserting “subsection (a)(1)(A)(ii)”;

(D) in paragraph (3)—

(i) by striking “Data” and inserting “Information”; and

(ii) by striking “paragraph (1) or (2)” and inserting “paragraph (1) or (2) of this subsection or under subsection (e)”; and

(E) in paragraph (4)—

(i) in subparagraph (A)(i), by inserting “, without consideration of costs or other nonrisk factors” after “health or the environment”; and

(ii) in subparagraph (C), by striking “, except that” and all that follows through “subparagraph (A)”;

(3) in subsection (e)—

(A) in the subsection heading, by striking “NOTICE” and inserting “REVIEW”; and

(B) by striking “before which” and all that follows through “subsection may begin”;
(4) in subsection (d)—

(A) by striking “test data” in paragraph (1)(B) and inserting “information”;

(B) by striking “data” each place it appears in paragraph (1)(C) and paragraph (2) and inserting “information”;

(C) in paragraph (2)(B), by striking “uses or intended uses of such substance” and inserting “uses of such substance identified in the notice”; and

(D) in paragraph (3)—

(i) by striking “for which the notification period prescribed by subsection (a), (b), or (c)” and inserting “for which the applicable review period”; and

(ii) by striking “such notification period” and inserting “such period”;

(5) in subsection (e)—

(A) in paragraph (1)(A)—

(i) in clause (i), by striking “; and” and inserting “; or”;

(ii) in clause (ii)(I), by inserting “without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed subpopulation
identified as relevant by the Administrator under the conditions of use;” after “health or the environment,”; and

(iii) in the matter after clause (ii)(II)—

(I) by striking “may issue a proposed order” and inserting “shall issue an order”;

(II) by striking “notification period applicable to the manufacturing or processing of such substance under subsection (a), (b), (c)” and inserting “applicable review period”; and

(III) by inserting “to the extent necessary to protect against an unreasonable risk of injury to health or the environment, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed or susceptible subpopulation identified as relevant by the Administrator under the conditions of use, and the submitter of the notice may commence manufacture of the chemical substance, or manufac-
nature or processing of the chemical substance for a significant new use, including while any required information is being developed, only in compliance with the order” before the period at the end;

(B) in paragraph (1)(B)—

(i) by striking “A proposed order” and inserting “An order”;

(ii) by striking “notification period applicable to the manufacture or processing of such substance under subsection (a), (b), (c)” and inserting “applicable review period”; and

(iii) by striking “of the proposed order” and inserting “of the order”;

(C) by striking paragraph (1)(C); and

(D) by striking paragraph (2);

(6) in subsection (f)—

(A) in paragraph (1)—

(i) by striking “finds that there is a reasonable basis to conclude that the manufacture, processing, distribution in commerce, use, or disposal of a chemical substance with” and inserting “determines
that a chemical substance or significant new use with”;

(ii) by striking “, or that any combination of such activities,”;

(iii) by striking “before a rule promulgated under section 6 can protect against such risk,” and inserting “, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed subpopulation identified as relevant by the Administrator under the conditions of use,”; and

(iv) by striking “notification period applicable under subsection (a), (b), or (e) to the manufacturing or processing of such substance” and inserting “applicable review period”;

(B) in paragraph (2), the matter following subparagraph (C), by striking “Section 6(d)(2)(B)” and inserting “Section 6(d)(3)(B)”;

(C) in paragraph (3)—

(i) in subparagraph (A)—

(I) by striking “Administrator may” and all that follows through
issue a proposed order to prohibit
the’’ and inserting ‘‘Administrator
may issue an order to prohibit or limit
the’’; and

(II) by striking ‘‘under para-
graph (1)’’ and all that follows
through ‘‘processing of such sub-
stance.’’ and inserting ‘‘under para-
graph (1). Such order shall take effect
on the expiration of the applicable re-
view period.’’;

(ii) by striking subparagraph (B) and
redesignating subparagraph (C) as sub-
paragraph (B);

(iii) in subparagraph (B), as so redes-
ignated—

(I) by striking ‘‘subparagraphs
(B) and (C)’’ and inserting ‘‘subpara-
graph (B)’’;

(II) by striking ‘‘clause (i) of’’;

and

(III) by striking ‘‘; and the provi-
sions of subparagraph (C) of sub-
section (e)(2) shall apply with respect
to an injunction issued under sub-
paragraph (B)”; and
(iv) by striking subparagraph (D);
and
(D) by adding at the end the following:
“(4) TREATMENT OF NONCONFORMING USES.—
Not later than 90 days after taking an action under
paragraph (2) or (3) or issuing an order under sub-
section (e) relating to a chemical substance with re-
spect to which the Administrator has made a deter-
mination under subsection (a)(3)(A) or (B), the Ad-
ministrator shall consider whether to promulgate a
rule pursuant to subsection (a)(2) that identifies as
a significant new use any manufacturing, processing,
use, distribution in commerce, or disposal of the
chemical substance that does not conform to the re-
strictions imposed by the action or order, and, as ap-
plicable, initiate such a rulemaking or publish a
statement describing the reasons of the Adminis-
trator for not initiating such a rulemaking.
“(5) WORKPLACE EXPOSURES.—To the extent
practicable, the Administrator shall consult with the
Assistant Secretary of Labor for Occupational Safe-
ty and Health prior to adopting any prohibition or
other restriction relating to a chemical substance
with respect to which the Administrator has made a
determination under subsection (a)(3)(A) or (B) to
address workplace exposures.”;

(7) by amending subsection (g) to read as fol-

lows:

“(g) STATEMENT ON ADMINISTRATOR FINDING.—If
the Administrator finds in accordance with subsection
(a)(3)(C) that a chemical substance or significant new use
is likely not to present an unreasonable risk of injury to
health or the environment, or in accordance with sub-
section (a)(3)(D) that the chemical substance is a low-haz-
ard substance, then notwithstanding any remaining por-
tion of the applicable review period, the submitter of the
notice may commence manufacture of the chemical sub-
stance or manufacture or processing for the significant
new use, and the Administrator shall make public a state-
ment of the Administrator’s finding. Such a statement
shall be submitted for publication in the Federal Register
as soon as is practicable before the expiration of such pe-
period. Publication of such statement in accordance with the
preceding sentence is not a prerequisite to the manufac-
turing or processing of the substance with respect to which
the statement is to be published.”;

(8) in subsection (h)—
(A) in paragraph (1)(A), by inserting “,
including an unreasonable risk to a potentially
exposed or susceptible subpopulation identified
by the Administrator for the specific conditions
of use identified in the application” after
“health or the environment”;

(B) in paragraph (2), by striking “data”
each place it appears and inserting “informa-
tion”; and

(C) in paragraph (4), by striking “. A rule
promulgated” and all that follows through “sec-
tion 6(c)” and inserting “, including an unrea-
sonable risk to a potentially exposed or suscep-
tible subpopulation identified by the Adminis-
trator under the conditions of use”; and

(9) by amending subsection (i) to read as fol-

laws:

“(i) DEFINITIONS.—(1) For purposes of this section,
the terms ‘manufacture’ and ‘process’ mean manufac-
turing or processing for commercial purposes.

“(2) For purposes of this Act, the term ‘requirement’
as used in this section shall not displace any statutory or
common law.

“(3) For purposes of this section, the term ‘applicable
review period’ means the period starting on the date the
Administrator receives a notice under subsection (a)(1) and ending 90 days after that date, or on such date as is provided for in subsection (b)(1) or (c).”.

SEC. 6. PRIORITIZATION, RISK EVALUATION, AND REGULATION OF CHEMICAL SUBSTANCES AND MIXTURES.

Section 6 of the Toxic Substances Control Act (15 U.S.C. 2605) is amended—

(1) by striking the section heading and inserting “PRIORITIZATION, RISK EVALUATION, AND REGULATION OF CHEMICAL SUBSTANCES AND MIXTURES”;  

(2) in subsection (a)—

(A) by striking “finds that there is a reasonable basis to conclude” and inserting “determines in accordance with subsection (b)(4)(A)”;

(B) by inserting “and subject to section 18, and in accordance with subsection (c)(2),” after “shall by rule”;  

(C) by striking “to protect adequately against such risk using the least burdensome requirements” and inserting “so that the chemical substance or mixture no longer presents such risk”;
(D) by inserting “or otherwise restricting” after “prohibiting” in paragraphs (1)(A) and (2)(A); 

(E) by inserting “minimum” before “warnings” both places it appears in paragraph (3); 

(F) by striking “and monitor or conduct tests” and inserting “or monitor or conduct tests” in paragraph (4); and 

(G) in paragraph (7)—

(i) by striking “such unreasonable risk of injury” and inserting “such determination”; and 

(ii) by striking “such risk of injury” and inserting “such determination”; 

(3) by amending subsection (b) to read as follows:

“(b) RISK EVALUATIONS.—

“(1) PRIORITIZATION FOR RISK EVALUATIONS.—

“(A) ESTABLISHMENT OF PROCESS.—Not later than 1 year after the date of enactment of the Frank R. Launtenberg Chemical Safety for the 21st Century Act, the Administrator shall establish, by rule, a risk-based screening process, including criteria for designating chemical
substances as high-priority substances for risk
evaluations or low-priority substances for which
risk evaluations are not warranted at the time.
The process to designate the priority of chem-
ical substances shall include a consideration of
the hazard and exposure potential of a chemical
substance or a category of chemical substances
(including consideration of persistence and bio-
accumulation, potentially exposed or susceptible
subpopulations and storage near significant
sources of drinking water), the conditions of use
or significant changes in the conditions of use
of the chemical substance, and the volume or
significant changes in the volume of the chem-
ical substance manufactured or processed.

“(B) IDENTIFICATION OF PRIORITIES FOR
RISK EVALUATION.—

“(i) HIGH-PRIORITY SUBSTANCES.—
The Administrator shall designate as a
high-priority substance a chemical sub-
stance that the Administrator concludes,
without consideration of costs or other
nonrisk factors, may present an unreason-
able risk of injury to health or the environ-
ment because of a potential hazard and a
potential route of exposure under the conditions of use, including an unreasonable risk to a potentially exposed or susceptible subpopulation identified as relevant by the Administrator.

“(ii) LOW-PRIORITY SUBSTANCES.—

Except as provided in clause (iii), the Administrator shall designate a chemical substance as a low-priority substance if the Administrator concludes, based on information sufficient to establish, without consideration of costs or other nonrisk factors, that such substance does not meet the standard identified in clause (i) for designating a chemical substance a high-priority substance.

“(iii) LOW-HAZARD SUBSTANCES.—

The Administrator may designate a low-priority substance as a low-hazard substance if the Administrator concludes, based on information sufficient to establish, without consideration of costs or other nonrisk factors or exposure, that the chemical substance poses no or low hazard to health or the environment, including any
hazard to a potentially exposed or susceptible subpopulation identified as relevant by the Administrator.

“(C) INFORMATION REQUEST AND REVIEW AND PROPOSED AND FINAL PRIORITIZATION DESIGNATION.—The rulemaking required in subparagraph (A) shall ensure that the time required to make a priority designation of a chemical substance be no shorter than nine months and no longer than 1 year, and that the process for such designations includes—

“(i) a requirement that the Administrator request interested persons to submit relevant information on a chemical substance that the Administrator has initiated the prioritization process on, before proposing a priority designation for the chemical substance, and provide 90 days for such information to be provided;

“(ii) a requirement that the Administrator publish each proposed designation of a chemical substance as a high- or low-priority substance, along with an identification of the information, analysis, and basis used to make the proposed designations,
and provide 90 days for public comment on each such proposed designation; and

“(iii) a process by which the Administrator may extend the deadline in clause (i) for up to three months in order to receive or evaluate information required to be submitted in accordance with section 4(a)(2)(B), subject to the limitation that if the information available to the Administrator at the end of such an extension remains insufficient to enable the designation of the chemical substance as a low-priority substance, the Administrator shall designate the chemical substance as a high-priority substance.

“(2) INITIAL RISK EVALUATIONS AND SUBSEQUENT DESIGNATIONS OF HIGH- AND LOW-PRIORITY SUBSTANCES.—

“(A) INITIAL RISK EVALUATIONS.—Not later than 180 days after the date of enactment of the Frank R. Launtenberg Chemical Safety for the 21st Century Act, the Administrator shall ensure that risk evaluations are being conducted on at least 10 chemical substances drawn from the 2014 update of the TSCA
Work Plan for Chemical Assessments and shall publish the list of such chemical substances during the 180 day period.

“(B) Additional risk evaluations.—Not later than three and one half years after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, the Administrator shall ensure that risk evaluations are being conducted on at least 20 high-priority substances and that at least 20 chemical substances have been designated as low-priority or low-hazard substances, subject to the limitation that at least 50 percent of all chemical substances on which risk evaluations are being conducted by the Administrator are drawn from the 2014 update of the TSCA Work Plan for Chemical Assessments.

“(C) Continuing designations and risk evaluations.—The Administrator shall continue to designate priority substances and conduct risk evaluations in accordance with this subsection at a pace consistent with the ability of the Administrator to complete risk evaluations in accordance with the deadlines under paragraph (4)(G).
“(D) PREFERENCE.—In designating high-priority substances, the Administrator shall give preference to—

“(i) chemical substances that are listed in the 2014 update of the TSCA Work Plan for Chemical Assessments as having a Persistence and Bioaccumulation Score of 3; and

“(ii) chemical substances that are listed in the 2014 update of the TSCA Work Plan for Chemical Assessments that are known human carcinogens and have high acute and chronic toxicity.

“(E) METALS AND METAL COMPOUNDS.—

In identifying priorities for risk evaluation and conducting risk evaluations of metals and metal compounds, the Administrator shall use the Framework for Metals Risk Assessment of the Office of the Science Advisor, Risk Assessment Forum, and dated March 2007, or a successor document that addresses metals risk assessment and is peer reviewed by the Science Advisory Board.

“(3) INITIATION OF RISK EVALUATIONS; DESIGNATIONS.—
“(A) Risk evaluation initiation.—
Upon designating a chemical substance as a high-priority substance, the Administrator shall initiate a risk evaluation on the substance.

“(B) Revision.—The Administrator may revise the designation of a low-priority substance or a low-hazard substance based on information made available to the Administrator.

“(C) Ongoing designations.—The Administrator shall designate at least one high-priority substance upon the completion of each risk evaluation (other than risk evaluations for chemical substances designated under paragraph (4)(C)(ii)).

“(4) Risk evaluation process and deadlines.—

“(A) In general.—The Administrator shall conduct risk evaluations pursuant to this paragraph to determine whether a chemical substance presents an unreasonable risk of injury to health or the environment, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed or susceptible subpopulation identified as
relevant to the risk evaluation by the Administrator, under the conditions of use.

“(B) ESTABLISHMENT OF PROCESS.—Not later than 1 year after the date of enactment of the Frank R. Launtenberg Chemical Safety for the 21st Century Act, the Administrator shall establish, by rule, a process to conduct risk evaluations in accordance with subparagraph (A).

“(C) REQUIREMENT.—The Administrator shall conduct and publish risk evaluations, in accordance with the rule promulgated under subparagraph (B), for a chemical substance—

“(i) that has been identified under paragraph (2)(A) or designated under paragraph (1)(B)(i); and

“(ii) subject to subparagraph (E), that a manufacturer of the chemical substance has requested, in a form and manner and using the criteria prescribed by the Administrator in the rule promulgated under subparagraph (B), be subjected to a risk evaluation.

“(D) SCOPE.—The Administrator shall, not later than 6 months after the initiation of
a risk evaluation, publish the scope of the risk
evaluation to be conducted, including the haz-
ards, exposures, conditions of use, and the po-
tentially exposed or susceptible subpopulations
the Administrator expects to consider, and, for
each designation of a high-priority substance,
ensure not less than 12 months between the ini-
tiation of the prioritization process for the
chemical substance and the publication of the
scope of the risk evaluation for the chemical
substance, and for risk evaluations conducted
on chemical substances that have been identi-
fied under paragraph (2)(A) or selected under
subparagraph (E)(iv)(II) of this paragraph, en-
sure not less than 3 months before the Admin-
istrator publishes the scope of the risk evalua-
tion.

“(E) LIMITATION AND CRITERIA.—

“(i) PERCENTAGE REQUIREMENTS.—
The Administrator shall ensure that, of the
number of chemical substances that under-
go a risk evaluation under clause (i) of
subparagraph (C), the number of chemical
substances undergoing a risk evaluation
under clause (ii) of subparagraph (C) is—
“(I) not less than 25 percent, if sufficient requests are made under clause (ii) of subparagraph (C); and

“(II) not more than 50 percent.

“(ii) Requested risk evaluations.—Requests for risk evaluations under subparagraph (C)(ii) shall be subject to the payment of fees pursuant to section 26(b), and the Administrator shall not expedite or otherwise provide special treatment to such risk evaluations.

“(iii) Preference.—In deciding whether to grant requests under subparagraph (C)(ii), the Administrator shall give preference to requests for risk evaluations on chemical substances for which the Administrator determines that restrictions imposed by 1 or more States have the potential to have a significant impact on interstate commerce or health or the environment.

“(iv) Exceptions.—(I) Chemical substances for which requests have been granted under subparagraph (C)(ii) and that are not drawn from the 2014 update
of the TSCA Work Plan for Chemical Assessments shall not be subject to section 18(b).

“(II) Requests for risk evaluations on chemical substances which are made under subparagraph (C)(ii) and that are drawn from the 2014 update of the TSCA Work Plan for Chemical Assessments shall be granted at the discretion of the Administrator and not be subject to clause (i)(II).

“(F) REQUIREMENTS.—In conducting a risk evaluation under this subsection, the Administrator shall—

“(i) integrate and assess available information on hazards and exposures for the conditions of use of the chemical substance, including information that is relevant to specific risks of injury to health or the environment and information on potentially exposed or susceptible subpopulations identified as relevant by the Administrator;

“(ii) describe whether aggregate or sentinel exposures to a chemical substance under the conditions of use were considered, and the basis for that consideration;
“(iii) not consider costs or other nonrisk factors;

“(iv) take into account, where relevant, the likely duration, intensity, frequency, and number of exposures under the conditions of use of the chemical substance; and

“(v) describe the weight of the scientific evidence for the identified hazard and exposure.

“(G) DEADLINES.—The Administrator—

“(i) shall complete a risk evaluation for a chemical substance as soon as practicable, but not later than 3 years after the date on which the Administrator initiates the risk evaluation under subparagraph (C); and

“(ii) may extend the deadline for a risk evaluation for not more than 6 months.

“(H) NOTICE AND COMMENT.—The Administrator shall provide no less than 30 days public notice and an opportunity for comment on a draft risk evaluation prior to publishing a final risk evaluation.”;}
(4) by amending subsection (c) to read as fol-

lows:

“(c) PROMULGATION OF SUBSECTION (a) RULES.—

“(1) DEADLINES.—If the Administrator deter-

mines that a chemical substance presents an unrea-

sonable risk of injury to health or the environment

in accordance with subsection (b)(4)(A), the Admin-

istrator—

“(A) shall propose in the Federal Register

a rule under subsection (a) for the chemical

substance not later than 1 year after the date

on which the final risk evaluation regarding the

chemical substance is published;

“(B) shall publish in the Federal Register

a final rule not later than 2 years after the date

on which the final risk evaluation regarding the

chemical substance is published; and

“(C) may extend the deadlines under this

paragraph for not more than two years, subject

to the condition that the aggregate length of ex-
tensions under this subparagraph and sub-

section (b)(4)(G)(ii) does not exceed two years,

and subject to the limitation that the Adminis-

trator may not extend a deadline for the publi-
cation of a proposed or final rule regarding a
chemical substance drawn from the 2014 update of the TSCA Work Plan for Chemical Assessments or a chemical substance that, with respect to persistence and bioaccumulation, scores high for 1 and either high or moderate for the other, pursuant to the TSCA Work Plan Chemicals Methods Document published by the Administrator in February 2012 (or a successor scoring system), without adequate public justification that demonstrates, following a review of the information reasonably available to the Administrator, that the Administrator cannot complete the proposed or final rule without additional information regarding the chemical substance.

“(2) REQUIREMENTS FOR RULE.—

“(A) STATEMENT OF EFFECTS.—In proposing and promulgating a rule under subsection (a) with respect to a chemical substance or mixture, the Administrator shall consider and publish a statement based on reasonably available information with respect to—

“(i) the effects of the chemical substance or mixture on health and the mag-
mitude of the exposure of human beings to
the chemical substance or mixture;

“(ii) the effects of the chemical sub-
stance or mixture on the environment and
the magnitude of the exposure of the envi-
ronment to such substance or mixture;

“(iii) the benefits of the chemical sub-
stance or mixture for various uses; and

“(iv) the reasonably ascertainable eco-
nomic consequences of the rule, includ-
ing consideration of—

“(I) the likely effect of the rule
on the national economy, small busi-
ness, technological innovation, the en-
vironment, and public health;

“(II) the costs and benefits of
the proposed and final regulatory ac-
tion and of the 1 or more primary al-
ternative regulatory actions considered
by the Administrator; and

“(III) the cost effectiveness of
the proposed regulatory action and of
the 1 or more primary alternative reg-
ulatory actions considered by the Ad-
ministrator.
“(B) Selecting requirements.—In selecting among prohibitions and other restrictions, the Administrator shall factor in, to the extent practicable, the considerations under subparagraph (A) in accordance with subsection (a).

“(C) Consideration of alternatives.—Based on the information published under subparagraph (A), in deciding whether to prohibit or restrict in a manner that substantially prevents a specific condition of use of a chemical substance or mixture, and in setting an appropriate transition period for such action, the Administrator shall consider, to the extent practicable, whether technically and economically feasible alternatives that benefit health or the environment, compared to the use so proposed to be prohibited or restricted, will be reasonably available as a substitute when the proposed prohibition or other restriction takes effect.

“(D) Replacement parts.—

“(i) In general.—The Administrator shall exempt replacement parts for complex durable goods and complex consumer goods
that are designed prior to the date of publication in the Federal Register of the rule under subsection (a), unless the Administrator finds that such replacement parts contribute significantly to the risk, identified in a risk evaluation conducted under subsection (b)(4)(A), to the general population or to an identified potentially exposed or susceptible subpopulation.

“(ii) DEFINITIONS.—In this subparagraph—

“(I) the term ‘complex consumer goods’ means electronic or mechanical devices composed of multiple manufactured components, with an intended useful life of 3 or more years, where the product is typically not consumed, destroyed, or discarded after a single use, and the components of which would be impracticable to redesign or replace; and

“(II) the term ‘complex durable goods’ means manufactured goods composed of 100 or more manufactured components, with an intended
useful life of 5 or more years, where
the product is typically not consumed,
destroyed, or discarded after a single
use.

“(E) ARTICLES.—In selecting among pro-
hibitions and other restrictions, the Adminis-
trator shall apply such prohibitions or other re-
strictions to an article or category of articles
containing the chemical substance or mixture
only to the extent necessary to address the
identified risks from exposure to the chemical
substance or mixture from the article or cat-
egory of articles so that the substance or mix-
ture does not present an unreasonable risk of
injury to health or the environment identified in
the risk evaluation conducted in accordance
with subsection (b)(4)(A).

“(3) PROCEDURES.—When prescribing a rule
under subsection (a) the Administrator shall proceed
in accordance with section 553 of title 5, United
States Code (without regard to any reference in such
section to sections 556 and 557 of such title), and
shall also—
“(A) publish a notice of proposed rule-making stating with particularity the reason for
the proposed rule;

“(B) allow interested persons to submit
written data, views, and arguments, and make
all such submissions publicly available;

“(C) promulgate a final rule based on the
matter in the rulemaking record; and

“(D) make and publish with the rule the
determination described in subsection (a).”;

(5) in subsection (d)—

(A) by redesignating paragraph (2) as
paragraph (3);

(B) by striking paragraph (1) and inserting
the following:

“(1) IN GENERAL.—In any rule under sub-
section (a), the Administrator shall—

“(A) specify the date on which it shall take
effect, which date shall be as soon as prac-
ticable;

“(B) except as provided in subparagraphs
(C) and (D), specify mandatory compliance
dates for all of the requirements under a rule
under subsection (a), which shall be as soon as
practicable, but not later than 5 years after the
date of promulgation of the rule, except in a case of a use exempted under subsection (g);

“(C) specify mandatory compliance dates for the start of ban or phase-out requirements under a rule under subsection (a), which shall be as soon as practicable, but not later than 5 years after the date of promulgation of the rule, except in the case of a use exempted under subsection (g);

“(D) specify mandatory compliance dates for full implementation of ban or phase-out requirements under a rule under subsection (a), which shall be as soon as practicable; and

“(E) provide for a reasonable transition period.

“(2) VARIABILITY.—As determined by the Administrator, the compliance dates established under paragraph (1) may vary for different affected persons.”; and

(C) in paragraph (3), as so redesignated by subparagraph (A) of this paragraph—

(i) in subparagraph (A)—

(I) by striking “upon its publication” and all that follows through “respecting such rule if” and inserting “,
and compliance with the proposed require-
ments to be mandatory, upon publica-
tion in the Federal Register of the proposed rule and until the com-
pliance dates applicable to such require-
ments in a final rule promul-
gated under section 6(a) or until the Administrator revokes such proposed rule, in accordance with subparagraph (B), if”; and

(II) in clause (i)(I), by inserting “without consideration of costs or other non-risk factors” after “effective date”; and

(ii) in subparagraph (B), by striking “, provide reasonable opportunity” and all that follows through the period at the end and inserting “in accordance with sub-
section (c), and either promulgate such rule (as proposed or with modifications) or revoke it.”;

(6) in subsection (e)(4), by striking “para-
graphs (2), (3), and (4)” and inserting “paragraph (3)”;} and
(7) by adding at the end the following new subsections:

“(g) Exemptions.—

“(1) Criteria for exemption.—The Administrator may, as part of a rule promulgated under subsection (a), or in a separate rule, grant an exemption from a requirement of a subsection (a) rule for a specific condition of use of a chemical substance or mixture, if the Administrator finds that—

“(A) the specific condition of use is a critical or essential use for which no technically and economically feasible safer alternative is available, taking into consideration hazard and exposure;

“(B) compliance with the requirement, as applied with respect to the specific condition of use, would significantly disrupt the national economy, national security, or critical infrastructure; or

“(C) the specific condition of use of the chemical substance or mixture, as compared to reasonably available alternatives, provides a substantial benefit to health, the environment, or public safety.
“(2) EXEMPTION ANALYSIS AND STATEMENT.—
In proposing an exemption under this subsection, the Administrator shall analyze the need for the exemption, and shall make public the analysis and a statement describing how the analysis was taken into account.

“(3) PERIOD OF EXEMPTION.—The Administrator shall establish, as part of a rule under this subsection, a time limit on any exemption for a time to be determined by the Administrator as reasonable on a case-by-case basis, and, by rule, may extend, modify, or eliminate an exemption if the Administrator determines, on the basis of reasonably available information and after adequate public justification, the exemption warrants extension or modification or is no longer necessary.

“(4) CONDITIONS.—As part of a rule promulgated under this subsection, the Administrator shall include conditions, including reasonable record-keeping, monitoring, and reporting requirements, to the extent that the Administrator determines the conditions are necessary to protect health and the environment while achieving the purposes of the exemption.
“(h) CHEMICALS THAT ARE PERSISTENT, BIO-
ACCUMULATIVE, AND TOXIC.—

“(1) EXPEDITED ACTION.—Not later than 3
years after the date of enactment of the Frank R.
Lautenberg Chemical Safety for the 21st Century
Act, the Administrator shall propose rules under
subsection (a) with respect to chemical substances
identified in the 2014 update of the TSCA Work
Plan for Chemical Assessments—

“(A) that the Administrator has a reason-
able basis to conclude are toxic and that with
respect to persistence and bioaccumulation
score high for one and either high or moderate
for the other, pursuant to the TSCA Work Plan
Chemicals Methods Document published by the
Administrator in February 2012 (or a successor
scoring system), and are not a metal or a metal
compound, and for which the Administrator has
not completed a Work Plan Problem Formula-
tion, initiated a review under section 5, or en-
tered into a consent agreement under section 4,
prior to the date of enactment of the Frank R.
Lautenberg Chemical Safety for the 21st Cen-
tury Act; and
“(B) exposure to which under the conditions of use is likely to the general population or to a potentially exposed or susceptible subpopulation identified by the Administrator, or the environment, on the basis of an exposure and use assessment conducted by the Administrator.

“(2) NO RISK EVALUATION REQUIRED.—The Administrator shall not be required to conduct risk evaluations on chemical substances that are subject to paragraph (1).

“(3) FINAL RULE.—Not later than 18 months after proposing a rule pursuant to paragraph (1), the Administrator shall promulgate a final rule under subsection (a).

“(4) SELECTING RESTRICTIONS.—In selecting among prohibitions and other restrictions promulgated in a rule under subsection (a) pursuant to paragraph (1), the Administrator shall address the risks of injury to health or the environment that the Administrator determines are presented by the chemical substance and shall reduce exposure to the substance to the extent practicable.

“(5) RELATIONSHIP TO SUBSECTION (b).—If, at any time prior to the date that is 90 days after
the date of enactment of the Frank R. Launtenberg
Chemical Safety for the 21st Century Act, the Ad-
ministrator makes a designation under subsection
(b)(1)(B)(i), or receives a request under subsection
(b)(4)(C)(ii) that meets the criteria prescribed by
the Administrator in the rule promulgated under
subsection (b)(4)(B), such chemical substance shall
not be subject to this subsection, except that in se-
lecting among prohibitions and other restrictions
promulgated in a rule pursuant to subsection (a),
the Administrator shall both ensure that the chem-
ical substance meets the rulemaking standard under
subsection (a) and reduce exposure to the substance
to the extent practicable.
“(i) FINAL AGENCY ACTION.—Under this section
and subject to section 18—
“(1) a determination by the Administrator
under subsection (b)(4)(A) that a chemical sub-
stance does not present an unreasonable risk of in-
jury to health or the environment shall be issued by
order and considered to be a final agency action, ef-
fective beginning on the date of issuance of the
order; and
“(2) a final rule promulgated under subsection
(a), including the associated determination by the
Administrator under subsection (b)(4)(A) that a chemical substance presents an unreasonable risk of injury to health or the environment, shall be considered to be a final agency action, effective beginning on the date of promulgation of the final rule.

“(j) DEFINITION.—For the purposes of this Act, the term ‘requirement’ as used in this section shall not displace statutory or common law.”.

SEC. 7. IMMINENT HAZARDS.

Section 7 of the Toxic Substances Control Act (15 U.S.C. 2606) is amended—

(1) in subsection (b)(1), by inserting “(as identified by the Administrator without consideration of costs or other nonrisk factors)” after “from the unreasonable risk”; and

(2) in subsection (f), by inserting “, without consideration of costs or other nonrisk factors” after “widespread injury to health or the environment”.

SEC. 8. REPORTING AND RETENTION OF INFORMATION.

(a) IN GENERAL.—Section 8 of the Toxic Substances Control Act (15 U.S.C. 2607) is amended—

(1) in subsection (a)—

(A) in paragraph (2), by striking the matter that follows subparagraph (G);
(B) in paragraph (3), by adding at the end the following:

“(C) Not later than 180 days after the date of enactment of the Frank R. Launtenberg Chemical Safety for the 21st Century Act, and not less frequently than once every 10 years thereafter, the Administrator, after consultation with the Administrator of the Small Business Administration, shall—

“(i) review the adequacy of the standards prescribed under subparagraph (B); and

“(ii) after providing public notice and an opportunity for comment, make a determination as to whether revision of the standards is warranted.”;

and

(C) by adding at the end the following:

“(4) CONTENTS.—The rules promulgated pursuant to paragraph (1)—

“(A) may impose differing reporting and recordkeeping requirements on manufacturers and processors; and

“(B) shall include the level of detail necessary to be reported, including the manner by which use and exposure information may be reported.
“(5) ADMINISTRATION.—In carrying out this section, the Administrator shall, to the extent feasible—

“(A) not require reporting which is unnecessary or duplicative;

“(B) minimize the cost of compliance with this section and the rules issued thereunder on small manufacturers and processors; and

“(C) apply any reporting obligations to those persons likely to have information relevant to the effective implementation of this title.

“(6) NEGOTIATED RULEMAKING.—(A) The Administrator shall enter into a negotiated rulemaking pursuant to subchapter III of chapter 5 of title 5, United States Code, to develop and publish, not later than 3 years after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, a proposed rule providing for limiting the reporting requirements, under this subsection, for manufacturers of any inorganic byproducts, when such byproducts, whether by the byproduct manufacturer or by any other person, are subsequently recycled, reused, or reprocessed.
“(B) Not later than 3 and one-half years after such date of enactment, the Administrator shall publish a final rule resulting from such negotiated rule-making.”; and

(2) in subsection (b), by adding at the end the following:

“(3) NOMENCLATURE.—

“(A) IN GENERAL.—In carrying out paragraph (1), the Administrator shall—

“(i) maintain the use of Class 2 nomenclature in use on the date of enactment of the Frank R. Launtenberg Chemical Safety for the 21st Century Act;

“(ii) maintain the use of the Soap and Detergent Association Nomenclature System, published in March 1978 by the Administrator in section 1 of addendum III of the document entitled ‘Candidate List of Chemical Substances’, and further described in the appendix A of volume I of the 1985 edition of the Toxic Substances Control Act Substances Inventory (EPA Document No. EPA–560/7–85–002a); and

“(iii) treat the individual members of the categories of chemical substances iden-
tified by the Administrator as statutory mixtures, as defined in Inventory descriptions established by the Administrator, as being included on the list established under paragraph (1).

“(B) MULTIPLE NOMENCLATURE LISTINGS.—If a manufacturer or processor demonstrates to the Administrator that a chemical substance appears multiple times on the list published under paragraph (1) under different CAS numbers, the Administrator may recognize the multiple listings as a single chemical substance.

“(C) RELATIONSHIP TO SECTION 5.—Nothing in subparagraph (B), nor any action of the Administrator pursuant to subparagraph (B), shall be construed as a basis to conclude that any chemical substance is not a new chemical substance.

“(4) CHEMICAL SUBSTANCES IN COMMERCE.—

“(A) RULES.—

“(i) IN GENERAL.—Not later than 1 year after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, the Administrator,
by rule, shall require manufacturers, and
may require processors, subject to the limi-
tations under subsection (a)(5)(A), to no-
tify the Administrator, by not later than
180 days after the date on which the final
rule is published in the Federal Register,
of each chemical substance on the list pub-
ished under paragraph (1) that the manu-
facturer or processor, as applicable, has
manufactured or processed for a non-
exempt commercial purpose during the 10-
year period ending on the day before the
date of enactment of the Frank R. Lauten-
berg Chemical Safety for the 21st Century
Act.

“(ii) Active substances.—The Ad-
ministrator shall designate chemical sub-
stances for which notices are received
under clause (i) to be active substances on
the list published under paragraph (1).

“(iii) Inactive substances.—The
Administrator shall designate chemical
substances for which no notices are re-
ceived under clause (i) to be inactive sub-
stances on the list published under paragraph (1).

“(iv) LIMITATION.—No chemical substance on the list published under paragraph (1) shall be removed from such list by reason of the implementation of this subparagraph, or be subject to section 5(a)(1)(A)(i) by reason of a change to active status under paragraph (5)(B).

“(B) CONFIDENTIAL CHEMICAL SUBSTANCES.—In promulgating a rule under subparagraph (A), the Administrator shall—

“(i) maintain the list under paragraph (1), which shall include a confidential portion and a nonconfidential portion consistent with this section and section 14;

“(ii) require any manufacturer or processor of a chemical substance on the confidential portion of the list published under paragraph (1) that seeks to maintain an existing claim for protection against disclosure of the specific chemical identity of the chemical substance as confidential pursuant to section 14 to submit
a notice under subparagraph (A) that includes such request;

“(iii) require the substantiation of those claims pursuant to section 14 and in accordance with the review plan described in subparagraph (C); and

“(iv) move any active chemical substance for which no request was received to maintain an existing claim for protection against disclosure of the specific chemical identity of the chemical substance as confidential from the confidential portion of the list published under paragraph (1) to the nonconfidential portion of that list.

“(C) REVIEW PLAN.—Not later than 1 year after the date on which the Administrator compiles the initial list of active substances pursuant to subparagraph (A), the Administrator shall promulgate a rule that establishes a plan to review all claims to protect the specific chemical identities of chemical substances on the confidential portion of the list published under paragraph (1) that are asserted pursuant to subparagraph (B).
“(D) Requirements of Review Plan.—

In establishing the review plan under subparagraph (C), the Administrator shall—

“(i) require, at a time specified by the Administrator, all manufacturers or processors asserting claims under subparagraph (B) to substantiate the claim, in accordance with section 14, unless the manufacturer or processor has substantiated the claim in a submission made to the Administrator during the 5-year period ending on the last day of the time period specified by the Administrator; and

“(ii) in accordance with section 14—

“(I) review each substantiation—

“(aa) submitted pursuant to clause (i) to determine if the claim qualifies for protection from disclosure; and

“(bb) submitted previously by a manufacturer or processor and relied on in lieu of the substantiation required pursuant to clause (i), if the substantiation has not been previously reviewed
by the Administrator, to determine if the claim warrants protection from disclosure;

“(II) approve, approve in part and deny in part, or deny each claim; and

“(III) except as provided in this section and section 14, protect from disclosure information for which the Administrator approves such a claim for a period of 10 years, unless, prior to the expiration of the period—

“(aa) the person notifies the Administrator that the person is withdrawing the claim, in which case the Administrator shall not protect the information from disclosure; or

“(bb) the Administrator otherwise becomes aware that the information does not qualify for protection from disclosure, in which case the Administrator shall take the actions described in section 14(g)(2).
“(E) TIMELINE FOR COMPLETION OF REVIEWS.—

“(i) IN GENERAL.—The Administrator shall implement the review plan so as to complete reviews of all claims specified in subparagraph (C) not later than 5 years after the date on which the Administrator compiles the initial list of active substances pursuant to subparagraph (A).

“(ii) CONSIDERATIONS.—

“(I) IN GENERAL.—The Administrator may extend the deadline for completion of the reviews for not more than 2 additional years, after an adequate public justification, if the Administrator determines that the extension is necessary based on the number of claims needing review and the available resources.

“(II) ANNUAL REVIEW GOAL AND RESULTS.—At the beginning of each year, the Administrator shall publish an annual goal for reviews and the number of reviews completed in the prior year.
“(5) ACTIVE AND INACTIVE SUBSTANCES.—

“(A) IN GENERAL.—The Administrator shall keep designations of active substances and inactive substances on the list published under paragraph (1) current.

“(B) CHANGE TO ACTIVE STATUS.—

“(i) IN GENERAL.—Any person that intends to manufacture or process for a nonexempt commercial purpose a chemical substance that is designated as an inactive substance shall notify the Administrator before the date on which the inactive substance is manufactured or processed.

“(ii) CONFIDENTIAL CHEMICAL IDENTITY.—If a person submitting a notice under clause (i) for an inactive substance on the confidential portion of the list published under paragraph (1) seeks to maintain an existing claim for protection against disclosure of the specific chemical identity of the inactive substance as confidential, the person shall, consistent with the requirements of section 14—

“(I) in the notice submitted under clause (i), assert the claim; and
“(II) by not later than 30 days after providing the notice under clause (i), substantiate the claim.

“(iii) ACTIVE STATUS.—On receiving a notification under clause (i), the Administrator shall—

“(I) designate the applicable chemical substance as an active substance;

“(II) pursuant to section 14, promptly review any claim and associated substantiation submitted pursuant to clause (ii) for protection against disclosure of the specific chemical identity of the chemical substance and approve, approve in part and deny in part, or deny the claim;

“(III) except as provided in this section and section 14, protect from disclosure the specific chemical identity of the chemical substance for which the Administrator approves a claim under subclause (II) for a period of 10 years, unless, prior to the expiration of the period—
“(aa) the person notifies the Administrator that the person is withdrawing the claim, in which case the Administrator shall not protect the information from disclosure; or

“(bb) the Administrator otherwise becomes aware that the information does not qualify for protection from disclosure, in which case the Administrator shall take the actions described in section 14(g)(2); and

“(IV) pursuant to section 6(b), review the priority of the chemical substance as the Administrator determines to be necessary.

“(C) CATEGORY STATUS.—The list of inactive substances shall not be considered to be a category for purposes of section 26(c).

“(6) INTERIM LIST OF ACTIVE SUBSTANCES.—Prior to the promulgation of the rule required under paragraph (4)(A), the Administrator shall designate the chemical substances reported under part 711 of title 40, Code of Federal Regulations (as in effect on
the date of enactment of the Frank R. Lautenberg
Chemical Safety for the 21st Century Act), during
the reporting period that most closely preceded the
date of enactment of the Frank R. Lautenberg
Chemical Safety for the 21st Century Act, as the in-
terim list of active substances for the purposes of
section 6(b).

“(7) PUBLIC INFORMATION.—Subject to this
subsection and section 14, the Administrator shall
make available to the public—

“(A) each specific chemical identity on the
nonconfidential portion of the list published
under paragraph (1) along with the Administra-
tor’s designation of the chemical substance as
an active or inactive substance;

“(B) the unique identifier assigned under
section 14, accession number, generic name,
and, if applicable, premanufacture notice case
number for each chemical substance on the con-
fidential portion of the list published under
paragraph (1) for which a claim of confiden-
tiality was received; and

“(C) the specific chemical identity of any
active substance for which—
“(i) a claim for protection against disclosure of the specific chemical identity of the active substance was not asserted, as required under this subsection or section 14;

“(ii) all claims for protection against disclosure of the specific chemical identity of the active substance have been denied by the Administrator; or

“(iii) the time period for protection against disclosure of the specific chemical identity of the active substance has expired.

“(8) LIMITATION.—No person may assert a new claim under this subsection or section 14 for protection from disclosure of a specific chemical identity of any active or inactive substance for which a notice is received under paragraph (4)(A)(i) or (5)(B)(i) that is not on the confidential portion of the list published under paragraph (1).

“(9) CERTIFICATION.—Under the rules promulgated under this subsection, manufacturers and processors, as applicable, shall be required—

“(A) to certify that each notice or substantiation the manufacturer or processor submits
complies with the requirements of the rule, and
that any confidentiality claims are true and cor-
correct; and

“(B) to retain a record documenting com-
pliance with the rule and supporting confiden-
tiality claims for a period of 5 years beginning
on the last day of the submission period.”.

(b) MERCURY INVENTORY.—Section 8(b) of the
Toxic Substances Control Act (15 U.S.C. 2607(b)) (as
amended by subsection (a)) is further amended by adding
at the end the following:

“(10) MERCURY.—

“(A) DEFINITION OF MERCURY.—In this
paragraph, notwithstanding section 3(2)(B), the
term ‘mercury’ means—

“(i) elemental mercury; and

“(ii) a mercury compound.

“(B) PUBLICATION.—Not later than April
1, 2017, and every 3 years thereafter, the Ad-
ministrator shall carry out and publish in the
Federal Register an inventory of mercury sup-
ply, use, and trade in the United States.

“(C) PROCESS.—In carrying out the inven-
tory under subparagraph (B), the Adminis-
trator shall—
“(i) identify any manufacturing processes or products that intentionally add mercury; and

“(ii) recommend actions, including proposed revisions of Federal law or regulations, to achieve further reductions in mercury use.

“(D) REPORTING.—

“(i) IN GENERAL.—To assist in the preparation of the inventory under subparagraph (B), any person who manufactures mercury or mercury-added products or otherwise intentionally uses mercury in a manufacturing process shall make periodic reports to the Administrator, at such time and including such information as the Administrator shall determine by rule promulgated not later than 2 years after the date of enactment of this paragraph.

“(ii) COORDINATION.—To avoid duplication, the Administrator shall coordinate the reporting under this subparagraph with the Interstate Mercury Education and Reduction Clearinghouse.
“(iii) EXEMPTION.—Clause (i) shall not apply to a person engaged in the generation, handling, or management of mercury-containing waste, unless that person manufactures or recovers mercury in the management of that waste.”

SEC. 9. RELATIONSHIP TO OTHER FEDERAL LAWS.

Section 9 of the Toxic Substances Control Act (15 U.S.C. 2608) is amended—

(1) in subsection (a)—

(A) in paragraph (1)—

(i) by striking “has reasonable basis to conclude” and inserting “determines”;

and

(ii) by inserting “, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed or susceptible subpopulation identified as relevant by the Administrator, under the conditions of use,” after “or the environment”;

(B) in paragraph (2)—

(i) in subparagraph (A), by inserting “, within the time period specified by the
Administrator in the report,” after “issues
an order”; and

(ii) in subparagraph (B), by inserting
“responds within the time period specified
by the Administrator in the report and”
before “initiates, within 90”; 

(C) by redesignating paragraph (3) as
paragraph (6); and

(D) by inserting after paragraph (2) the
following:

“(3) The Administrator shall take the actions de-
scribed in paragraph (4) if the Administrator makes a re-
port under paragraph (1) with respect to a chemical sub-
stance or mixture and the agency to which the report was
made does not—

“(A) issue the order described in paragraph
(2)(A) within the time period specified by the Ad-
ministrator in the report; or

“(B)(i) respond under paragraph (1) within the
timeframe specified by the Administrator in the re-
port; and

“(ii) initiate action within 90 days of publica-
tion in the Federal Register of the response de-
scribed in clause (i).
“(4) If an agency to which a report is submitted under paragraph (1) does not take the actions described in subparagraph (A) or (B) of paragraph (3), the Administrator shall—

“(A) initiate or complete appropriate action under section 6; or

“(B) take any action authorized or required under section 7, as applicable.

“(5) This subsection shall not relieve the Administrator of any obligation to take any appropriate action under section 6(a) or 7 to address risks from the manufacture, processing, distribution in commerce, use, or disposal of a chemical substance or mixture, or any combination of those activities, that are not identified in a report issued by the Administrator under paragraph (1).”;

(2) in subsection (b)—

(A) by striking “The Administrator shall coordinate” and inserting “(1) The Administrator shall coordinate”; and

(B) by adding at the end the following:

“(2) In making a determination under paragraph (1) that it is in the public interest for the Administrator to take an action under this title with respect to a chemical substance or mixture rather than under another law administered in whole or in part by the Administrator, the
Administrator shall consider, based on information reasonably available to the Administrator, all relevant aspects of the risk described in paragraph (1) and a comparison of the estimated costs and efficiencies of the action to be taken under this title and an action to be taken under such other law to protect against such risk.”; and

(3) by adding at the end the following:

“(e) EXPOSURE INFORMATION.—In addition to the requirements of subsection (a), if the Administrator obtains information related to exposures or releases of a chemical substance or mixture that may be prevented or reduced under another Federal law, including a law not administered by the Administrator, the Administrator shall make such information available to the relevant Federal agency or office of the Environmental Protection Agency.”.

SEC. 10. EXPORTS OF ELEMENTAL MERCURY.

(a) Prohibition on Export of Certain Mercury Compounds.—Section 12(c) of the Toxic Substances Control Act (15 U.S.C. 2611(c)) is amended—

(1) in the subsection heading, by inserting “AND MERCURY COMPOUNDS” after “MERCURY”;

and

(2) by adding at the end the following:
“(7) Prohibition on export of certain mercury compounds.—

“(A) In general.—Effective January 1, 2020, the export of the following mercury compounds is prohibited:

“(i) Mercury (I) chloride or calomel.

“(ii) Mercury (II) oxide.

“(iii) Mercury (II) sulfate.

“(iv) Mercury (II) nitrate.

“(v) Cinnabar or mercury sulphide.

“(vi) Any mercury compound that the Administrator adds to the list published under subparagraph (B) by rule, on determining that exporting that mercury compound for the purpose of regenerating elemental mercury is technically feasible.

“(B) Publication.—Not later than 90 days after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, and as appropriate thereafter, the Administrator shall publish in the Federal Register a list of the mercury compounds that are prohibited from export under this paragraph.
“(C) Petition.—Any person may petition the Administrator to add a mercury compound to the list published under subparagraph (B).

“(D) Environmentally sound disposal.—This paragraph does not prohibit the export of mercury compounds on the list published under subparagraph (B) to member countries of the Organization for Economic Cooperation and Development for environmentally sound disposal, on the condition that no mercury or mercury compounds so exported are to be recovered, recycled, or reclaimed for use, or directly reused, after such export.

“(E) Report.—Not later than 5 years after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, the Administrator shall evaluate any exports of mercury compounds on the list published under subparagraph (B) for disposal that occurred after such date of enactment and shall submit to Congress a report that—

“(i) describes volumes and sources of mercury compounds on the list published under subparagraph (B) exported for disposal;
“(ii) identifies receiving countries of such exports;

“(iii) describes methods of disposal used after such export;

“(iv) identifies issues, if any, presented by the export of mercury compounds on the list published under subparagraph (B);

“(v) includes an evaluation of management options in the United States for mercury compounds on the list published under subparagraph (B), if any, that are commercially available and comparable in cost and efficacy to methods being utilized in such receiving countries; and

“(vi) makes a recommendation regarding whether Congress should further limit or prohibit the export of mercury compounds on the list published under subparagraph (B) for disposal.

“(F) EFFECT ON OTHER LAW.—Nothing in this paragraph shall be construed to affect the authority of the Administrator under the Solid Waste Disposal Act (42 U.S.C. 6901 et seq.).”.
(b) **Temporary Generator Accumulation**.—Section 5 of the Mercury Export Ban Act of 2008 (42 U.S.C. 6939f) is amended—

1. in subsection (a)(2), by striking “2013” and inserting “2019”;

2. in subsection (b)—
   
   (A) in paragraph (1)—
   
   (i) by redesignating subparagraphs (A), (B), and (C), as clauses (i), (ii), and (iii), respectively and indenting appropriately;

   (ii) in the first sentence, by striking “After consultation” and inserting the following:

   “(A) ASSESSMENT AND COLLECTION.—

   After consultation”;

   (iii) in the second sentence, by striking “The amount of such fees” and inserting the following:

   “(B) AMOUNT.—The amount of the fees described in subparagraph (A)”;

   (iv) in subparagraph (B) (as so designated)—

   (I) in clause (i) (as so redesignated), by striking “publicly avail-
able not later than October 1, 2012’’

and inserting “publicly available not
later than October 1, 2018’’;

(II) in clause (ii) (as so redesign-
ated), by striking “and”;

(III) in clause (iii) (as so redesign-
ated), by striking the period at the
end and inserting ‘‘, subject to clause
(iv); and’’; and

(IV) by adding at the end the fol-
lowing:

“(iv) for generators temporarily accu-
mulating elemental mercury in a facility
subject to subparagraphs (B) and (D)(iv)
of subsection (g)(2) if the facility des-
ignated in subsection (a) is not operational
by January 1, 2019, shall be adjusted to
subtract the cost of the temporary accumu-
lation during the period in which the facil-
ity designated under subsection (a) is not
operational.”; and

(v) by adding at the end the following:

“(C) CONVEYANCE OF TITLE AND PERMIT-
TING.—If the facility designated in subsection
(a) is not operational by January 1, 2020, the Secretary—

“(i) shall immediately accept the conveyance of title to all elemental mercury that has accumulated in facilities in accordance with subsection (g)(2)(D), before January 1, 2020, and deliver the accumulated mercury to the facility designated under subsection (a) on the date on which the facility becomes operational;

“(ii) shall pay any applicable Federal permitting costs, including the costs for permits issued under section 3005(c) of the Solid Waste Disposal Act (42 U.S.C. 6925(c)); and

“(iii) shall store, or pay the cost of storage of, until the time at which a facility designated in subsection (a) is operational, accumulated mercury to which the Secretary has title under this subparagraph in a facility that has been issued a permit under section 3005(c) of the Solid Waste Disposal Act (42 U.S.C. 6925(c)).”; and
(B) in paragraph (2), in the first sentence,
by striking “paragraph (1)(C)” and inserting
“paragraph (1)(B)(iii)”; and
(3) in subsection (g)(2)—
(A) in the undesignated material at the end, by striking “This subparagraph” and insert-
ing the following:
“(C) Subparagraph (B)”;
(B) in subparagraph (C) (as designated by subparagraph (A)), by inserting “of that sub-
paragraph” before the period at the end; and
(C) by adding at the end the following:
“(D) A generator producing elemental mercury incidentally from the beneficiation or processing of ore or related pollution control ac-
tivities may accumulate the mercury produced onsite that is destined for a facility designated by the Secretary under subsection (a) for more than 90 days without a permit issued under section 3005(e) of the Solid Waste Disposal Act (42 U.S.C. 6925(c)), and shall not be subject to the storage prohibition of section 3004(j) of that Act (42 U.S.C. 6924(j)), if—
“(i) the Secretary is unable to accept the mercury at a facility designated by the
Secretary under subsection (a) for reasons beyond the control of the generator;

“(ii) the generator certifies in writing to the Secretary that the generator will ship the mercury to a designated facility when the Secretary is able to accept the mercury;

“(iii) the generator certifies in writing to the Secretary that the generator is storing only mercury the generator has produced or recovered onsite and will not sell, or otherwise place into commerce, the mercury; and

“(iv) the generator has obtained an identification number under section 262.12 of title 40, Code of Federal Regulations, and complies with the requirements described in paragraphs (1) through (4) of section 262.34(a) of title 40, Code of Federal Regulations (as in effect on the date of enactment of this subparagraph).

“(E) MANAGEMENT STANDARDS FOR TEMPORARY STORAGE.—Not later than January 1, 2017, the Secretary, after consultation with the Administrator of the Environmental Protection
Agency and State agencies in affected States, shall develop and make available guidance that establishes procedures and standards for the management and short-term storage of elemental mercury at a generator covered under subparagraph (D), including requirements to ensure appropriate use of flasks or other suitable containers. Such procedures and standards shall be protective of health and the environment and shall ensure that the elemental mercury is stored in a safe, secure, and effective manner. A generator may accumulate mercury in accordance with subparagraph (D) immediately upon enactment of this subparagraph, and notwithstanding that guidance called for by this paragraph has not been developed or made available.”.

(c) INTERIM STATUS.—Section 5(d)(1) of the Mercury Export Ban Act of 2008 (42 U.S.C. 6939f(d)(1)) is amended—

(1) in the fourth sentence, by striking “in existence on or before January 1, 2013,”; and

(2) in the last sentence, by striking “January 1, 2015” and inserting “January 1, 2020”.

SEC. 11. CONFIDENTIAL INFORMATION.

Section 14 of the Toxic Substances Control Act (15 U.S.C. 2613) is amended to read as follows:

“SEC. 14. CONFIDENTIAL INFORMATION.

“(a) IN GENERAL.—Except as provided in this section, the Administrator shall not disclose information that is exempt from disclosure pursuant to subsection (a) of section 552 of title 5, United States Code, by reason of subsection (b)(4) of that section—

“(1) that is reported to, or otherwise obtained by, the Administrator under this Act; and

“(2) for which the requirements of subsection (c) are met.

In any proceeding under section 552(a) of title 5, United States Code, to obtain information the disclosure of which has been denied because of the provisions of this subsection, the Administrator may not rely on section 552(b)(3) of such title to sustain the Administrator’s action.

“(b) INFORMATION NOT PROTECTED FROM DISCLOSURE.—

“(1) MIXED CONFIDENTIAL AND NONCONFIDENTIAL INFORMATION.—Information that is protected from disclosure under this section, and which is mixed with information that is not protected from disclosure under this section, does not lose its pro-
tection from disclosure notwithstanding that it is mixed with information that is not protected from disclosure.

“(2) INFORMATION FROM HEALTH AND SAFETY STUDIES.—Subsection (a) does not prohibit the disclosure of—

“(A) any health and safety study which is submitted under this Act with respect to—

“(i) any chemical substance or mixture which, on the date on which such study is to be disclosed has been offered for commercial distribution; or

“(ii) any chemical substance or mixture for which testing is required under section 4 or for which notification is required under section 5; and

“(B) any information reported to, or otherwise obtained by, the Administrator from a health and safety study which relates to a chemical substance or mixture described in clause (i) or (ii) of subparagraph (A).

This paragraph does not authorize the disclosure of any information, including formulas (including molecular structures) of a chemical substance or mixture, that discloses processes used in the manufac-
turing or processing of a chemical substance or mixture or, in the case of a mixture, the portion of the mixture comprised by any of the chemical substances in the mixture.

“(3) OTHER INFORMATION NOT PROTECTED FROM DISCLOSURE.—Subsection (a) does not prohibit the disclosure of—

“(A) any general information describing the manufacturing volumes, expressed as specific aggregated volumes or, if the Administrator determines that disclosure of specific aggregated volumes would reveal confidential information, expressed in ranges; or

“(B) a general description of a process used in the manufacture or processing and industrial, commercial, or consumer functions and uses of a chemical substance, mixture, or article containing a chemical substance or mixture, including information specific to an industry or industry sector that customarily would be shared with the general public or within an industry or industry sector.

“(4) BANS AND PHASE-OUTS.—

“(A) IN GENERAL.—If the Administrator promulgates a rule pursuant to section 6(a)
that establishes a ban or phase-out of a chemical substance or mixture, the protection from disclosure of any information under this section with respect to the chemical substance or mixture shall be presumed to no longer apply, subject to subsection (g)(1)(E) and subparagraphs (B) and (C) of this paragraph.

“(B) LIMITATIONS.—

“(i) CRITICAL USE.—In the case of a chemical substance or mixture for which a specific condition of use is subject to an exemption pursuant to section 6(g), if the Administrator establishes a ban or phase-out described in subparagraph (A) with respect to the chemical substance or mixture, the presumption against protection under such subparagraph shall only apply to information that relates solely to any conditions of use of the chemical substance or mixture to which the exemption does not apply.

“(ii) EXPORT.—In the case of a chemical substance or mixture for which there is manufacture, processing, or distribution in commerce that meets the conditions of sec-
tion 12(a)(1), if the Administrator estab-
ishes a ban or phase-out described in sub-
paragraph (A) with respect to the chemical
substance or mixture, the presumption
against protection under such subpara-
graph shall only apply to information that
relates solely to any other manufacture,
processing, or distribution in commerce of
the chemical substance or mixture for the
conditions of use subject to the ban or
phase-out, unless the Administrator makes
the determination in section 12(a)(2).

“(iii) SPECIFIC CONDITIONS OF
USE.—In the case of a chemical substance
or mixture for which the Administrator es-
tablishes a ban or phase-out described in
subparagraph (A) with respect to a specific
condition of use of the chemical substance
or mixture, the presumption against pro-
tection under such subparagraph shall only
apply to information that relates solely to
the condition of use of the chemical sub-
stance or mixture for which the ban or
phase-out is established.

“(C) REQUEST FOR NONDISCLOSURE.—
“(i) IN GENERAL.—A manufacturer or processor of a chemical substance or mixture subject to a ban or phase-out described in this paragraph may submit to the Administrator, within 30 days of receiving a notification under subsection (g)(2)(A), a request, including documentation supporting such request, that some or all of the information to which the notice applies should not be disclosed or that its disclosure should be delayed, and the Administrator shall review the request under subsection (g)(1)(E).

“(ii) EFFECT OF NO REQUEST OR DENIAL.—If no request for nondisclosure or delay is submitted to the Administrator under this subparagraph, or the Administrator denies such a request under subsection (g)(1)(A), the information shall not be protected from disclosure under this section.

“(5) CERTAIN REQUESTS.—If a request is made to the Administrator under section 552(a) of title 5, United States Code, for information reported to or otherwise obtained by the Administrator under this
Act that is not protected from disclosure under this subsection, the Administrator may not deny the request on the basis of section 552(b)(4) of title 5, United States Code.

“(c) REQUIREMENTS FOR CONFIDENTIALITY CLAIMS.—

“(1) ASSERTION OF CLAIMS.—

“(A) IN GENERAL.—A person seeking to protect from disclosure any information that person submits under this Act (including information described in paragraph (2)) shall assert to the Administrator a claim for protection from disclosure concurrent with submission of the information, in accordance with such rules regarding a claim for protection from disclosure as the Administrator has promulgated or may promulgate pursuant to this title.

“(B) INCLUSION.—An assertion of a claim under subparagraph (A) shall include a statement that the person has—

“(i) taken reasonable measures to protect the confidentiality of the information;

“(ii) determined that the information is not required to be disclosed or otherwise
made available to the public under any other Federal law;

“(iii) a reasonable basis to conclude that disclosure of the information is likely to cause substantial harm to the competitive position of the person; and

“(iv) a reasonable basis to believe that the information is not readily discoverable through reverse engineering.

“(C) ADDITIONAL REQUIREMENTS FOR CLAIMS REGARDING CHEMICAL IDENTITY INFORMATION.—In the case of a claim under subparagraph (A) for protection from disclosure of a specific chemical identity, the claim shall include a structurally descriptive generic name for the chemical substance that the Administrator may disclose to the public, subject to the condition that such generic name shall—

“(i) be consistent with guidance developed by the Administrator under paragraph (4)(A); and

“(ii) describe the chemical structure of the chemical substance as specifically as practicable while protecting those features of the chemical structure—
“(I) that are claimed as confidential; and

“(II) the disclosure of which would be likely to cause substantial harm to the competitive position of the person.

“(2) INFORMATION GENERALLY NOT SUBJECT TO SUBSTANTIATION REQUIREMENTS.—Subject to subsection (f), the following information shall not be subject to substantiation requirements under paragraph (3):

“(A) Specific information describing the processes used in manufacture or processing of a chemical substance, mixture, or article.

“(B) Marketing and sales information.

“(C) Information identifying a supplier or customer.

“(D) In the case of a mixture, details of the full composition of the mixture and the respective percentages of constituents.

“(E) Specific information regarding the use, function, or application of a chemical substance or mixture in a process, mixture, or article.
"(F) Specific production or import volumes of the manufacturer or processor.

"(G) Prior to the date on which a chemical substance is first offered for commercial distribution, the specific chemical identity of the chemical substance, including the chemical name, molecular formula, Chemical Abstracts Service number, and other information that would identify the specific chemical substance, if the specific chemical identity was claimed as confidential at the time it was submitted in a notice under section 5.

"(3) Substantiation requirements.—Except as provided in paragraph (2), a person asserting a claim to protect information from disclosure under this section shall substantiate the claim, in accordance with such rules as the Administrator has promulgated or may promulgate pursuant to this section.

"(4) Guidance.—The Administrator shall develop guidance regarding—

"(A) the determination of structurally descriptive generic names, in the case of claims for the protection from disclosure of specific chemical identity; and
“(B) the content and form of the statements of need and agreements required under paragraphs (4), (5), and (6) of subsection (d).

“(5) CERTIFICATION.—An authorized official of a person described in paragraph (1)(A) shall certify that the statement required to assert a claim submitted pursuant to paragraph (1)(B), and any information required to substantiate a claim submitted pursuant to paragraph (3), are true and correct.

“(d) EXCEPTIONS TO PROTECTION FROM DISCLOSURE.—Information described in subsection (a)—

“(1) shall be disclosed to an officer or employee of the United States—

“(A) in connection with the official duties of that person under any Federal law for the protection of health or the environment; or

“(B) for a specific Federal law enforcement purpose;

“(2) shall be disclosed to a contractor of the United States and employees of that contractor—

“(A) if, in the opinion of the Administrator, the disclosure is necessary for the satisfactory performance by the contractor of a contract with the United States for the performance of work in connection with this Act; and
“(B) subject to such conditions as the Administrator may specify;

“(3) shall be disclosed if the Administrator determines that disclosure is necessary to protect health or the environment against an unreasonable risk of injury to health or the environment, without consideration of costs or other nonrisk factors, including an unreasonable risk to a potentially exposed or susceptible subpopulation identified as relevant by the Administrator under the conditions of use;

“(4) shall be disclosed to a State, political subdivision of a State, or tribal government, on written request, for the purpose of administration or enforcement of a law, if such entity has 1 or more applicable agreements with the Administrator that are consistent with the guidance developed under subsection (c)(4)(B) and ensure that the entity will take appropriate measures, and has adequate authority, to maintain the confidentiality of the information in accordance with procedures comparable to the procedures used by the Administrator to safeguard the information;

“(5) shall be disclosed to a health or environmental professional employed by a Federal or State agency or tribal government or a treating physician
or nurse in a nonemergency situation if such person
provides a written statement of need and agrees to
sign a written confidentiality agreement with the Ad-
ministrator, subject to the conditions that—

“(A) the statement of need and confiden-
tiality agreement are consistent with the guid-
ance developed under subsection (c)(4)(B);

“(B) the statement of need shall be a
statement that the person has a reasonable
basis to suspect that—

“(i) the information is necessary for,
or will assist in—

“(I) the diagnosis or treatment of
1 or more individuals; or

“(II) responding to an environ-
mental release or exposure; and

“(ii) 1 or more individuals being diag-
nosed or treated have been exposed to the
chemical substance or mixture concerned,
or an environmental release of or exposure
to the chemical substance or mixture con-
cerned has occurred; and

“(C) the person will not use the informa-
tion for any purpose other than the health or
environmental needs asserted in the statement
of need, except as otherwise may be authorized by the terms of the agreement or by the person who has a claim under this section with respect to the information, except that nothing in this title prohibits the disclosure of any such information through discovery, subpoena, other court order, or any other judicial process otherwise allowed under applicable Federal or State law;

“(6) shall be disclosed in the event of an emergency to a treating or responding physician, nurse, agent of a poison control center, public health or environmental official of a State, political subdivision of a State, or tribal government, or first responder (including any individual duly authorized by a Federal agency, State, political subdivision of a State, or tribal government who is trained in urgent medical care or other emergency procedures, including a police officer, firefighter, or emergency medical technician) if such person requests the information, subject to the conditions that such person shall—

“(A) have a reasonable basis to suspect that—

“(i) a medical, public health, or environmental emergency exists;
“(ii) the information is necessary for, or will assist in, emergency or first-aid diagnosis or treatment; or

“(iii) 1 or more individuals being diagnosed or treated have likely been exposed to the chemical substance or mixture concerned, or a serious environmental release of or exposure to the chemical substance or mixture concerned has occurred; and

“(B) if requested by a person who has a claim with respect to the information under this section—

“(i) provide a written statement of need and agree to sign a confidentiality agreement, as described in paragraph (5); and

“(ii) submit to the Administrator such statement of need and confidentiality agreement as soon as practicable, but not necessarily before the information is disclosed;

“(7) may be disclosed if the Administrator determines that disclosure is relevant in a proceeding under this Act, subject to the condition that the dis-
closure is made in such a manner as to preserve con-

fidentiality to the extent practicable without impair-

ing the proceeding; and

“(8) shall be disclosed if the information is re-

quired to be made public under any other provision

of Federal law.

“(e) DURATION OF PROTECTION FROM DISCLOS-

URE.—

“(1) IN GENERAL.—Subject to paragraph (2),

subsection (f)(3), and section 8(b), the Adminis-

trator shall protect from disclosure information de-

dscribed in subsection (a)—

“(A) in the case of information described

in subsection (e)(2), until such time as—

“(i) the person that asserted the claim

notifies the Administrator that the person

is withdrawing the claim, in which case the

information shall not be protected from

disclosure under this section; or

“(ii) the Administrator becomes aware

that the information does not qualify for

protection from disclosure under this sec-

tion, in which case the Administrator shall

take any actions required under sub-

sections (f) and (g); and
“(B) in the case of information other than information described in subsection (e)(2)—

“(i) for a period of 10 years from the date on which the person asserts the claim with respect to the information submitted to the Administrator; or

“(ii) if applicable before the expiration of such 10-year period, until such time as—

“(I) the person that asserted the claim notifies the Administrator that the person is withdrawing the claim, in which case the information shall not be protected from disclosure under this section; or

“(II) the Administrator becomes aware that the information does not qualify for protection from disclosure under this section, in which case the Administrator shall take any actions required under subsections (f) and (g).

“(2) EXTENSIONS.—

“(A) IN GENERAL.—In the case of information other than information described in sub-
section (c)(2), not later than the date that is 60
days before the expiration of the period de-
scribed in paragraph (1)(B)(i), the Adminis-
trator shall provide to the person that asserted
the claim a notice of the impending expiration
of the period.

“(B) Request.—

“(i) In General.—Not later than the
date that is 30 days before the expiration
of the period described in paragraph
(1)(B)(i), a person reasserting the relevant
claim shall submit to the Administrator a
request for extension substantiating, in ac-
cordance with subsection (c)(3), the need
to extend the period.

“(ii) Action by Administrator.—
Not later than the date of expiration of the
period described in paragraph (1)(B)(i),
the Administrator shall, in accordance with
subsection (g)(1)—

“(I) review the request submitted
under clause (i);

“(II) make a determination re-
garding whether the claim for which
the request was submitted continues
to meet the relevant requirements of
this section; and

“(III)(aa) grant an extension of
10 years; or

“(bb) deny the request.

“(C) No limit on number of extensions.—There shall be no limit on the number
of extensions granted under this paragraph, if
the Administrator determines that the relevant
request under subparagraph (B)(i)—

“(i) establishes the need to extend the
period; and

“(ii) meets the requirements estab-
lished by the Administrator.

“(f) Review and resubstantiation.—

“(1) Discretion of administrator.—The
Administrator may require any person that has
claimed protection for information from disclosure
under this section, whether before, on, or after the
date of enactment of the Frank R. Launtenberg
Chemical Safety for the 21st Century Act, to re-
assert and substantiate or resubstantiate the claim
in accordance with this section—
“(A) after the chemical substance is designated as a high-priority substance under section 6(b);

“(B) for any chemical substance designated as an active substance under section 8(b)(5)(B)(iii); or

“(C) if the Administrator determines that disclosure of certain information currently protected from disclosure would be important to assist the Administrator in conducting risk evaluations or promulgating rules under section 6.

“(2) REVIEW REQUIRED.—The Administrator shall review a claim for protection of information from disclosure under this section and require any person that has claimed protection for that information, whether before, on, or after the date of enactment of the Frank R. Launtenberg Chemical Safety for the 21st Century Act, to reassert and substantiate or resubstantiate the claim in accordance with this section—

“(A) as necessary to determine whether the information qualifies for an exemption from disclosure in connection with a request for in-
formation received by the Administrator under section 552 of title 5, United States Code;

“(B) if the Administrator has a reasonable basis to believe that the information does not qualify for protection from disclosure under this section; or

“(C) for any chemical substance the Administrator determines under section 6(b)(4)(A) presents an unreasonable risk of injury to health or the environment.

“(3) PERIOD OF PROTECTION.—If the Administrator requires a person to reassert and substantiate or resubstantiate a claim under this subsection, and determines that the claim continues to meet the relevant requirements of this section, the Administrator shall protect the information subject to the claim from disclosure for a period of 10 years from the date of such determination, subject to any subsequent requirement by the Administrator under this subsection.

“(g) DUTIES OF ADMINISTRATOR.—

“(1) DETERMINATION.—

“(A) IN GENERAL.—Except for claims regarding information described in subsection (c)(2), the Administrator shall, subject to sub-
paragraph (C), not later than 90 days after the
receipt of a claim under subsection (e), and not
later than 30 days after the receipt of a request
for extension of a claim under subsection (e) or
a request under subsection (b)(4)(C), review
and approve, approve in part and deny in part,
or deny the claim or request.

“(B) REASONS FOR DENIAL.—If the Ad-
ministrator denies or denies in part a claim or
request under subparagraph (A) the Adminis-
trator shall provide to the person that asserted
the claim or submitted the request a written
statement of the reasons for the denial or de-
nial in part of the claim or request.

“(C) SUBSETS.—The Administrator
shall—

“(i) except with respect to information
described in subsection (e)(2)(G), review
all claims or requests under this section for
the protection from disclosure of the spe-
cific chemical identity of a chemical sub-
stance; and

“(ii) review a representative subset,
comprising at least 25 percent, of all other
claims or requests for protection from disclosure under this section.

“(D) Effect of Failure to Act.—The failure of the Administrator to make a decision regarding a claim or request for protection from disclosure or extension under this section shall not have the effect of denying or eliminating a claim or request for protection from disclosure.

“(E) Determination of Requests under Subsection (b)(4)(C).—With respect to a request submitted under subsection (b)(4)(C), the Administrator shall, with the objective of ensuring that information relevant to the protection of health and the environment is disclosed to the extent practicable, determine whether the documentation provided by the person rebuts what shall be the presumption of the Administrator that the public interest in the disclosure of the information outweighs the public or proprietary interest in maintaining the protection for all or a portion of the information that the person has requested not be disclosed or for which disclosure be delayed.

“(2) Notification.—
“(A) IN GENERAL.—Except as provided in subparagraph (B) and subsections (b), (d), and (e), if the Administrator denies or denies in part a claim or request under paragraph (1), concludes, in accordance with this section, that the information does not qualify for protection from disclosure, intends to disclose information pursuant to subsection (d), or promulgates a rule under section 6(a) establishing a ban or phase-out with respect to a chemical substance or mixture, the Administrator shall notify, in writing, the person that asserted the claim or submitted the request of the intent of the Administrator to disclose the information or not protect the information from disclosure under this section. The notice shall be furnished by certified mail (return receipt requested), by personal delivery, or by other means that allows verification of the fact and date of receipt.

“(B) DISCLOSURE OF INFORMATION.—Except as provided in subparagraph (C), the Administrator shall not disclose information under this subsection until the date that is 30 days after the date on which the person that asserted
the claim or submitted the request receives notification under subparagraph (A).

“(C) EXCEPTIONS.—

“(i) FIFTEEN DAY NOTIFICATION.—

For information the Administrator intends to disclose under subsections (d)(3), (d)(4), (d)(5), and (j), the Administrator shall not disclose the information until the date that is 15 days after the date on which the person that asserted the claim or submitted the request receives notification under subparagraph (A), except that, with respect to information to be disclosed under subsection (d)(3), if the Administrator determines that disclosure of the information is necessary to protect against an imminent and substantial harm to health or the environment, no prior notification shall be necessary.

“(ii) NOTIFICATION AS SOON AS PRACTICABLE.—For information the Administrator intends to disclose under paragraph (6) of subsection (d), the Administrator shall notify the person that submitted the information that the information has been
disclosed as soon as practicable after disclosure of the information.

“(iii) NO NOTIFICATION REQUIRED.—Notification shall not be required—

“(I) for the disclosure of information under paragraphs (1), (2), (7), or (8) of subsection (d); or

“(II) for the disclosure of information for which—

“(aa) the Administrator has provided to the person that asserted the claim a notice under subsection (e)(2)(A); and

“(bb) such person does not submit to the Administrator a request under subsection (e)(2)(B) on or before the deadline established in subsection (e)(2)(B)(i).

“(D) APPEALS.—

“(i) ACTION TO RESTRAIN DISCLOSURE.—If a person receives a notification under this paragraph and believes the information is protected from disclosure under this section, before the date on which the information is to be disclosed
pursuant to subparagraph (B) or (C) the
person may bring an action to restrain dis-
closure of the information in—

“(I) the United States district
court of the district in which the com-
plainant resides or has the principal
place of business; or

“(II) the United States District
Court for the District of Columbia.

“(ii) NO DISCLOSURE.—

“(I) IN GENERAL.—Subject to
subsection (d), the Administrator shall
not disclose information that is the
subject of an appeal under this para-
graph before the date on which the
applicable court rules on an action
under clause (i).

“(II) EXCEPTION.—Subclause (I)
shall not apply to disclosure of infor-
mation described under subsections
(d)(4) and (j).

“(3) REQUEST AND NOTIFICATION SYSTEM.—
The Administrator, in consultation with the Director
of the Centers for Disease Control and Prevention,
shall develop a request and notification system that,
in a format and language that is readily accessible
and understandable, allows for expedient and swift
access to information disclosed pursuant to para-
graphs (5) and (6) of subsection (d).

“(4) UNIQUE IDENTIFIER.—The Administrator
shall—

“(A)(i) develop a system to assign a
unique identifier to each specific chemical iden-
tity for which the Administrator approves a re-
quest for protection from disclosure, which shall
not be either the specific chemical identity or a
structurally descriptive generic term; and

“(ii) apply that identifier consistently to all
information relevant to the applicable chemical
substance;

“(B) annually publish and update a list of
chemical substances, referred to by their unique
identifiers, for which claims to protect the spe-
cific chemical identity from disclosure have been
approved, including the expiration date for each
such claim;

“(C) ensure that any nonconfidential infor-
mation received by the Administrator with re-
spect to a chemical substance included on the
list published under subparagraph (B) while the
specific chemical identity of the chemical sub-
stance is protected from disclosure under this
section identifies the chemical substance using
the unique identifier; and

“(D) for each claim for protection of a spe-
cific chemical identity that has been denied by
the Administrator or expired, or that has been
withdrawn by the person who asserted the
claim, and for which the Administrator has
used a unique identifier assigned under this
paragraph to protect the specific chemical iden-
tity in information that the Administrator has
made public, clearly link the specific chemical
identity to the unique identifier in such infor-
mation to the extent practicable.

“(h) CRIMINAL PENALTY FOR WRONGFUL DISCLOS-
URE.—

“(1) INDIVIDUALS SUBJECT TO PENALTY.—

“(A) IN GENERAL.—Subject to subpara-
graph (C) and paragraph (2), an individual de-
scribed in subparagraph (B) shall be fined
under title 18, United States Code, or impris-
oned for not more than 1 year, or both.
“(B) DESCRIPTION.—An individual referred to in subparagraph (A) is an individual who—

“(i) pursuant to this section, obtained possession of, or has access to, information protected from disclosure under this section; and

“(ii) knowing that the information is protected from disclosure under this section, willfully discloses the information in any manner to any person not entitled to receive that information.

“(C) EXCEPTION.—This paragraph shall not apply to any medical professional (including an emergency medical technician or other first responder) who discloses any information obtained under paragraph (5) or (6) of subsection (d) to a patient treated by the medical professional, or to a person authorized to make medical or health care decisions on behalf of such a patient, as needed for the diagnosis or treatment of the patient.

“(2) OTHER LAWS.—Section 1905 of title 18, United States Code, shall not apply with respect to the publishing, divulging, disclosure, or making
known of, or making available, information reported
to or otherwise obtained by the Administrator under
this Act.

“(i) APPLICABILITY.—

“(1) IN GENERAL.—Except as otherwise pro-
vided in this section, section 8, or any other applica-
ble Federal law, the Administrator shall have no au-
thority—

“(A) to require the substantiation or re-
substantiation of a claim for the protection
from disclosure of information reported to or
otherwise obtained by the Administrator under
this Act prior to the date of enactment of the
Frank R. Launtenberg Chemical Safety for the
21st Century Act; or

“(B) to impose substantiation or re-
substantiation requirements, with respect to the
protection of information described in sub-
section (a), under this Act that are more exten-
sive than those required under this section.

“(2) ACTIONS PRIOR TO PROMULGATION OF
RULES.—Nothing in this Act prevents the Adminis-
trator from reviewing, requiring substantiation or re-
substantiation of, or approving, approving in part, or
denying any claim for the protection from disclosure
of information before the effective date of such rules applicable to those claims as the Administrator may promulgate after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act.

“(j) ACCESS BY CONGRESS.—Notwithstanding any limitation contained in this section or any other provision of law, all information reported to or otherwise obtained by the Administrator (or any representative of the Administrator) under this Act shall be made available, upon written request of any duly authorized committee of the Congress, to such committee.”.

SEC. 12. PENALTIES.

Section 16 of the Toxic Substances Control Act (15 U.S.C. 2615) is amended—

(1) in subsection (a)(1), by striking “$25,000” and inserting “$37,500”; and

(2) in subsection (b)—

(A) by striking “Any person” and inserting the following:

“(1) IN GENERAL.—Any person”;

(B) by striking “$25,000” and inserting “$50,000”; and

(C) by adding at the end the following:
“(2) IMMINENT DANGER OF DEATH OR SERIOUS BODILY INJURY.—

“(A) IN GENERAL.—Any person who knowingly and willfully violates any provision of section 15 or 409, and who knows at the time of the violation that the violation places an individual in imminent danger of death or serious bodily injury, shall be subject on conviction to a fine of not more than $250,000, or imprisonment for not more than 15 years, or both.

“(B) ORGANIZATIONS.—Notwithstanding the penalties described in subparagraph (A), an organization that commits a knowing violation described in subparagraph (A) shall be subject on conviction to a fine of not more than $1,000,000 for each violation.

“(C) INCORPORATION OF CORRESPONDING PROVISIONS.—Subparagraphs (B) through (F) of section 113(c)(5) of the Clean Air Act (42 U.S.C. 7413(e)(5)(B)—(F)) shall apply to the prosecution of a violation under this paragraph.”.

SEC. 13. STATE-FEDERAL RELATIONSHIP.

Section 18 of the Toxic Substances Control Act (15 U.S.C. 2617) is amended—
(1) by amending subsection (a) to read as follows:

“(a) IN GENERAL.—

“(1) ESTABLISHMENT OR ENFORCEMENT.—Except as otherwise provided in subsections (c), (d), (e), (f), and (g), and subject to paragraph (2), no State or political subdivision of a State may establish or continue to enforce any of the following:

“(A) DEVELOPMENT OF INFORMATION.—A statute or administrative action to require the development of information about a chemical substance or category of chemical substances that is reasonably likely to produce the same information required under section 4, 5, or 6 in—

“(i) a rule promulgated by the Administrator;

“(ii) a consent agreement entered into by the Administrator; or

“(iii) an order issued by the Administrator.

“(B) CHEMICAL SUBSTANCES FOUND NOT TO PRESENT AN UNREASONABLE RISK OR RESTRICTED.—A statute, criminal penalty, or administrative action to prohibit or otherwise restrict the manufacture, processing, or distribu-
tion in commerce or use of a chemical sub-
stance—

“(i) for which the determination de-
scribed in section 6(i)(1) is made, con-
sistent with the scope of the risk evalua-
tion under section (6)(b)(4)(D); or

“(ii) for which a final rule is promul-
gated under section 6(a), after the effective
date of the rule issued under section 6(a)
for the chemical substance, consistent with
the scope of the risk evaluation under sec-
tion (6)(b)(4)(D).

“(C) SIGNIFICANT NEW USE.—A statute or
administrative action requiring the notification
of a use of a chemical substance that the Ad-
ministrator has specified as a significant new
use and for which the Administrator has re-
quired notification pursuant to a rule promul-
gated under section 5.

“(2) EFFECTIVE DATE OF PREEMPTION.—
Under this subsection, Federal preemption of stat-
utes and administrative actions applicable to specific
chemical substances shall not occur until the effec-
tive date of the applicable action described in para-
graph (1) taken by the Administrator.”;
(2) by amending subsection (b) to read as follows:

“(b) New Statutes, Criminal Penalties, or Administrative Actions Creating Prohibitions or Other Restrictions.—

“(1) In general.—Except as provided in subsections (c), (d), (e), (f), and (g), beginning on the date on which the Administrator defines the scope of a risk evaluation for a chemical substance under section 6(b)(4)(D) and ending on the date on which the deadline established pursuant to section 6(b)(4)(G) for completion of the risk evaluation expires, or on the date on which the Administrator publishes the risk evaluation under section 6(b)(4)(C), whichever is earlier, no State or political subdivision of a State may establish a statute, criminal penalty, or administrative action prohibiting or otherwise restricting the manufacture, processing, distribution in commerce, or use of such chemical substance that is a high-priority substance designated under 6(b)(1)(B)(i), such chemical substance that has been identified under section 6(b)(2)(A) (except for the first 10 chemical substances so identified), or such chemical substance that has been selected for risk evaluation under section 6(b)(4)(E)(iv)(II).
“(2) Effect of subsection.—This subsection does not restrict the authority of a State or political subdivision of a State to continue to enforce any statute enacted, or administrative action taken, prior to the date on which the Administrator defines and publishes the scope of a risk evaluation under section 6(b)(4)(D).”; and

(3) by adding at the end the following:

“(c) Scope of preemption.—Federal preemption under subsections (a) and (b) of statutes, criminal penalties, and administrative actions applicable to specific chemical substances shall apply only to—

“(1) with respect to subsection (a)(1)(A), the chemical substances or category of chemical substances subject to a rule, order, or consent agreement under section 4;

“(2) with respect to subsections (a)(1)(B) and (b), the hazards, exposures, risks, and uses or conditions of use of such chemical substances consistent with the scope of the risk evaluation under 6(b)(4)(D); or

“(3) with respect to subsection (a)(1)(C), the uses of such chemical substances that the Administrator has specified as significant new uses and for
which the Administrator has required notification pursuant to a rule promulgated under section 5.

“(d) EXCEPTIONS.—

“(1) NO PREEMPTION OF STATUTES AND ADMINISTRATIVE ACTIONS.—

“(A) IN GENERAL.—Nothing in this Act, nor any amendment made by the Frank R. Lautenberg Chemical Safety for the 21st Century Act, nor any rule, standard of performance, risk evaluation, or scientific assessment implemented pursuant to this Act, shall affect the right of a State or a political subdivision of a State to adopt or enforce any rule, standard of performance, risk evaluation, scientific assessment, or any other protection for public health or the environment that—

“(i) is adopted or authorized under the authority of any other Federal law or adopted to satisfy or obtain authorization or approval under any other Federal law;

“(ii) implements a reporting, monitoring, or other information obligation for the chemical substance not otherwise required by the Administrator under this Act or required under any other Federal law;
“(iii) is adopted pursuant to authority under a law of the State or political subdivision of the State related to water quality, air quality, or waste treatment or disposal, except to the extent that the action—

“(I) imposes a restriction on the manufacture, processing, distribution in commerce, or use of a chemical substance; and

“(II)(aa) addresses the same hazards and exposures, with respect to the same conditions of use as are included in the scope of the risk evaluation published pursuant to section 6(b)(4)(D), but is inconsistent with the action of the Administrator; or

“(bb) would cause a violation of the applicable action by the Administrator under section 5 or 6; or

“(iv) subject to subparagraph (B), is identical to a requirement prescribed by the Administrator.

“(B) IDENTICAL REQUIREMENTS.—
“(i) IN GENERAL.—The penalties and other sanctions applicable under a law of a State or political subdivision of a State in the event of noncompliance with the identical requirement shall be no more stringent than the penalties and other sanctions available to the Administrator under section 16 of this Act.

“(ii) PENALTIES.—In the case of an identical requirement—

“(I) a State or political subdivision of a State may not assess a penalty for a specific violation for which the Administrator has assessed an adequate penalty under section 16; and

“(II) if a State or political subdivision of a State has assessed a penalty for a specific violation, the Administrator may not assess a penalty for that violation in an amount that would cause the total of the penalties assessed for the violation by the State or political subdivision of a State and the Administrator combined to exceed
the maximum amount that may be assessed for that violation by the Administrator under section 16.

“(2) Applicability to certain rules or orders.—

“(A) Prior rules and orders.—Nothing in this section shall be construed as modifying the preemptive effect under this section, as in effect on the day before the effective date of the Frank R. Launtenberg Chemical Safety for the 21st Century Act, of any rule or order promulgated or issued under this Act prior to that effective date.

“(B) Certain chemical substances and mixtures.—With respect to a chemical substance or mixture for which any rule or order was promulgated or issued under section 6 prior to the effective date of the Frank R. Launtenberg Chemical Safety for the 21st Century Act with respect to manufacturing, processing, distribution in commerce, use, or disposal of the chemical substance or mixture, nothing in this section shall be construed as modifying the preemptive effect of this section as in effect prior to the enactment of the Frank
R. Lautenberg Chemical Safety for the 21st Century Act of any rule or order that is promulgated or issued with respect to such chemical substance or mixture under section 6 after that effective date, unless the latter rule or order is with respect to a chemical substance or mixture containing a chemical substance and follows a designation of that chemical substance as a high-priority substance under section 6(b)(1)(B)(i), the identification of that chemical substance under section 6(b)(2)(A), or the selection of that chemical substance for risk evaluation under section 6(b)(4)(E)(iv)(II).

“(e) PRESERVATION OF CERTAIN LAWS.—

“(1) IN GENERAL.—Nothing in this Act, subject to subsection (g) of this section, shall—

“(A) be construed to preempt or otherwise affect the authority of a State or political subdivision of a State to continue to enforce any action taken or requirement imposed or requirement enacted relating to a specific chemical substance before April 22, 2016, under the authority of a law of the State or political subdivision of the State that prohibits or otherwise restricts manufacturing, processing, distribution
in commerce, use, or disposal of a chemical sub-
stance; or

“(B) be construed to preempt or otherwise
affect any action taken pursuant to a State law
that was in effect on August 31, 2003.

“(2) Effect of subsection.—This sub-
section does not affect, modify, or alter the relation-
ship between Federal law and laws of a State or po-
litical subdivision of a State pursuant to any other
Federal law.

“(f) Waivers.—

“(1) Discretionary exemptions.—Upon ap-
lication of a State or political subdivision of a
State, the Administrator may, by rule, exempt from
subsection (a), under such conditions as may be pre-
scribed in the rule, a statute, criminal penalty, or
administrative action of that State or political sub-
division of the State that relates to the effects of ex-
posure to a chemical substance under the conditions
of use if the Administrator determines that—

“(A) compelling conditions warrant grant-
ing the waiver to protect health or the environ-
ment;

“(B) compliance with the proposed require-
ment of the State or political subdivision of the
State would not unduly burden interstate commerce in the manufacture, processing, distribution in commerce, or use of a chemical substance;

“(C) compliance with the proposed requirement of the State or political subdivision of the State would not cause a violation of any applicable Federal law, rule, or order; and

“(D) in the judgment of the Administrator, the proposed requirement of the State or political subdivision of the State is designed to address a risk of a chemical substance, under the conditions of use, that was identified—

“(i) consistent with the best available science;

“(ii) using supporting studies conducted in accordance with sound and objective scientific practices; and

“(iii) based on the weight of the scientific evidence.

“(2) REQUIRED EXEMPTIONS.—Upon application of a State or political subdivision of a State, the Administrator shall exempt from subsection (b) a statute or administrative action of a State or political subdivision of a State that relates to the effects
of exposure to a chemical substance under the conditions of use if the Administrator determines that—

“(A)(i) compliance with the proposed requirement of the State or political subdivision of the State would not unduly burden interstate commerce in the manufacture, processing, distribution in commerce, or use of a chemical substance;

“(ii) compliance with the proposed requirement of the State or political subdivision of the State would not cause a violation of any applicable Federal law, rule, or order; and

“(iii) the State or political subdivision of the State has a concern about the chemical substance or use of the chemical substance based in peer-reviewed science; or

“(B) no later than the date that is 18 months after the date on which the Administrator has initiated the prioritization process for a chemical substance under the rule promulgated pursuant to section 6(b)(1)(A), or the date on which the Administrator publishes the scope of the risk evaluation for a chemical substance under section 6(b)(4)(D), whichever is sooner, the State or political subdivision of the
State has enacted a statute or proposed or finalized an administrative action intended to prohibit or otherwise restrict the manufacture, processing, distribution in commerce, or use of the chemical substance.

“(3) **DETERMINATION OF A WAIVER REQUEST.**—The duty of the Administrator to grant or deny a waiver application shall be nondelegable and shall be exercised—

“(A) not later than 180 days after the date on which an application under paragraph (1) is submitted; and

“(B) not later than 110 days after the date on which an application under paragraph (2) is submitted.

“(4) **FAILURE TO MAKE A DETERMINATION.**—If the Administrator fails to make a determination under paragraph (3)(B) during the 110-day period beginning on the date on which an application under paragraph (2) is submitted, the statute or administrative action of the State or political subdivision of the State that was the subject of the application shall not be considered to be an existing statute or administrative action for purposes of subsection (b)
by reason of the failure of the Administrator to make a determination.

“(5) NOTICE AND COMMENT.—Except in the case of an application approved under paragraph (9), the application of a State or political subdivision of a State under this subsection shall be subject to public notice and comment.

“(6) FINAL AGENCY ACTION.—The decision of the Administrator on the application of a State or political subdivision of a State shall be—

“(A) considered to be a final agency action; and

“(B) subject to judicial review.

“(7) DURATION OF WAIVERS.—A waiver granted under paragraph (2) or approved under paragraph (9) shall remain in effect until such time as the Administrator publishes the risk evaluation under section 6(b).

“(8) JUDICIAL REVIEW OF WAIVERS.—Not later than 60 days after the date on which the Administrator makes a determination on an application of a State or political subdivision of a State under paragraph (1) or (2), any person may file a petition for judicial review in the United States Court of Appeals
for the District of Columbia Circuit, which shall have exclusive jurisdiction over the determination.

“(9) APPROVAL.—

“(A) AUTOMATIC APPROVAL.—If the Administrator fails to meet the deadline established under paragraph (3)(B), the application of a State or political subdivision of a State under paragraph (2) shall be automatically approved, effective on the date that is 10 days after the deadline.

“(B) REQUIREMENTS.—Notwithstanding paragraph (6), approval of a waiver application under subparagraph (A) for failure to meet the deadline under paragraph (3)(B) shall not be considered final agency action or be subject to judicial review or public notice and comment.

“(g) SAVINGS.—

“(1) NO PREEMPTION OF COMMON LAW OR STATUTORY CAUSES OF ACTION FOR CIVIL RELIEF OR CRIMINAL CONDUCT.—

“(A) IN GENERAL.—Nothing in this Act, nor any amendment made by the Frank R. Lautenberg Chemical Safety for the 21st Century Act, nor any standard, rule, requirement, standard of performance, risk evaluation, or sci-
entific assessment implemented pursuant to this
Act, shall be construed to preempt, displace, or
supplant any State or Federal common law
rights or any State or Federal statute creating
a remedy for civil relief, including those for civil
damage, or a penalty for a criminal conduct.

“(B) CLARIFICATION OF NO PREEM-
PTION.—Notwithstanding any other provision of
this Act, nothing in this Act, nor any amend-
ments made by the Frank R. Lautenberg
Chemical Safety for the 21st Century Act, shall
preempt or preclude any cause of action for
personal injury, wrongful death, property dam-
age, or other injury based on negligence, strict
liability, products liability, failure to warn, or
any other legal theory of liability under any
State law, maritime law, or Federal common
law or statutory theory.

“(2) NO EFFECT ON PRIVATE REMEDIES.—

“(A) IN GENERAL.—Nothing in this Act,
nor any amendments made by the Frank R.
Lautenberg Chemical Safety for the 21st Cen-
tury Act, nor any rules, regulations, require-
ments, risk evaluations, scientific assessments,
or orders issued pursuant to this Act shall be
interpreted as, in either the plaintiff’s or defendant’s favor, dispositive in any civil action.

“(B) AUTHORITY OF COURTS.—This Act does not affect the authority of any court to make a determination in an adjudicatory proceeding under applicable State or Federal law with respect to the admission into evidence or any other use of this Act or rules, regulations, requirements, standards of performance, risk evaluations, scientific assessments, or orders issued pursuant to this Act.”.

SEC. 14. JUDICIAL REVIEW.

Section 19(a) of the Toxic Substances Control Act (15 U.S.C. 2618(a)) is amended—

(1) in paragraph (1), by adding at the end the following:

“(C)(i) Not later than 60 days after the publication of a designation under section 6(b)(1)(B)(ii) or (iii), any person may commence a civil action to challenge the designation.

“(ii) The United States Court of Appeals for the District of Columbia Circuit shall have exclusive jurisdiction over a civil action filed under this sub-paragraph.”; and

(2) by striking paragraph (3).
SEC. 15. CITIZENS’ CIVIL ACTIONS.

Section 20(b) of the Toxic Substances Control Act (15 U.S.C. 2619(b)) is amended—

(1) in paragraph (1)(B), by striking “or” at the end; and

(2) in paragraph (2), by striking the period at the end and inserting the following: “, except that no prior notification shall be required in the case of a civil action brought to compel a decision by the Administrator pursuant to section 18(f)(3)(B); or

“(3) in the case of a civil action brought to compel a decision by the Administrator pursuant to section 18(f)(3)(B), after the date that is 60 days after the deadline specified in section 18(f)(3)(B).”.

SEC. 16. STUDIES.

Section 25 of the Toxic Substances Control Act (15 U.S.C. 2624) is repealed.

SEC. 17. ADMINISTRATION OF THE ACT.

Section 26 of the Toxic Substances Control Act (15 U.S.C. 2625) is amended—

(1) in subsection (b)(1)—

(A) by striking “of a reasonable fee”;

(B) by striking “data under section 4 or 5 to defray the cost of administering this Act” and inserting “information under section 4 or a notice or other information to be reviewed by
the Administrator under section 5, or who man-
ufactures or processes a chemical substance
that is the subject of a risk evaluation under
section 6(b), of a fee that is sufficient and not
more than reasonably necessary to defray the
cost related to such chemical substance of ad-
ministering sections 4, 5, and 6, and collecting,
processing, reviewing, and providing access to
and protecting from disclosure as appropriate
under section 14 information on chemical sub-
stances under this title, including contractor
costs incurred by the Administrator”;  
(C) by striking “Such rules shall not pro-
vide for any fee in excess of $2,500 or, in the
case of a small business concern, any fee in ex-
cess of $100.”; and
(D) by striking “submit the data and the
cost to the Administrator of reviewing such
data” and inserting “pay such fee and the cost
to the Administrator of carrying out the activi-
ties described in this paragraph”;
(2) in subsection (b)—
(A) in paragraph (2), by striking “para-
graph (1)” and inserting “paragraph (4)”;
and
(B) by adding at the end the following:
“(3) FUND.—

“(A) ESTABLISHMENT.—There is established in the Treasury of the United States a fund, to be known as the TSCA Service Fee Fund (in this paragraph referred to as the ‘Fund’), consisting of such amounts as are deposited in the Fund under this paragraph.

“(B) COLLECTION AND DEPOSIT OF FEES.—Subject to the conditions of subparagraph (C), the Administrator shall collect the fees described in this subsection and deposit those fees in the Fund.

“(C) USE OF FUNDS BY ADMINISTRATOR.—Fees authorized under this section shall be collected and available for obligation only to the extent and in the amount provided in advance in appropriations Acts, and shall be available without fiscal year limitation for use in defraying the costs of the activities described in paragraph (1).

“(D) ACCOUNTING AND AUDITING.—

“(i) ACCOUNTING.—The Administrator shall biennially prepare and submit to the Committee on Environment and Public Works of the Senate and the Committee on Energy and Commerce of the House of Representatives a report that includes an accounting of the fees paid to
the Administrator under this paragraph and amounts disbursed from the Fund for the period covered by the report, as reflected by financial statements provided in accordance with sections 3515 and 3521 of title 31, United States Code.

“(ii) AUDITING.—

“(I) IN GENERAL.—For the purpose of section 3515(c) of title 31, United States Code, the Fund shall be considered a component of a covered executive agency.

“(II) COMPONENTS OF AUDIT.—The annual audit required in accordance with sections 3515 and 3521 of title 31, United States Code, of the financial statements of activities carried out using amounts from the Fund shall include an analysis of—

“(aa) the fees collected and amounts disbursed under this subsection;

“(bb) the reasonableness of the fees in place as of the date of the audit to meet current and projected costs of administering the provisions
of this title for which the fees may be
used; and

“(cc) the number of requests for
a risk evaluation made by manufac-
turers under section 6(b)(4)(C)(ii).

“(III) **FEDERAL RESPONSIBILITY.**—
The Inspector General of the Environ-
mental Protection Agency shall conduct
the annual audit described in subclause
(II) and submit to the Administrator a re-
port that describes the findings and any
recommendations of the Inspector General
resulting from the audit.

“(4) **AMOUNT AND ADJUSTMENT OF FEES; RE-
FUNDS.**—In setting fees under this section, the Adminis-
trator shall—

“(A) prescribe lower fees for small business
concerns, after consultation with the Administrator
of the Small Business Administration;

“(B) set the fees established under paragraph
(1) at levels such that the fees will, in aggregate,
provide a sustainable source of funds to annually de-
fray—

“(i) the lower of—
“(I) 25 percent of the costs to the Administrator of carrying out sections 4, 5, and 6, and of collecting, processing, reviewing, and providing access to and protecting from disclosure as appropriate under section 14 information on chemical substances under this title, other than the costs to conduct and complete risk evaluations under section 6(b); or

“(II) $25,000,000 (subject to adjustment pursuant to subparagraph (F)); and

“(ii) the costs of risk evaluations specified in subparagraph (D);

“(C) reflect an appropriate balance in the assessment of fees between manufacturers and processors, and allow the payment of fees by consortia of manufacturers or processors;

“(D) notwithstanding subparagraph (B)—

“(i) except as provided in clause (ii), for chemical substances for which the Administrator has granted a request from a manufacturer pursuant to section 6(b)(4)(C)(ii), establish the fee at a level sufficient to defray the full costs to the Administrator of conducting the risk evaluation under section 6(b);
“(ii) for chemical substances for which the Administrator has granted a request from a manufacturer pursuant to section 6(b)(4)(C)(ii), and which are included in the 2014 update of the TSCA Work Plan for Chemical Assessments, establish the fee at a level sufficient to defray 50 percent of the costs to the Administrator of conducting the risk evaluation under section 6(b); and

“(iii) apply fees collected pursuant to clauses (i) and (ii) only to defray the costs described in those clauses;

“(E) prior to the establishment or amendment of any fees under paragraph (1), consult and meet with parties potentially subject to the fees or their representatives, subject to the condition that no obligation under the Federal Advisory Committee Act (5 U.S.C. App.) or subchapter II of chapter 5 of title 5, United States Code, is applicable with respect to such meetings;

“(F) beginning with the fiscal year that is 3 years after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, and every 3 years thereafter, after consultation with parties potentially subject to the fees and their
representatives pursuant to subparagraph (E), increase or decrease the fees established under paragraph (1) as necessary to adjust for inflation and to ensure that funds deposited in the Fund are sufficient to defray—

“(i) approximately but not more than 25 percent of the costs to the Administrator of carrying out sections 4, 5, and 6, and of collecting, processing, reviewing, and providing access to and protecting from disclosure as appropriate under section 14 information on chemical substances under this title, other than the costs to conduct and complete risk evaluations requested under section 6(b)(4)(C)(ii); and

“(ii) the costs of risk evaluations specified in subparagraph (D); and

“(G) if a notice submitted under section 5 is not reviewed or such a notice is withdrawn, refund the fee or a portion of the fee if no substantial work was performed on the notice.

“(5) MINIMUM AMOUNT OF APPROPRIATIONS.—Fees may not be assessed for a fiscal year under this section unless the amount of appropriations for the Chemical Risk Review and Reduction program project of the Environmental Protection Agency for the fiscal year (excluding
the amount of any fees appropriated for the fiscal year) are equal to or greater than the amount of appropriations for that program project for fiscal year 2014.

“(6) TERMINATION.—The authority provided by this subsection shall terminate at the conclusion of the fiscal year that is 10 years after the date of enactment of the Frank R. Launenberg Chemical Safety for the 21st Century Act unless otherwise reauthorized or modified by Congress.”; and

(3) by adding at the end the following:

“(h) SCIENTIFIC STANDARDS.—In carrying out sections 4, 5, and 6, to the extent that the Administrator makes a decision based on science, the Administrator shall use scientific information, technical procedures, measures, methods, protocols, methodologies, or models, employed in a manner consistent with the best available science, and shall consider as applicable—

“(1) the extent to which the scientific information, technical procedures, measures, methods, protocols, methodologies, or models employed to generate the information are reasonable for and consistent with the intended use of the information;

“(2) the extent to which the information is relevant for the Administrator’s use in making a decision about a chemical substance or mixture;
“(3) the degree of clarity and completeness with which the data, assumptions, methods, quality assurance, and analyses employed to generate the information are documented;

“(4) the extent to which the variability and uncertainty in the information, or in the procedures, measures, methods, protocols, methodologies, or models, are evaluated and characterized; and

“(5) the extent of independent verification or peer review of the information or of the procedures, measures, methods, protocols, methodologies, or models.

“(i) WEIGHT OF SCIENTIFIC EVIDENCE.—The Administrator shall make decisions under sections 4, 5, and 6 based on the weight of the scientific evidence.

“(j) AVAILABILITY OF INFORMATION.—Subject to section 14, the Administrator shall make available to the public—

“(1) all notices, determinations, findings, rules, consent agreements, and orders of the Administrator under this title;

“(2) any information required to be provided to the Administrator under section 4;

“(3) a nontechnical summary of each risk evaluation conducted under section 6(b);
“(4) a list of the studies considered by the Administrator in carrying out each such risk evaluation, along with the results of those studies; and

“(5) each designation of a chemical substance under section 6(b), along with an identification of the information, analysis, and basis used to make the designations.

“(k) Reasonably Available Information.—In carrying out sections 4, 5, and 6, the Administrator shall take into consideration information relating to a chemical substance or mixture, including hazard and exposure information, under the conditions of use, that is reasonably available to the Administrator.

“(l) Policies, Procedures, and Guidance.—

“(1) Development.—Not later than 2 years after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, the Administrator shall develop any policies, procedures, and guidance the Administrator determines are necessary to carry out the amendments to this Act made by the Frank R. Lautenberg Chemical Safety for the 21st Century Act.

“(2) Review.—Not later than 5 years after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, and not
less frequently than once every 5 years thereafter, the Administrator shall—

“(A) review the adequacy of the policies, procedures, and guidance developed under paragraph (1), including with respect to animal, nonanimal, and epidemiological test methods and procedures for assessing and determining risk under this title; and

“(B) revise such policies, procedures, and guidance as the Administrator determines necessary to reflect new scientific developments or understandings.

“(3) Testing of Chemical Substances and Mixtures.—The policies, procedures, and guidance developed under paragraph (1) applicable to testing chemical substances and mixtures shall—

“(A) address how and when the exposure level or exposure potential of a chemical substance or mixture would factor into decisions to require new testing, subject to the condition that the Administrator shall not interpret the lack of exposure information as a lack of exposure or exposure potential; and

“(B) describe the manner in which the Administrator will determine that additional infor-
information is necessary to carry out this title, includ-
ing information relating to potentially ex-
posed or susceptible populations.

“(4) CHEMICAL SUBSTANCES WITH COMPLETED RISK ASSESSMENTS.—With respect to a chemical substance listed in the 2014 update to the TSCA Work Plan for Chemical Assessments for which the Administrator has published a completed risk assessment prior to the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, the Administrator may publish proposed and final rules under section 6(a) that are consistent with the scope of the completed risk assessment for the chemical substance and consistent with other applicable requirements of section 6.

“(5) GUIDANCE.—Not later than 1 year after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, the Administrator shall develop guidance to assist interested persons in developing and submitting draft risk evaluations which shall be considered by the Administrator. The guidance shall, at a minimum, address the quality of the information submitted and the process to be followed in developing draft risk evaluations for consideration by the Administrator.
“(m) REPORT TO CONGRESS.—

“(1) INITIAL REPORT.—Not later than 6 months after the date of enactment of the Frank R. Lautenberg Chemical Safety for the 21st Century Act, the Administrator shall submit to the Committees on Energy and Commerce and Appropriations of the House of Representatives and the Committees on Environment and Public Works and Appropriations of the Senate a report containing an estimation of—

“(A) the capacity of the Environmental Protection Agency to conduct and publish risk evaluations under section 6(b)(4)(C)(i) and(ii), and the resources necessary to conduct the minimum number of risk evaluations required under section 6(b)(2);

“(B) the capacity of the Environmental Protection Agency to conduct and publish risk evaluations under section 6(b)(4)(A)(ii), the likely demand for such risk evaluations, and the anticipated schedule for accommodating that demand;

“(C) the capacity of the Environmental Protection Agency to promulgate rules under section 6(a) as required based on risk evalua-
tions conducted and published under section 6(b); and

“(D) the actual and anticipated efforts of the Environmental Protection Agency to increase the Agency’s capacity to conduct and publish risk evaluations under section 6(b).

“(2) SUBSEQUENT REPORTS.—The Administrator shall update and resubmit the report described in paragraph (1) not less frequently than once every 5 years.

“(n) ANNUAL PLAN.—

“(1) IN GENERAL.—The Administrator shall inform the public regarding the schedule and the resources necessary for the completion of each risk evaluation as soon as practicable after initiating the risk evaluation.

“(2) PUBLICATION OF PLAN.—At the beginning of each calendar year, the Administrator shall publish an annual plan that—

“(A) identifies the chemical substances for which risk evaluations are expected to be initiated or completed that year and the resources necessary for their completion;
“(B) describes the status of each risk evaluation that has been initiated but not yet completed; and

“(C) if the schedule for completion of a risk evaluation has changed, includes an updated schedule for that risk evaluation.

“(o) CONSULTATION WITH SCIENCE ADVISORY COMMITTEE ON CHEMICALS.—

“(1) ESTABLISHMENT.—Not later than 1 year after the date of enactment of the Frank R. Launtenberg Chemical Safety for the 21st Century Act, the Administrator shall establish an advisory committee, to be known as the Science Advisory Committee on Chemicals (referred to in this subsection as the ‘Committee’).

“(2) PURPOSE.—The purpose of the Committee shall be to provide independent advice and expert consultation, at the request of the Administrator, with respect to the scientific and technical aspects of issues relating to the implementation of this title.

“(3) COMPOSITION.—The Committee shall be composed of representatives of such science, government, labor, public health, public interest, animal protection, industry, and other groups as the Administrator determines to be advisable, including rep-
representatives that have specific scientific expertise in
the relationship of chemical exposures to women,
children, and other potentially exposed or susceptible
subpopulations.

“(4) SCHEDULE.—The Administrator shall con-
vene the Committee in accordance with such sched-
ule as the Administrator determines to be appro-
priate, but not less frequently than once every 2
years.

“(p) PRIOR ACTIONS.—

“(1) RULES, ORDERS, AND EXEMPTIONS.—
Nothing in the Frank R. Lautenberg Chemical Saf-
ety for the 21st Century Act eliminates, modifies, or
withdraws any rule promulgated, order issued, or ex-
emption established pursuant to this Act before the
date of enactment of the Frank R. Lautenberg

“(2) PRIOR-INITIATED EVALUATIONS.—Nothing
in this Act prevents the Administrator from initi-
ating a risk evaluation regarding a chemical sub-
stance, or from continuing or completing such risk
evaluation, prior to the effective date of the policies,
procedures, and guidance required to be developed
by the Administrator pursuant to the amendments
made by the Frank R. Lautenberg Chemical Safety

“(3) ACTIONS COMPLETED PRIOR TO COMPLE-
TION OF POLICIES, PROCEDURES, AND GUIDANCE.—
Nothing in this Act requires the Administrator to re-
vise or withdraw a completed risk evaluation, deter-
mination, or rule under this Act solely because the
action was completed prior to the development of a
policy, procedure, or guidance pursuant to the
amendments made by the Frank R. Lautenberg
Chemical Safety for the 21st Century Act.”.

SEC. 18. STATE PROGRAMS.

Section 28 of the Toxic Substances Control Act (15
U.S.C. 2627) is amended by striking subsections (c) and
(d).

SEC. 19. CONFORMING AMENDMENTS.

(a) TABLE OF CONTENTS.—The table of contents in
section 1 of the Toxic Substances Control Act is amend-
ed—

(1) by striking the item relating to section 6

and inserting the following:

“Sec. 6. Prioritization, risk evaluation, and regulation of chemical substances
and mixtures.”;

(2) by striking the item relating to section 10

and inserting the following:

“Sec. 10. Research, development, collection, dissemination, and utilization of in-
formation.”;
(3) by striking the item relating to section 14
and inserting the following:

“Sec. 14. Confidential information.”; and

(4) by striking the item relating to section 25.

(b) SECTION 2.—Section 2(b)(1) of the Toxic Sub-
stances Control Act (15 U.S.C. 2601(b)(1)) is amended
by striking “data” both places it appears and inserting
“information”.

c) SECTION 3.—Section 3 of the Toxic Substances
Control Act (15 U.S.C. 2602) is amended—

(1) in paragraph (8) (as redesignated by section
3 of this Act), by striking “data” and inserting “in-
formation”; and

(2) in paragraph (15) (as redesignated by sec-
section 3 of this Act)—

(A) by striking “standards” and inserting
“protocols and methodologies”;

(B) by striking “test data” both places it
appears and inserting “information”; and

(C) by striking “data” each place it ap-
pears and inserting “information”.

d) SECTION 4.—Section 4 of the Toxic Substances
Control Act (15 U.S.C. 2603) is amended—

(1) in subsection (b)—

(A) in paragraph (1)—
(i) in the paragraph heading, by adding “, ORDER, OR CONSENT AGREEMENT” at the end; and

(ii) by striking “rule” each place it appears and inserting “rule, order, or consent agreement”;

(B) in paragraph (2)(B), by striking “rules” and inserting “rules, orders, and consent agreements”;

(C) in paragraph (3)(A), by striking “rule” and inserting “rule or order”; and

(D) in paragraph (4)—

(i) by striking “rule under subsection (a)” each place it appears and inserting “rule, order, or consent agreement under subsection (a)”;

(ii) by striking “repeals the rule” each place it appears and inserting “repeals the rule or order or modifies the consent agreement to terminate the requirement”; and

(iii) by striking “repeals the application of the rule” and inserting “repeals or modifies the application of the rule, order, or consent agreement”;}
(2) in subsection (c)—

(A) in paragraph (1), by striking “rule” and inserting “rule or order”;

(B) in paragraph (2)—

(i) in subparagraph (A), by striking “a rule under subsection (a) or for which data is being developed pursuant to such a rule” and inserting “a rule, order, or consent agreement under subsection (a) or for which information is being developed pursuant to such a rule, order, or consent agreement”;

(ii) in subparagraph (B), by striking “such rule or which is being developed pursuant to such rule” and inserting “such rule, order, or consent agreement or which is being developed pursuant to such rule, order, or consent agreement”; and

(iii) in the matter following subparagraph (B), by striking “the rule” and inserting “the rule or order”; 

(C) in paragraph (3)(B)(i), by striking “rule promulgated” and inserting “rule, order, or consent agreement”; and

(D) in paragraph (4)—
(i) by striking “rule promulgated” each place it appears and inserting “rule, order, or consent agreement”; 

(ii) by striking “such rule” each place it appears and inserting “such rule, order, or consent agreement”; and 

(iii) in subparagraph (B), by striking “the rule” and inserting “the rule or order”; 

(3) in subsection (d), by striking “rule” and inserting “rule, order, or consent agreement”; and 

(4) in subsection (g), by striking “rule” and inserting “rule, order, or consent agreement”.

(e) Section 5.—Section 5 of the Toxic Substances Control Act (15 U.S.C. 2604) is amended—

(1) in subsection (b)—

(A) in paragraph (1)(A)—

(i) by striking “rule promulgated” and inserting “rule, order, or consent agreement”; and 

(ii) by striking “such rule” and inserting “such rule, order, or consent agreement”;
(B) in paragraph (1)(B), by striking “rule promulgated” and inserting “rule or order”; and

(C) in paragraph (2)(A)(ii), by striking “rule promulgated” and inserting “rule, order, or consent agreement”; and

(2) in subsection (d)(2)(C), by striking “rule” and inserting “rule, order, or consent agreement”.

(f) SECTION 7.—Section 7(a) of the Toxic Substances Control Act (15 U.S.C. 2606(a)) is amended—

(1) in paragraph (1), in the matter following subparagraph (C), by striking “a rule under section 4, 5, 6, or title IV or an order under section 5 or title IV” and inserting “a determination under section 5 or 6, a rule under section 4, 5, or 6 or title IV, an order under section 4, 5, or 6 or title IV, or a consent agreement under section 4”; and

(2) in paragraph (2), by striking “subsection 6(d)(2)(A)(i)” and inserting “section 6(d)(3)(A)(i)”.

(g) SECTION 8.—Section 8(a) of the Toxic Substances Control Act (15 U.S.C. 2607(a)) is amended—

(1) in paragraph (2)(E), by striking “data” and inserting “information”; and

(2) in paragraph (3)(A)(ii)(I), by striking “or an order in effect under section 5(e)” and inserting
(h) SECTION 9.—Section 9 of the Toxic Substances Control Act (15 U.S.C. 2608) is amended—

(1) in subsection (a), by striking “section 6” each place it appears and inserting “section 6(a)”; and

(2) in subsection (d), by striking “Health, Education, and Welfare” and inserting “Health and Human Services”.

(i) SECTION 10.—Section 10 of the Toxic Substances Control Act (15 U.S.C. 2609) is amended—

(1) in the section heading, by striking “DATA” and inserting “INFORMATION”;

(2) by striking “Health, Education, and Welfare” each place it appears and inserting “Health and Human Services”;

(3) in subsection (b)—

(A) in the subsection heading, by striking “DATA” and inserting “INFORMATION”;

(B) by striking “data” and inserting “information” in paragraph (1);

(C) by striking “data” and inserting “information” in paragraph (2)(A); and
(D) by striking “a data” and inserting “an information” in paragraph (2)(B); and
(4) in subsection (g), by striking “data” and inserting “information”.

(j) SECTION 11.—Section 11(b)(2) of the Toxic Substances Control Act (15 U.S.C. 2610(b)(2)) is amended—
(1) by striking “data” each place it appears and inserting “information”; and
(2) in subparagraph (E), by striking “rule promulgated” and inserting “rule promulgated, order issued, or consent agreement entered into”.

(k) SECTION 12.—Section 12(b)(1) of the Toxic Substances Control Act (15 U.S.C. 2611(b)(1)) is amended by striking “data” both places it appears and inserting “information”.

(l) SECTION 15.—Section 15(1) of the Toxic Substances Control Act (15 U.S.C. 2614(1)) is amended by striking “(A) any rule” and all that follows through “or (D)” and inserting “any requirement of this title or any rule promulgated, order issued, or consent agreement entered into under this title, or”.

(m) SECTION 19.—Section 19 of the Toxic Substances Control Act (15 U.S.C. 2618) is amended—
(1) in subsection (a)—
(A) in paragraph (1)(A)—
(i) by striking “Not later than 60 days after the date of the promulgation of a rule under section 4(a), 5(a)(2), 5(b)(4), 6(a), 6(e), or 8, or under title II or IV” and inserting “Except as otherwise provided in this title, not later than 60 days after the date on which a rule is promulgated under this title, title II, or title IV, or the date on which an order is issued under section 4, 5(e), 5(f), or 6(i)(1),”;

(ii) by striking “such rule” and inserting “such rule or order”; and

(iii) by striking “such a rule” and inserting “such a rule or order”;

(B) in paragraph (1)(B)—

(i) by striking “Courts” and inserting “Except as otherwise provided in this title, courts”; and

(ii) by striking “subparagraph (A) or (B) of section 6(b)(1)” and inserting “this title, other than an order under section 4, 5(e), 5(f), or 6(i)(1),”; and

(C) in paragraph (2)—

(i) by striking “rulemaking record” and inserting “record”; and
(ii) by striking “based the rule” and
inserting “based the rule or order”;

(2) in subsection (b)—

(A) by striking “review a rule” and insert-
ing “review a rule, or an order under section 4,
5(e), 5(f), or 6(i)(1),”;

(B) by striking “such rule” and inserting
“such rule or order”;

(C) by striking “the rule” and inserting
“the rule or order”;

(D) by striking “new rule” each place it
appears and inserting “new rule or order”; and

(E) by striking “modified rule” and insert-
ing “modified rule or order”; and

(3) in subsection (c)—

(A) in paragraph (1)—

(i) in subparagraph (A)—

(I) by striking “a rule” and in-
serting “a rule or order”; and

(II) by striking “such rule” and
inserting “such rule or order”;

(ii) in subparagraph (B)—

(I) in the matter preceding clause

(i), by striking “a rule” and inserting

“a rule or order”;
(II) by amending clause (i) to read as follows:

“(i) in the case of review of—

“(I) a rule under section 4(a), 5(b)(4), 6(a) (including review of the associated determination under section 6(b)(4)(A)), or 6(e), the standard for review prescribed by paragraph (2)(E) of such section 706 shall not apply and the court shall hold unlawful and set aside such rule if the court finds that the rule is not supported by substantial evidence in the rule-making record taken as a whole; and

“(II) an order under section 4, 5(e), 5(f), or 6(i)(1), the standard for review prescribed by paragraph (2)(E) of such section 706 shall not apply and the court shall hold unlawful and set aside such order if the court finds that the order is not supported by substantial evidence in the record taken as a whole; and”;

(III) by striking clauses (ii) and (iii) and the matter after clause (iii) and inserting the following:

“(ii) the court may not review the contents and adequacy of any statement of basis and purpose required by section 553(e) of title 5, United States
Code, to be incorporated in the rule or order, except as part of the record, taken as a whole.”; and

(iii) by striking subparagraph (C);

and

(B) in paragraph (2), by striking “any rule” and inserting “any rule or order”.

(n) SECTION 20.—Section 20(a)(1) of the Toxic Substances Control Act (15 U.S.C. 2619(a)(1)) is amended by striking “order issued under section 5” and inserting “order issued under section 4 or 5”.

(o) SECTION 21.—Section 21 of the Toxic Substances Control Act (15 U.S.C. 2620) is amended—

(1) in subsection (a), by striking “order under section 5(e) or (6)(b)(2)” and inserting “order under section 4 or 5(e) or (f)”;

(2) in subsection (b)—

(A) in paragraph (1), by striking “order under section 5(e), 6(b)(1)(A), or 6(b)(1)(B)” and inserting “order under section 4 or 5(e) or (f)”;

(B) in paragraph (4)(B)—

(i) in the matter preceding clause (i), by striking “order under section 5(e) or 6(b)(2)” and inserting “order under section 4 or 5(e) or (f)”;
(ii) in clause (i), by striking “order under section 5(e)” and inserting “order under section 4 or 5(e) or (f)”; and

(iii) in clause (ii), by striking “section 6 or 8 or an order under section 6(b)(2)” and inserting “section 6(a) or 8 or an order under section 5(f)”.

(p) SECTION 24.—Section 24(b)(2)(B) of the Toxic Substances Control Act (15 U.S.C. 2623(b)(2)(B)) is amended—

(1) by inserting “and” at the end of clause (i);

(2) by striking clause (ii); and

(3) by redesignating clause (iii) as clause (ii).

(q) SECTION 26.—Section 26 of the Toxic Substances Control Act (15 U.S.C. 2625) is amended—

(1) in subsection (e), by striking “Health, Education, and Welfare” each place it appears and inserting “Health and Human Services”; and

(2) in subsection (g)(1), by striking “data” and inserting “information”.

(r) SECTION 27.—Section 27(a) of the Toxic Substances Control Act (15 U.S.C. 2626(a)) is amended—

(1) by striking “Health, Education, and Welfare” and inserting “Health and Human Services”;
(2) by striking “test data” both places it appears and inserting “information”;
(3) by striking “rules promulgated” and inserting “rules, orders, or consent agreements”; and
(4) by striking “standards” and inserting “protocols and methodologies”.

(s) SECTION 30.—Section 30(2) of the Toxic Substances Control Act (15 U.S.C. 2629(2)) is amended by striking “rule” and inserting “rule, order, or consent agreement”.

SEC. 20. NO RETROACTIVITY.

Nothing in sections 1 through 19, or the amendments made by sections 1 through 19, shall be interpreted to apply retroactively to any State, Federal, or maritime legal action filed before the date of enactment of this Act.

SEC. 21. TREVOR’S LAW.

(a) PURPOSES.—The purposes of this section are—

(1) to provide the appropriate Federal agencies with the authority to help conduct investigations into potential cancer clusters;
(2) to ensure that Federal agencies have the authority to undertake actions to help address cancer clusters and factors that may contribute to the creation of potential cancer clusters; and
(3) to enable Federal agencies to coordinate with other Federal, State, and local agencies, institutes of higher education, and the public in investigating and addressing cancer clusters.

(b) Designation and Investigation of Potential Cancer Clusters.—Part P of title III of the Public Health Service Act (42 U.S.C. 280g et seq.) is amended by adding at the end the following:

“SEC. 399V–6. DESIGNATION AND INVESTIGATION OF POTENTIAL CANCER CLUSTERS.

“(a) DEFINITIONS.—In this section:

“(1) CANCER CLUSTER.—The term ‘cancer cluster’ means the incidence of a particular cancer within a population group, a geographical area, and a period of time that is greater than expected for such group, area, and period.

“(2) PARTICULAR CANCER.—The term ‘particular cancer’ means one specific type of cancer or a type of cancers scientifically proven to have the same cause.

“(3) POPULATION GROUP.—The term ‘population group’ means a group, for purposes of calculating cancer rates, defined by factors such as race, ethnicity, age, or gender.
“(b) CRITERIA FOR DESIGNATION OF POTENTIAL CANCER CLUSTERS.—

“(1) DEVELOPMENT OF CRITERIA.—The Secretary shall develop criteria for the designation of potential cancer clusters.

“(2) REQUIREMENTS.—The criteria developed under paragraph (1) shall consider, as appropriate—

“(A) a standard for cancer cluster identification and reporting protocols used to determine when cancer incidence is greater than would be typically observed;

“(B) scientific screening standards that ensure that a cluster of a particular cancer involves the same type of cancer, or types of cancers;

“(C) the population in which the cluster of a particular cancer occurs by factors such as race, ethnicity, age, and gender, for purposes of calculating cancer rates;

“(D) the boundaries of a geographic area in which a cluster of a particular cancer occurs so as not to create or obscure a potential cluster by selection of a specific area; and
“(E) the time period over which the number of cases of a particular cancer, or the calculation of an expected number of cases, occurs.

“(c) GUIDELINES FOR INVESTIGATION OF POTENTIAL CANCER CLUSTERS.—The Secretary, in consultation with the Council of State and Territorial Epidemiologists and representatives of State and local health departments, shall develop, publish, and periodically update guidelines for investigating potential cancer clusters. The guidelines shall—

“(1) recommend that investigations of cancer clusters—

“(A) use the criteria developed under subsection (b);

“(B) use the best available science; and

“(C) rely on a weight of the scientific evidence;

“(2) provide standardized methods of reviewing and categorizing data, including from health surveillance systems and reports of potential cancer clusters; and

“(3) provide guidance for using appropriate epidemiological and other approaches for investigations.

“(d) INVESTIGATION OF CANCER CLUSTERS.—
“(1) **SECRETARY DISCRETION.**—The Secretary—

“(A) in consultation with representatives of
the relevant State and local health departments,
shall consider whether it is appropriate to con-
duct an investigation of a potential cancer clus-
ter; and

“(B) in conducting investigations shall
have the discretion to prioritize certain poten-
tial cancer clusters, based on the availability of
resources.

“(2) **COORDINATION.**—In investigating poten-
tial cancer clusters, the Secretary shall coordinate
with agencies within the Department of Health and
Human Services and other Federal agencies, such as
the Environmental Protection Agency.

“(3) **BIOMONITORING.**—In investigating poten-
tial cancer clusters, the Secretary shall rely on all
appropriate biomonitoring information collected
under other Federal programs, such as the National
Health and Nutrition Examination Survey. The Sec-
retary may provide technical assistance for relevant
biomonitoring studies of other Federal agencies.

“(e) **DUTIES.**—The Secretary shall—
“(1) ensure that appropriate staff of agencies within the Department of Health and Human Services are prepared to provide timely assistance, to the extent practicable, upon receiving a request to investigate a potential cancer cluster from a State or local health authority;

“(2) maintain staff expertise in epidemiology, toxicology, data analysis, environmental health and cancer surveillance, exposure assessment, pediatric health, pollution control, community outreach, health education, laboratory sampling and analysis, spatial mapping, and informatics;

“(3) consult with community members as investigations into potential cancer clusters are conducted, as the Secretary determines appropriate;

“(4) collect, store, and disseminate reports on investigations of potential cancer clusters, the possible causes of such clusters, and the actions taken to address such clusters; and

“(5) provide technical assistance for investigating cancer clusters to State and local health departments through existing programs, such as the Epi-Aids program of the Centers for Disease Control and Prevention and the Assessments of Chemical
TITLE II—RURAL HEALTHCARE CONNECTIVITY

SEC. 201. SHORT TITLE.

This title may be cited as the “Rural Healthcare Connectivity Act of 2016”.

SEC. 202. TELECOMMUNICATIONS SERVICES FOR SKILLED NURSING FACILITIES.

(a) IN GENERAL.—Section 254(h)(7)(B) of the Communications Act of 1934 (47 U.S.C. 254(h)(7)(B)) is amended—

(1) in clause (vi), by striking “and” at the end;

(2) by redesignating clause (vii) as clause (viii);

(3) by inserting after clause (vi) the following:

“(vii) skilled nursing facilities (as defined in section 1819(a) of the Social Security Act (42 U.S.C. 1395i–3(a))); and”;

and

(4) in clause (viii), as redesignated, by striking “clauses (i) through (vi)” and inserting “clauses (i) through (vii)”.

(b) SAVINGS CLAUSE.—Nothing in subsection (a) shall be construed to affect the aggregate annual cap on Federal universal service support for health care providers
under section 54.675 of title 47, Code of Federal Regulations, or any successor regulation.

(c) EFFECTIVE DATE.—The amendments made by subsection (a) shall apply beginning on the date that is 180 days after the date of the enactment of this Act.