DRAFT for Review Purposes

Feasibility Assessment for Epidemiological Studies at Pease International Tradeport, Portsmouth, New Hampshire

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Contents

Summary	2
Introduction	8
Site history	9
Community concerns	11
Exposure assessment	12
Summary of literature review	14
Adult cancers and other adult diseases	14
Health effects in children	14
Sources of adverse outcome data for the Pease population	15
Sources of exposure data	17
Feasibility of an epidemiological study of children at the Pease Tradeport	18
Feasibility of an epidemiological study of adults at the Pease Tradeport	32
Feasibility of an epidemiological study of former military service and civilian workers at the former Pease Air Force base	
Other study designs and health-related endpoints	42
References	45
Tables	59
Appendix	76
Literature review	77
Description of sample size calculations	93
Other sites with PFAS-contaminated drinking water from the UCMR-3	. 100
Appendix tables	. 106

Summary

This report describes the activities and the conclusions of ATSDR's feasibility assessment of possible future drinking water epidemiological studies at the Pease International Tradeport, Portsmouth, New Hampshire ("Pease"). The drinking water at Pease was contaminated with perfluoroalkyl substances (PFAS), in particular perfluorooctane sulfonate (PFOS) and perfluorohexane sulfonate (PFHxS), from the use of aqueous film-forming foam (AFFF) at the former Pease Air Force Base. The base used AFFF for firefighting training and to extinguish flammable liquid fires. In 2015, the New Hampshire Department of Health and Human Services (NH DHHS) established a PFAS blood testing program at Pease. A total of 1,578 persons submitted a blood sample for analysis. The results from the blood testing program indicated that the exposed population had higher serum levels of PFOS and PFHxS than did the U.S. population.

In March 2016, ATSDR established a community assistance panel (CAP) as a mechanism for the community to voice its concerns and provide input on decisions concerning potential health activities at Pease. A key concern expressed by the community was the lack of information on the possible short-term and long-term health effects to children and adults exposed to the PFAS contaminants in the drinking water at Pease. Specifically, the community was concerned about cancers, elevated lipids, effects on thyroid and immune function, and developmental delays in children.

ATSDR then assessed whether epidemiological studies focusing on populations at Pease were feasible and whether such studies could answer the concerns of the community. When evaluating whether an epidemiological study would be scientifically feasible, ATSDR used three main criteria:

- 1. Meaningful and credible results a study should have sufficient validity and precision, be capable of detecting health-related effects, and be as responsive as possible to the community's questions and concerns. Ideally, a study should also be capable of detecting health-related effects, for example a 20% to 100% increase in risk with sufficient statistical power (i.e., statistical power ≥80%).
- 2. Scientific importance a study should evaluate biologically plausible diseases and other health-related endpoints (also called "effect biomarkers") and improve our understanding of possible health effects of PFAS exposures.
- 3. Public health significance a study should provide a basis for determining if PFAS exposures increase the risks for specific adverse health effects, and if so, what public health actions are necessary to reduce the risks. The study should also be relevant to other populations with similar exposures.

The feasibility assessment is guided by these three criteria and does not address considerations of financial or operational feasibility. Feasibility was also assessed in terms of whether sufficient participation (sample size) could be obtained from within the Pease community to achieve sufficient statistical power for the health-related endpoints being considered, or whether the study would need to be expanded to other communities beyond the Pease population.

ATSDR reviewed the epidemiological literature on PFAS exposures to identify the health-related endpoints that have been studied and current data gaps, in particular, for the effects of PFHxS. The literature review also was used to identify adverse effect sizes observed in the PFAS studies for PFAS serum levels similar to those found in the Pease population.

The literature review found that most information on potential health effects concerned exposures to perfluorooctanoic acid (PFOA). In particular, numerous studies have been conducted of West Virginia and Ohio residents and workers exposed to PFOA from a chemical plant (the "C8" studies) [Frisbee 2009]. Studies of other workforces also were primarily focused on PFOA exposures. The literature review found that less information was available about the potential health effects of PFOS exposures, and very little information was available on the potential health effects of exposures to PFHxS. Because the primary contaminants in the drinking water at the Pease Tradeport were PFOS and PFHxS, epidemiological studies of the Pease populations have the potential to fill key knowledge gaps and address the community's concerns.

The literature review identified many health-related endpoints evaluated in previous epidemiological studies of PFAS exposures. These included cancers, lipids, effects on thyroid and immune function, and developmental delays. They also included effects on kidney and liver function and sex hormones, and diseases such as endometriosis, ulcerative colitis and osteoporosis. Many of these health-related endpoints were also previously raised by the community and the Pease CAP.

In considering possible study designs, ATSDR focused on the methods used in previous epidemiological research of PFAS exposures. Adopting study design methods consistent with previous research would facilitate the interpretation and synthesis of findings across studies. The literature review found that most of the epidemiological studies of PFAS exposures were cross-sectional and evaluated serum PFAS measurements. Some studies also evaluated cumulative PFAS serum levels that were estimated from modeling methods. ATSDR concluded that any study of populations exposed to the PFAS-contaminated drinking water at the Pease Tradeport should be cross-sectional and evaluate measured serum PFAS measurements as well as estimated cumulative PFAS serum levels. ATSDR also concluded that methods used to evaluate health-related endpoints in the Pease Tradeport populations should be consistent with methods used in previous epidemiological research of PFAS exposures.

Potential Study Designs

A. Cross-sectional study of children

The first design is a cross-sectional study of children who were exposed to the PFAS-contaminated drinking water while attending the two day-care centers at Pease. Inclusion would be limited to children who attended the day-care centers any time before June 2014, and who would be in the age range of 4–16 years at the time the study begins. During the 2015 blood testing program at Pease, 370 children aged 1–13 years contributed blood samples. If a study were to begin in 2018, these children would be ages 4–16 years. The study would involve re-contacting these participants and obtaining new blood samples. To increase the sample size, the study would also recruit and obtain blood samples from children who attended the day-care centers at Pease, but who did not participate in the New Hampshire blood testing program. Because PFAS-contaminated drinking water exposures could occur to children in utero and during breastfeeding if the mother worked at the Pease Tradeport, the study would include these additional children if the exposures began prior to June 2014 and their ages are 4 – 16 years at the time the study begins.

A comparison group of children, who did not attend day care at the Pease Tradeport and whose parents did not work at the Pease Tradeport or have occupational exposures to PFAS, would be recruited and blood samples collected. The comparison group would be sampled from the Portsmouth public schools and selected to have similar demographics as the Pease children.

Based on the health-related endpoints included in the final study, blood samples could be used to evaluate PFAS serum levels and several biomarkers of effect, including lipids, thyroid function, kidney function, immune function, and sex hormones. The children could also be assessed for neurological endpoints such as intelligence quotient (IQ), learning problems, and attention-deficit/hyperactivity disorder (ADHD) behaviors.

Calculations were conducted assuming a sample size of 350 exposed children who attended day care at the Pease Tradeport and 175 unexposed children from the Portsmouth area who did not attend day care at the Pease Tradeport. Additional sample size calculations assumed a sample size of 500 exposed children and 250 unexposed children. The sample size calculations also assumed a simple comparison of exposed versus unexposed children. A second approach was to determine the sample sizes needed to detect effects found in other PFAS studies of children with serum PFAS levels similar to those observed in the Pease children population. For some health-related endpoints, there was insufficient information to conduct any sample size calculations.

Based on sample size considerations, health-related endpoints were grouped into three categories: 1) feasible to study, 2) possible to study (but would require a larger sample size than 350 exposed children and 175 unexposed children), and 3) not feasible to study using the Pease children population unless additional populations exposed to PFAS-contaminated drinking water from other affected communities are included in the study.

Health-related endpoints feasible to study in children at Pease

- Mean difference in lipids (total cholesterol, LDL, HDL, triglycerides)
- Mean difference in estimated glomerular filtration rate (eGFR), a measure of kidney function
- Insulin-like growth factor -1 (a measure of growth hormone deficiency)
- Overweight/Obesity

<u>Health-related endpoints that may be possible to study in children at Pease</u> (although a larger sample size from the Pease community will likely be needed)

- Mean difference in uric acid, a measure of kidney function
- Elevated total cholesterol (hypercholesterolemia)
- Elevated uric acid (hyperuricemia)
- IQ/neurobehavioral
- Thyroid function
- Sex hormones
- Asthma and atopic dermatitis (immune function)
- Rhinitis (stuffy, runny nose)
- Antibody responses to rubella, mumps and diphtheria vaccines

<u>Health-related endpoints not feasible to study using the Pease children population</u> (in order to address these health endpoints, populations from other sites beyond the Pease community with PFAS-contaminated drinking water would need to be included along with the Pease children population)

- Attention deficit/hyperactivity disorder (ADHD)
- Autism spectrum disorder
- Delayed puberty
- Thyroid disease
- Childhood cancers

To evaluate exposure-response trends, the study participants would need to be split into tertiles or quartiles based on their serum PFAS levels. This might require a larger sample size for some of the health-related endpoints listed as feasible to study.

B. Cross-sectional study of adults

The second cross-sectional study design would involve obtaining blood samples from adults aged ≥ 18 years who worked anytime at the Pease Tradeport during January 2008–May 2014. This study would evaluate PFAS serum levels, lipids, thyroid function, liver function, kidney function, and immune function. The study would also evaluate diseases such as kidney disease, liver disease, cardiovascular disease, thyroid disease, ulcerative colitis, rheumatoid arthritis, osteoporosis, osteoarthritis, and endometriosis. In the 2015 blood testing program at Pease, 1,182 adults aged ≥ 18 years participated, and 1,083 (91.6%) adults reported that they last worked at Pease during 2008–2014.

Calculations were conducted assuming a sample size of 1,500 adults exposed while employed at the Pease Tradeport and 1,500 unexposed adults from the Portsmouth area who never worked at the Pease Tradeport. The sample size calculations also assumed a simple comparison of exposed versus unexposed adults. A second approach was to determine the sample sizes needed to detect effects found in other PFAS studies of adults with serum PFAS levels similar to those observed in the Pease adult population.

Based on sample size considerations, health-related endpoints were grouped into three categories: 1) feasible to study, 2) possible to study (but would require a larger sample size than 1,500 exposed and 1,500 unexposed adults), and 3) not feasible to study using the Pease adult population unless additional populations exposed to PFAS-contaminated drinking water are included in the study.

Health-related endpoints feasible to study in adults at Pease

- Mean difference in lipids (total cholesterol, LDL, HDL, triglycerides)
- Elevated total cholesterol (hypercholesterolemia)
- Mean difference in uric acid, a measure of kidney function
- Elevated uric acid (hyperuricemia)
- Thyroid disease (unconfirmed)
- Cardiovascular disease
- Hypertension
- Osteoarthritis and osteoporosis

• Mean differences in serum immunoglobin (IgA, IgE, IgG, IgM), and C-reactive protein (an indicator of inflammation); increase in antinuclear antibodies (an indicator of autoimmune reaction); alterations in specific cytokines

<u>Health-related endpoints that may be possible to study in adults at Pease</u> (although a larger sample size from the Pease community may be needed)

- Liver function
- Thyroid disease (confirmed)
- Thyroid function
- Endometriosis
- Pregnancy-induced hypertension

<u>Health endpoints not feasible to study using the Pease adult population</u> (i.e., populations from other sites beyond the Pease community with PFAS-contaminated drinking water would need to be included to evaluate these health-related endpoints)

- Liver disease
- Kidney disease
- Ulcerative colitis
- Rheumatoid arthritis
- Lupus
- Multiple sclerosis
- Kidney cancer (and other adult cancers)

To evaluate exposure-response trends, the study participants would need to be split into tertiles or quartiles based on their serum PFAS levels. This might require a larger sample size for some of the health endpoints listed as feasible to study.

C. Mortality study of former military service and civilian worker personnel

A third study design that was considered would evaluate mortality and cancer incidence among former military service and civilian worker personnel at the former Pease Air Force Base and other military bases where drinking water was contaminated with PFOS and PFHxS from the use of AFFF. Comparison military bases would also need to be identified that had no PFAS-contaminated drinking water or drinking water contamination from other chemicals above the U.S. Environmental Protection Agency's maximum contaminant levels (MCLs). Personal identifier information (e.g., Social Security number, name, date of birth, sex) necessary for data linkage with the national death index and state and federal cancer registries could be obtained from the Defense Manpower Data Center.

However, based on sample size considerations, ATSDR concluded that it is not feasible to conduct a mortality or cancer incidence study that is limited to the military service and civilian workers who were

stationed or worked at the Pease Air Force Base. Such a study would require, in addition to the Pease Air Force Base populations, several thousands of exposed populations from military bases where PFAS-contaminated drinking water occurred, as well as several thousands of comparison populations from military bases that did not have drinking water contamination.

Conclusions

The feasibility assessment concluded that it is possible to evaluate some health-related endpoints if a sufficient number of children and adults from the Pease population participate. Other health-related endpoints would require larger numbers of exposed individuals and would require the inclusion of populations from other sites who were exposed to PFAS-contaminated drinking water. The feasibility assessment concluded that a third study design, a mortality and cancer incidence study of former military service and civilian worker personnel, would not be feasible solely with the population at Pease.

No single study of the Pease population will provide definitive answers to the community about whether their exposures to the PFAS-contaminated drinking water caused their health problems. All epidemiological studies of environmental exposures and health outcomes have limitations and uncertainties. Whether a study will find an association between an environmental exposure and health effects cannot be known prior to conducting the study. The ability of a study of the Pease population to provide useful information will depend to a great extent on the success of recruiting sufficient number of study participants.

The feasibility assessment is still a draft. It will be finalized once the Pease Community Assistance Panel (CAP) and the larger Pease Tradeport community have the opportunity to review and make comments on the assessment. ATSDR will then revise the assessment based on the comments received. The feasibility of successfully evaluating particular health-related endpoints (or effect biomarkers) could change depending on final study design and goals.

Introduction

This draft report describes the approach and the conclusions of the Agency for Toxic Substance and Disease Registry's (ATSDR's) feasibility assessment of possible drinking water epidemiological studies at the Pease International Tradeport ("Pease"), Portsmouth, New Hampshire. The purpose of the feasibility assessment was to determine whether epidemiological studies are reasonable to conduct at Pease and whether data exist to conduct scientifically credible epidemiological studies. This draft feasibility assessment report for possible future studies at Pease International Tradeport is being distributed to the Pease Community Assistance Panel (CAP) for members' review and input. Input from the CAP is intended to help ATSDR ensure the proposed research is relevant to community concerns. The report is a DRAFT document that may be edited based on CAP input; it is not intended to be a protocol or systematic literature review. The final study design, including sample size, the health endpoints that can be considered and the development of the study protocol itself, including the statistical analysis approach have yet to be determined. The Pease CAP will have an opportunity to review and provide input on a draft of the study design before it is finalized. The draft feasibility assessment does not represent a commitment by ATSDR to conduct research at Pease International Tradeport, given that funding and staffing to conduct the described research are not available at this time.

Three criteria were used to determine whether epidemiological studies are warranted at Pease:

- 1. **Meaningful and credible results** —a study should have sufficient validity and precision, be capable of detecting health-related effects, and be as responsive as possible to the community's questions and concerns. Ideally, a study should also be capable of detecting health-related effects, for example a 20% to 100% increase in risk with sufficient statistical power (i.e., statistical power ≥80%). To achieve sufficient validity, a study should minimize biases such as selection bias and confounding bias. Sufficient precision can be achieved by a sample size that has at least 80% statistical power to detect health-related effect sizes observed in other studies for PFAS serum levels similar to those in the Pease population.
- 2. Scientific importance a study should evaluate biologically plausible diseases and other health-related endpoints (also called "effect biomarkers") and improve our understanding of possible health effects of PFAS exposures and fill important data gaps. Evidence for the biological plausibility of a health-related endpoint can come from animal studies of PFAS exposures, information on how PFAS exposures cause adverse effects (i.e., mechanistic information), and epidemiological studies. Since PFHxS and PFOS serum levels were elevated in the Pease population compared to national data, a Pease study should focus on data gaps concerning the health effects of exposures to these chemicals. The feasibility assessment included a literature search of epidemiological studies of PFAS exposures to identify the health-related endpoints evaluated in these studies and the data gaps that exist on the health effects of PFHxS and PFOS.
- 3. **Public health significance** a study should provide a basis for determining if PFAS exposures increase the risks for specific adverse health effects, and if so, what public health actions are necessary to reduce the risks. In particular, the study should provide a basis for early medical intervention for health outcomes that are not routinely evaluated in physical exams. The study should also be relevant to other populations with similar exposures.

In addition to the above criteria, a feasibility assessment must address specific questions:

- 1. Can the study population be enumerated and selected to minimize selection bias? (Selection bias occurs when the probability of selection is related both to exposure status and to disease status.)
- 2. Is there an appropriate comparison population?
- 3. Is there a complete exposure pathway, well-defined exposed population, and ability to assign levels of exposure with adequate accuracy?
- 4. Is there justification for studying the specific health outcome(s) being considered? (e.g., is there suggestive biological evidence? A finding in a previous study?)
- 5. Can the health effect(s) be validly ascertained or measured?
- 6. Is the exposed population sufficiently large so that risks can be estimated with precision?
- 7. Can information be obtained on other risk factors that need to be taken into account?
- 8. Can a study answer the questions of concern to the Pease community?

Site history

The Pease International Tradeport is located in Portsmouth, New Hampshire. It contains over 250 companies employing more than 9,525 people. In 1993, companies began to operate at the Pease Tradeport. Two day-care centers are located at the Tradeport. One of the day-care centers estimated that about 695 children attended the center during 1996–2016. The other day-care center could not easily compile total enrollment statistics, but its capacity is 220 children, they usually enroll about 180–195 children at a time, and they have been operating for almost 7 years. As of July 2015, the estimated population of Portsmouth was 21,530 (http://www.census.gov/quickfacts/table/PST045215/3362900). According to the 2010 census, 4.7% were children younger than 5 years, 11.9% were children ages 6–17 years, 67.5% were adults ages 18–64 years, and 15.9% were adults ages 65 years and older. Additionally, 51.5% of the population were female, 91.5% were white, and 95.6% of persons ages 25 years and older were high school graduates.

The area on which the Tradeport is located was originally built in 1951 as part of the Pease Air Force Base. In October 1989, 3,465 military personnel were assigned to the base, accompanied by 4,746 dependents. The Air Force estimated that 537 civilian employees worked on-base at that time (ATSDR 1999). During 1970–1990, an average of 3,000 personnel and their families were assigned to the base at any one time. Before 1970, the base supported a maximum of 5,000 personnel (ATSDR 1999).

Three major supply wells provided drinking water to the base: the Haven, Smith, and Harrison wells. Before 1981, the wells fed directly into the distribution system so that a particular area of base would primarily receive water from the nearest well. After 1981, the water from the three wells were mixed together and treated before entering the distribution system. These same three supply wells provided drinking water to the Pease Tradeport after it opened.

In 1977, water from the base wells was found to contain trichloroethylene (TCE). Two of the three wells serving the base were contaminated. The maximum concentrations of TCE measured in the Haven and Harrison supply wells were 391 micrograms per liter (μ g/L) and 28.5 μ g/L, respectively. After the discovery of the contamination, those wells were shut down and the city of Portsmouth supplied drinking water to the base during 1977–1978. In the fall of 1978, the wells were back in operation. TCE levels in the Haven well fluctuated between 50 μ g/L and 115 μ g/L from the fall of 1978 through January

1980, then fell below 50 μ g/L, with an occasional spike above 50 μ g/L through October 1980. From November 1980 through July 1981, TCE levels averaged about 30 μ g/L, then fell to around 10 μ g/L from August 1981 through May 1983. Levels continued to decline, but did not remain consistently below the current U.S. Environmental Protection Agency (EPA) maximum contaminant level (MCL) in drinking water of 5 μ g/L until January 1986 (ATSDR 1999).

The base officially closed in October 1991, and most of the property was transferred to the Pease Development Authority (PDA). During 1993, the business and aviation industrial park began operation. The City of Portsmouth entered into a long-term lease and operation agreement with the PDA to operate and maintain the public water system serving the Tradeport.

From approximately 1970 until the base closed, aqueous film-forming foam (AFFF) was used to extinguish and prevent flammable liquid fires. AFFF was also used during firefighting training at the base. Through 2001, perfluoroalkyl substances (PFAS) were used in the manufacturing of AFFF, including perfluorooctanoic acid (PFOA), perfluorooctane sulfonate (PFOS), and perfluorohexane sulfonate (PFHxS). AFFF containing PFAS likely leached into the soil and groundwater and migrated to the three supply wells serving the Pease Tradeport. It is not known when these wells were contaminated with PFAS, but it is possible that the contamination began before the opening of the Tradeport, when the Air Force base was still in operation.

The Haven, Smith and Harrison wells have also served the Tradeport. In addition, the City of Portsmouth has the capability to supply water to the Tradeport via its main distribution system. Monthly pumping records for the three wells were provided by the City of Portsmouth, Department of Public Works. Up through 1999, the Haven well on average provided about 56% of the total water supply at the Tradeport, with the Smith well providing 44% and the Harrison well out of service. In 2000-2001, the Haven well supplied 88% of the supply and the Smith well supplied 12%. From 2003 until it was taken out of service in May 2014, the Haven well on average supplied about half the water supply. By 2006, the Harrison well was back in service and the Smith and Harrison wells together supplied on average about half of the water supply at the Tradeport. After May 2014, the Smith and Harrison wells supplied 56% of the Tradeport water supply and the City of Portsmouth provided the other 44%.

In 2009, EPA established provisional health advisory levels for PFOS and PFOA of 0.2 μ g/L and 0.4 μ g/L, respectively [US EPA 2009]. In 2013, sampling of monitoring wells at the former Pease Air Force Base fire training areas detected PFOS and PFOA above these EPA provisional health advisory levels. In May 2016, EPA established a new lifetime health advisory for PFOS and PFOA that said the combined concentrations of PFOS and PFOA in drinking water should not exceed 0.07 μ g/L [US EPA 2016a]. No drinking water health advisory level has been established for PFHxS or other PFAS chemicals. While the EPA has a lifetime health advisory for PFOS and PFOA, no federal regulatory standards for these contaminants have been issued.

In April and May 2014, the three supply wells serving the Tradeport were sampled for PFAS. In the April sampling, the Haven well had PFOS, PFOA, and PFHxS levels of 2.5 μ g/L, 0.35 μ g/L, and 0.83 μ g/L, respectively. In the May sampling, the Haven well had PFOS, PFOA, and PFHxS levels of 2.4 μ g/L, 0.32 μ g/L, and 0.96 μ g/L. Other PFASs were also detected in the Haven well. The Harrison well had much lower levels of these contaminants with maximum PFOS, PFOA, and PFHxS levels of 0.048 μ g/L, and 0.036 μ g/L, respectively. The Smith well had maximum levels of PFOS and PFHxS of 0.018 μ g/L and 0.013 μ g/L, respectively, with an estimated level of PFOA of about 0.004 μ g/L.

No samples of the Pease Tradeport distribution system for PFAS are available from the period when the Haven well was in operation. We can use a simple mixing model to estimate the PFAS levels in the distribution system, assuming that contamination concentrations are approximately uniform throughout the system. The model takes into account the pumping rates for each of the three wells, the total water demand, and the concentrations of PFAS in the wells during the April and May 2014 sampling. Using this simple approach, the estimated levels of PFOS, PFOA, and PFHxS in the Pease Tradeport distribution system in April 2014 would be approximately 1.4 μ g/L, 0.2 μ g/L, and 0.5 μ g/L, respectively.

In April 2015, the City of Portsmouth created a community advisory board (CAB) to address the PFAS contamination in the Tradeport drinking water. The CAB was established to act as a liaison between the affected community and the New Hampshire Department of Health and Human Services (NH DHHS), to represent the diverse views of the affected community, to review the blood testing conducted by NH DHHS, and to provide input into future direction of the blood testing program (CAB 2015). The CAB held 14 public meetings during May through December 1, 2015, and disbanded after issuing its final report of its activities on December 21, 2015. Among the recommendations of the CAB in its final report were the following:

- 1. Establish a community body to coordinate ongoing issues with ATSDR, NH DHHS, and the U.S. Air Force's Restoration Advisory Board at Pease and to provide an effective mechanism for communication with all persons working or cared for at the Pease Tradeport.
- 2. A new community body should, along with its partner agencies, provide health education to the public regarding environmental chemical exposures and how exposures and risks can be reduced.

In February 2016, ATSDR began recruiting community volunteers to serve as members of a Pease community assistance panel (CAP). Technical advisors who could help CAP members in reviewing the scientific information on PFAS and proposed health activities were also recruited. The purpose of the CAP was to provide a mechanism for the community to participate directly in ATSDR's health activities related to the exposures to the contaminated drinking water at the Tradeport. The CAP would provide input concerning possible health activities proposed by ATSDR. CAP members would also work with ATSDR to gather and review community health concerns, provide information on how people might have been exposed to hazardous substances, and inform ATSDR about ways to involve the community. The first public meeting of the CAP was held in May 2016 in Portsmouth. The second public meeting was held in September 2016. ATSDR has also convened monthly calls with the CAP.

Community concerns

The final report of the CAB, issued on December 21, 2015, noted that "...the lack of any definitive information regarding the possible health effects of PFC [perfluorinated compound] exposure remains a source of frustration and concern." [CAB 2015] The report concluded, "There is a great need to better understand what if any health effects might result for PFC exposure, and at what levels of exposure these risks might be manifested."

In an email sent to ATSDR in November 2015, the CAB asked that ATSDR consider the following question: "What, if any, long-term health effects, such as specific cancers, elevated blood lipids, thyroid

function, immune function and developmental delays, are associated with the PFC exposure at Pease? This question should be broken down with regard to specific populations including children, nursing/pregnant women, firefighters, and adult exposed workers." This question was reiterated at the first in-person CAP meeting in May 2016. Some CAP members, as parents, were very concerned about the health of their children who were exposed at a critical, early age of development while attending the two day-care centers at the Pease Tradeport. They noted the lack of pediatric studies associated with PFAS exposure and wanted ATSDR to consider testing the exposed children for health endpoints such as lipids. CAP members also voiced concern about the exposed adult population, especially former military service personnel and civilian workers at the former Pease Air Force Base. Concern was also expressed for firefighters who were exposed to contaminated drinking water at Pease and also directly to AFFF as part of their firefighting duties. CAP members expressed their desire for a longitudinal approach (compared to a cross-sectional approach) to evaluate short-term and long-term health conditions, including cancers.

Exposure assessment

Using the information currently available on PFAS concentrations in the supply wells during April and May 2014, supply well pumping data, the total demand in the system, and assuming that PFAS concentrations in the supply wells during the April—May 2014 sampling reflect historical concentrations (given the persistence of these chemicals in the environment), a simple but crude assessment of PFAS drinking water exposures could be conducted. However, to accurately estimate historical PFAS concentrations in the Haven, Harrison, and Smith supply wells and the distribution system they served, both during the operation of the Air Force base and the Tradeport, would require the following steps:

- 1. Obtain information on the locations and use of AFFF at the Air Force base, including accidental releases.
- 2. Model the migration of contaminants from the soil where AFFF was used or released to the groundwater and then to the supply wells.
- 3. Model the PFAS concentrations throughout the distribution system. Historical reconstruction of PFAS concentrations in the drinking water distribution system would be needed to assess exposures to service personnel and civilian employees who were at the Air Force base during its operations, and to workers and day-care attendees at the Tradeport.

Another important source of information on exposures at the Pease Tradeport was the NH DHHS PFAS blood testing program conducted during April–October 2015. A person was eligible for this program if he or she had worked at, lived on, or attended childcare at the Pease Tradeport or Pease Air Force Base, or lived in a home near the Pease Tradeport that was served by a PFAS-contaminated private well. A total of 1,578 persons volunteered to submit a blood sample for PFASs testing [NH DHHS 2016]. This was a convenience (or volunteer) sample, not a statistically based sample. Nevertheless, the testing program provided important information on the extent and magnitude of exposures to the PFAS-contaminated drinking water at the Pease Tradeport.

Table 1 shows the serum concentrations of PFOS, PFOA, PFHxS, and perfluorononanoic acid (PFNA) for the 366 children younger than 12 years at the time of testing and comparison values from studies conducted in Texas [Schecter 2012] and California (Wu 2015). Data from the National Health and

Nutrition Examination Survey (NHANES) are not available for children younger than 12 years. NHANES testing for serum PFAS was restricted to those ages 12 years and older. The California study [Wu 2015] conducted a random sample of households in northern California and obtained blood samples from 68 children ages 2−8 years for PFAS analyses during December 2007–November 2009. The parents of the children had higher education levels than the general population. The Texas study [Schecter 2012] analyzed serum samples collected from 300 children ages ≤12 years at a children's hospital during 2009. Whether the children in the Texas study were healthy or receiving treatment for illness was not reported. None of the California and Texas children were known to be exposed to PFAS-contaminated drinking water. The children in both studies were considered to be representative of general population exposures to PFAS via diet and consumer products.

Table 1 shows that the median and geometric mean serum PFHxS and PFOS levels in the Pease children (ages <12 years) are considerably higher than background median and geometric mean levels seen in the Texas and California studies. For PFOA, the Pease children have slightly higher levels than the reference group in the Texas study, but lower than in the California study. However, the comparisons with Texas and California results might not be appropriate given the difference in sampling years. Nationally, serum levels of PFOS and PFOA have been declining sharply over time. For example, in the 1999–2000 NHANES cycle, the geometric mean serum PFOA level for persons aged \geq 12 years was 5.2 $\mu g/L$. By the 2013–2014 cycle, it had declined to 1.9 $\mu g/L$. Serum PFOS declined even more sharply, from 30.4 $\mu g/L$ during the 1999–2000 cycle to 5.0 $\mu g/L$ in the 2013–2014 cycle. PFHxS also declined, but more gradually, from 2.1 $\mu g/L$ during the 1999–2000 cycle to 1.3 $\mu g/L$ in the 2013–2014 cycle. In the NHANES 2013–2014 cycle, children ages 12–19 years had geometric mean PFOA, PFOS, and PFHxS serum levels of 1.66 $\mu g/L$, 3.54 $\mu g/L$, and 1.27 $\mu g/L$, respectively. Therefore, the most appropriate PFAS comparison values for the Pease blood testing program would be serum levels obtained near in time to the Pease sampling (i.e., 2015). Such comparison values are not currently available.

Table 2 shows the serum concentrations of PFOS, PFOA, PFHxS, and PFNA for the 1,212 participants ages 12 years and older at the time of testing and comparison values from NHANES for 2013–2014 (the most recent years data are currently available). Table 2 indicates that, similar to the children at Pease, the median and geometric mean serum levels of PFHxS and PFOS among those ages ≥12 years are considerably higher than those in the NHANES 2013–2014 cycle. The median and geometric mean serum PFOA among those at Pease were also slightly elevated compared with NHANES results.

In analyses conducted by NH DHHS, geometric mean PFHxS serum levels were higher for persons who drank \geq 4 cups of water per day compared to those who drank <4 cups per day. Of all the PFAS serum levels measured, water consumption had the strongest effect on PFHxS serum levels. In particular, water consumption had the highest effect on PFHxS serum levels among persons aged \leq 19 years (β = 0.31, SE = 0.15, marginal effect = 36.4%). Geometric mean PFOS and PFOA serum levels were also higher among persons who drank \geq 4 cups of water per day compared with those who drank <4 cups per day [NH DHHS 2016]. Linear trends were observed for geometric mean serum levels of PFOS, PFOA, and PFHxS and increasing time spent at the Pease Tradeport. The trend was strongest for PFOS and PFHxS [NH DHHS 2016].

Summary of literature review

ATSDR reviewed published health studies to identify health-related endpoints that have been studied and the data gaps that exist, in particular, for the effects of PFHxS and PFOS. The literature review also was used to identify adverse effect sizes observed in the PFAS studies for PFAS serum levels similar to those found in the Pease population.

The Appendix has a listing of the epidemiological literature on PFAS exposures and adult cancers, other adult diseases, and adverse outcomes in children. Tables 3 and 4 provide a summary. In these tables, a "+" indicates that at least one study had a finding for a specific PFAS chemical that suggests an increased risk of an adverse outcome (e.g., an odds ratio [OR] or risk ratio [RR] of ≥1.20), and a "*" indicates that no study has been conducted for that PFAS chemical. In these tables, an "I" indicates that the findings from studies have not suggested an increased risk for an adverse outcome (e.g., all odds ratios or risk ratios are <1.20) but the information is too limited to conclude that there is no association between the PFAS exposure and the adverse outcome.

These tables are for illustrative purposes, to indicate where data gaps exist and therefore additional research may be needed. Tables 3 and 4, and the tables and descriptions of the studies in the appendix, should not be interpreted as implying causation or as an assessment of the weight of evidence for an association. Currently, epidemiological research on the health effects of PFAS exposures is at an early stage. This is particularly true for PFHxS in addition to PFAS chemicals other than PFOA and PFOS. However, even for PFOA and PFOS, additional research on all the health-related endpoints mentioned in these tables will be needed to provide sufficient evidence for causal assessments and to address community health concerns.

Adult cancers and other adult diseases

Based on its assessment of the epidemiological literature, ATSDR concluded that there was limited or no information concerning associations with PFAS exposures and most cancers and other adult diseases (Table 3). In particular, very few studies have evaluated PFHxS exposures and cancers and other adult diseases. Although more information is available for PFOS exposures and cancers and other adult diseases than for PFHxS exposures, the information is still very limited and therefore inadequate to determine whether PFOS exposures increase the risk for most of the adult diseases evaluated. Although more information is available on PFOA exposure, the information is still too limited to determine whether a causal association exists between PFOA and specific cancers and other adult disease. Therefore, additional research on the effects of PFHxS, PFOS, and PFOA would be needed to determine whether exposures increase the risk for many adult cancers and non-cancer diseases.

Health effects in children

There is some evidence that PFAS exposures are associated with decreased birth weight, small fetus size for gestational age, measures of intrauterine growth retardation, and preterm birth. In particular, two meta-analyses have found an overall decrease in birthweight associated with PFOA and PFOS [Verner 2015; Bach 2015]. However, the findings across studies are inconsistent for these outcomes and for other adverse birth outcomes, and few studies have evaluated PFHxS. Several studies of infants have

found that prenatal PFAS exposures affect thyroid function, but only two studies have evaluated thyroid function in older children. A few studies have found elevated uric acid with PFAS exposures, but the possibility of reverse causation cannot be ruled out. Four studies of PFAS exposures and testosterone and other sex hormones have been conducted. However, the findings have not been consistent across studies and further research is needed. Three of the studies did find that PFAS exposures decreased testosterone in boys or girls. There is some evidence from four studies that PFAS exposures might be associated with ADHD, but findings have not been consistent across studies. Evaluating the evidence for PFAS exposures and neurobehavioral outcomes is difficult for several reasons: 1) the studies used different methods to measure the outcomes, 2) studies are inconsistent in the outcomes evaluated, and 3) too few studies have been conducted. A few studies have found associations between PFAS exposures and a decline in antibody response to specific vaccines, but only two studies evaluated the same vaccine (i.e., rubella). In summary, there are considerable data gaps concerning the health effects in children of PFAS exposures. This is because of the small number of studies conducted, inconsistencies in methods and findings across studies, and limited sample sizes in some studies. As for other adverse outcomes, few studies have evaluated the effects on children of PFHxS exposures.

Sources of adverse outcome data for the Pease population

The adverse outcomes of interest for PFAS exposure that can be ascertained from the birth certificate are pregnancy-induced hypertension, diabetes, small for gestational age (SGA), low birth weight, birth weight, preterm birth, and gestational age. Although the birth certificate has a checklist for congenital anomalies, the most reliable data on birth defects are provided by population-based birth defect registries. Birth defects registries exist in 41 states, including New Hampshire. The New Hampshire Birth Conditions Program (NHBCP), based at the Geisel School of Medicine at Dartmouth College, began collecting data on births occurring in-state to New Hampshire residents in 2003 (http://www.cdc.gov/ncbddd/birthdefects/states/newhampshire.html). Data reported on 46 different birth defects are ascertained for infants aged ≤1 year are collected through active surveillance methods. Congenital hypothyroidism data can be obtained from the newborn screening program. Newborn screening for congenital hypothyroidism is conducted in every state, including New Hampshire.

The birth certificate has information on sex of the child, plurality, gestational and pre-pregnancy diabetes, previous preterm birth, parity and gravidity, cigarette smoking before and during pregnancy, principal source of payment for the delivery (a measure of socio-economic status), date of last pregnancy, date of last normal menses, date of first and last prenatal care visit and total number of prenatal care visits, race/ethnicity of the mother and father, education of the mother and father, parents' names and address, mother's marital status, labor and delivery complications, and whether the infant is being breastfed at discharge. The New Hampshire Division of Vital Records Administration collects information on births in New Hampshire from hospitals and midwives, birth certificates, and interstate exchange agreements for births occurring out-of-state to New Hampshire residents (http://www.dhhs.nh.gov/dphs/hsdm/birth/).

Mortality information is available from the National Death Index (NDI) operated by the National Center for Health Statistics (NCHS), Centers for Disease Control and Prevention. Currently, 2014 data are complete and available for searches. "Early release data" for 2015 are ≥90% complete (98% complete for New Hampshire) and also available for searches. NDI "plus" provides information on cause of death (underlying, contributing and all other causes of death listed on the death certificate) and date and state of death based on death certificate data provided by the states. The NDI has data starting from 1979.

New Hampshire death certificate data are available from the New Hampshire Division of Vital Records Administration, which collects information on deaths of New Hampshire residents and deaths occurring in New Hampshire (http://www.dhhs.nh.gov/dphs/hsdm/death/index.htm). Information on deaths of New Hampshire residents that occur out-of-state is captured through interstate exchange agreements. Information on underlying cause of death and up to 14 contributing causes of death is collected. Complete data are available approximately 24–48 months after the close of a calendar year.

Population-based cancer registries exist in all 50 states and Washington, DC. The New Hampshire State Cancer Registry (NHSCR) is a statewide, population-based cancer surveillance program that has collected incidence data on all cancer cases diagnosed or treated in the state since 1985 (http://geiselmed.dartmouth.edu/nhscr/). NHSCR, which is contracted to the Geisel School of Medicine at Dartmouth College, currently collects data from the larger hospitals in the state. NHSCR also receives case reports from physician practices, free standing radiation oncology centers, pathology laboratories and other sources. NHSCR staff assist hospitals with fewer than 100 cases per year with reporting. Through interstate data exchange agreements, NHSCR also receives case reports for New Hampshire residents who are diagnosed outside the state.

The New Hampshire Uniform Hospital Discharge Data Set (UHDDS) collects discharge data from all health care facilities in the state (acute care hospitals, specialty hospitals, freestanding hospital emergency facilities, and walk-in urgent care centers), as required by law (http://www.dhhs.nh.gov/dphs/hsdm/hospital/index.htm). Discharge data from Maine, Massachusetts, and Vermont hospitals for New Hampshire residents are included in the UHDDS via interstate data exchange agreements. The dataset includes transfers of NH residents. Chronic diseases such as asthma, chronic obstructive pulmonary disease, angina, hypertension, congestive heart failure, hypoglycemia, and diabetes are included in the UHDDS. Limitations of this dataset are that discharges are not deduplicated and one person with multiple admissions might falsely increase the number of persons hospitalized. Additionally, state law requires health care professionals to report information on chronic health conditions relating to children, infectious diseases, immunizations, and autism to NH DHHS (http://www.healthinfolaw.org/state-topics/30,67/f_topics).

To ascertain autism or ADHD reliably, a review of school special education records and medical records from providers that conduct developmental evaluations of children or provide treatment is necessary. In Portsmouth, records are available from three elementary schools (serving grades K–5), one middle school (serving grades 6–8), and one high school (serving grades 9–12). Projected enrollment for the 2016–17 school year was 988 students in the elementary schools, 516 students in the middle school, and 1,183 students in the high school (http://cityofportsmouth.com/school/FY16BudgetBooklet.pdf). In school year 2015–2016, the Portsmouth Public Schools provided special education services to 416 students. Among those students, 121 (29.1%) had an orthopedic impairment, 36 (8.7%) had a speech/language impairment, 32 (7.7%) had a developmental delay, 25 (6.0%) had autism, 17 (4.1%) had an emotional disturbance, 11 (2.6%) had some other disability, and 174 (41.8%) were classified as having a "specific learning disability."

Various studies have focused on West Virginia and Ohio residents and workers exposed to PFOA from a chemical plant (the "C8" studies) [Frisbee 2009]. In a C8 study that evaluated ADHD, affected persons were identified via questionnaire, which included a question requesting information on medications used [Stein 2011]. For chronic diseases, the C8 studies relied primarily on self-reported information from questionnaires with attempted confirmation of self-reports by obtaining medical records.

Sources of exposure data

An important source of exposure information is PFAS biomonitoring. Measuring serum levels of PFAS chemicals provides information on the amount of these chemicals that has entered the body from all sources. At Pease, 1,578 persons volunteered to submit blood samples for PFAS analyses during the NH DHHS biomonitoring program in 2015. In the C8 study, blood samples for PFAS analyses were obtained from 66,899 persons during the 13-month baseline period, 2005–2006 [Frisbee 2009]. Biomonitoring for PFAS is useful in estimating past exposures, given the long half-lives of PFOS (approximately 5.4 years) and PFHxS (approximately 8.5 years). Although biomonitoring integrates PFAS exposures from all sources, including diet and consumer products, PFAS levels in serum from populations exposed to PFAS-contaminated drinking water will mostly reflect the drinking water exposures, unless the person is or was also exposed occupationally (e.g., firefighters, PFAS manufacturing workers).

The use of PFAS biomonitoring in epidemiological studies has some limitations. A key limitation is the issue of "reverse causation," in which the disease under investigation (e.g., kidney disease or kidney function) affects the elimination of PFAS in the body, causing higher serum levels of PFAS. Other problems include potential confounding by a factor that is both a risk factor for the disease of interest and a factor influencing serum PFAS levels (e.g., parity in the evaluation of adverse birth outcomes). Another limitation is that biomonitoring results, by themselves, might not provide sufficient information to estimate historical exposures. Estimating historical exposures is necessary to assess cumulative exposure and to characterize periods of special vulnerability to PFAS exposures, such as prenatal or early childhood exposures.

Modeling methods are used to reconstruct historical PFAS serum levels. The results of PFAS biomonitoring can be used to validate estimates of PFAS serum levels obtained from modeling. C8 researchers have successfully used physiologically based pharmacokinetic modeling of absorption, distribution, metabolism, and excretion of PFOA in the body in conjunction with drinking water contaminant levels, estimates of water intake, and residential history to predict historical and current PFOA serum levels [Shin 2011]. Researchers have also been able to simulate PFOS serum levels using information on drinking water levels and PBPK modeling [Loccisano 2011]. Therefore, reconstruction of historical PFOS serum levels is also feasible. However, reconstruction of PFOA and PFOS serum levels is limited by various uncertainties. These include lack of accurate information on individual consumption of drinking water and length of time exposed and limited information on factors that produce inter-individual variability (e.g., gender, age) and pre-existing medical conditions (e.g., compromised renal function) [Loccisano 2011]. Nevertheless, the ability to predict serum PFOS and PFOA levels based on drinking water contamination levels can substitute for, and enhance, the information provided by PFAS biomonitoring.

Issues concerning cross-sectional study designs

Cross-sectional studies are especially suitable for assessing effect biomarkers and the prevalences of nonfatal diseases, in particular, diseases with no clear point of onset [Checkoway 2004]. However, if the cross-sectional study concurrently measures the exposure and the outcome (i.e., the disease or effect biomarker), then it might be difficult to determine whether the exposure caused the outcome or whether the outcome influenced the measured exposure level [Flanders 1992, 2016]. For example, as discussed above, the concurrent measurement of serum PFAS levels and kidney function biomarkers might raise

the question of "reverse causation" because kidney function can affect the levels of PFAS in serum. This issue can be addressed by estimating exposures based on the historical reconstruction modeling of serum PFAS levels. In addition, it might be possible to estimate exposures during critical vulnerable periods (e.g., in utero exposure) through the modeling of historical serum PFAS levels. However, the modeling of historical PFAS serum levels is subject to uncertainties and data limitations, as discussed above, and published methods are available only to model serum levels of PFOA and PFOS.

Other issues concerning cross-sectional study designs are similar to those that confront other observational study designs, such as cohort studies. These issues include: 1) the ability to clearly define, enumerate and recruit (without introducing selection bias) the exposed and comparison populations, 2) the comparability of the exposed and comparison populations on risk factors other than the PFAS exposures, 3) accurate exposure assessment, and 4) accurate measurement of effect biomarkers and ascertainment of diseases.

Based on its review of the literature, ATSDR concludes that several health-related endpoints could be considered for studies of the Pease population. It is also clear that exposures to the PFAS-contaminated drinking water have occurred in the Pease population, as documented by the observed serum PFAS levels in the NH DHHS PFAS blood testing program. Therefore, it is reasonable to conduct epidemiological studies of the Pease population. However, whether it is feasible to study a specific health-related endpoint depends to a great extent on the size of the exposed population that can be recruited into a study. The usual approach to determine the necessary size of the study population for each health-related endpoint is to conduct sample size calculations.

All epidemiological studies of environmental exposures and health outcomes have limitations and uncertainties. Whether a study will find an association between an environmental exposure and health effects cannot be known prior to conducting the study. No single study of the Pease population will provide definitive answers to the community about whether their exposures to the PFAS-contaminated drinking water caused their health problems. The ability of a study of the Pease population to provide useful information will depend to a great extent on the success of recruiting a sufficient number of study participants.

Feasibility of an epidemiological study of children at the Pease Tradeport

The first population that ATSDR considered for an epidemiological study was the children who attended the two day-care centers at the Pease Tradeport. One reason to focus on children is that they are more vulnerable to environmental exposures, in particular exposures to potential endocrine-disrupting chemicals. In addition, there is serious concern in the community about the possible health effects to children from the drinking water exposures, which was conveyed to ATSDR by the Pease CAP. Finally, a study of children who attended daycare at the Pease Tradeport is the most feasible epidemiological study to conduct. The population is less transient than an adult population and the adverse health endpoints of interest do not require as large a sample size as adult chronic conditions.

The public health significance of conducting a study of these children consists of 1) the possibility of early intervention if early signs of adverse health effects, including developmental delays, are observed and 2) the relevance of a study at Pease for other populations exposed to drinking water primarily

contaminated with PFOS and PFHxS. A study of children at Pease would have scientific importance because of key data gaps concerning PFAS exposure effects on sex hormones and on neurobehavioral, immunological, and thyroid function. Animal studies support the biological plausibility of immune effects. Animal data also suggest that PFAS might be developmental neurotoxicants that can alter cognitive function and reduce learning ability. PFAS also have endocrine-disruptive properties and could interfere with thyroid function and sex hormones. A study of children at Pease would be responsive to the community's concerns and has the potential (from the perspective of statistical power) to provide meaningful and credible results for some of the adverse outcomes of interest. However, a study limited to the population of children who attended the Pease Tradeport day-care centers would likely not be sufficiently large for some of the possible adverse outcomes of interest (e.g., higher prevalences of rare diseases or very subtle changes in biomarkers of effect that have been observed in research conducted elsewhere).

A. Study population

The population of interest could be persons who attended day care at the Pease Tradeport before June 2014 and are in the age range of 4-16 years at the start of the study. The end of the period was selected because the Haven well was taken out of service in May 2014. Because PFAS-contaminated drinking water exposures could occur to children in utero and during breastfeeding if the mother worked at the Pease Tradeport, the study would include these additional children if the exposures began prior to June 2014 and their ages are 4-16 years at the time the study begins.

The age range for the Pease children study was determined by taking into account the age ranges in previous PFAS studies and the age range appropriate for the candidate endpoints. Previous epidemiological studies of children exposed to PFAS included varying age ranges. Because of data limitations (i.e., no PFAS serum data for those aged <12 years), the studies that used NHANES data evaluated those aged 12–18 years or 12–19 years. Some of the C8 studies limited participant ages to those <12 years; other C8 studies included persons up to 18 years of age. The upper age limit for many of the Taiwan children studies of PFAS was 15 years. An age range of 4–16 years would overlap the age ranges in these studies.

The chosen age range also reflected the focus of the study (i.e., children exposed to the PFAS-contaminated drinking water while attending daycare at the Pease Tradeport). The younger age limit of 4 years was chosen because intelligence quotient (IQ) testing is available for those aged 4 years and older. (For example, the Wechsler Preschool and Primary Scale of Intelligence test has an age band of 4 years to 7 years, 7 months that overlaps the Wechsler test for those aged 6–16 years.) The Strengths and Difficulties Questionnaire (SDQ), a behavioral screening questionnaire used in a Faroes study [Oulhote 2016], a Taiwan study [Lien 2016] and a Danish study [Fei 2011] has an age range of 4 – 16 years. The upper age limit of 16 years was chosen for three reasons:

- 1. Age at puberty was a candidate endpoint and virtually all of the children in a C8 study achieved puberty by age 16 years.
- 2. The IQ and SDQ testing instruments for children have an upper age limit of 16 years.
- 3. Children aged >16 years would have been last exposed (i.e., last attended daycare) more than 10 years ago.

Table 5 provides the data on serum PFOS, PFOA, and PFHxS for the 370 children who participated in the 2015 NH DHHs testing program at Pease and who were aged 1–13 years at the time of blood draw. These children would be aged 4–16 years in 2018. The geometric mean serum PFHxS in these children was $3.80 \,\mu\text{g/L}$, approximately three times higher than the serum levels reported in the Texas [Schecter 2012] and California [Wu 2015] studies and in the NHANES data for 2013–2014.

We currently do not know how many children attended daycare at the Pease Tradeport before June 2014 and who would be in the 4–16 years age range in 2018. The Discovery Child Enrichment Center is located at the Pease Tradeport and began operation in 1994. Its yearly enrollment is approximately 149 children ages 6 weeks to 5 years. Computerized records at this day-care center start in 1996. A preliminary records search by the director of the Discovery Child Enrichment Center identified 695 children who attended the daycare during 1996–2015 and who would be aged of 6–18 years in 2018. Based on the results of this search, the number of children who attended this day care prior to June 2014 and would be between the ages of 4 and 16 years in 2018 could be within the range of 250 – 450 individuals.

The Great Bay Kids' Company is also located at the Pease Tradeport and began operation in 2010. Its annual enrollment is approximately 270 children aged \leq 12 years. Assuming that most of the children enrolled would be \leq 5 years of age, and that most of the children attend daycare for 4 years, about 300 children might have attended this daycare during the period of interest and would be aged 4–16 years in 2018.

Assuming that a minimum of about 500 children attended the two day-care centers at Pease before June 2014 and would be aged 4–16 years in 2018, and assuming a reasonable participation rate of 70%, it would be possible to recruit 350 Pease children into the study. It would also be feasible to recruit at least 175 children in the same age range from the public schools in Portsmouth, NH, who were unexposed to the PFAS-contaminated drinking water at the Pease Tradeport and whose parents did not work at the Pease Tradeport or have occupational exposures to PFAS. It is reasonable to assume that participation rates would be high because of strong interest in the community concerning the Pease Tradeport situation. Moreover, the Pease CAP members have pledged to support recruitment efforts if and when a study is to be conducted. Pease CAP members have strong ties and are active in the Portsmouth community. If the actual number of children who attended the two day-care centers prior to June 2014 and would be aged 4 – 16 years in 2018 is in the range of 650 – 750, then as many as 500 children could be recruited from the Pease population. It should also be possible to recruit at least 250 children in the same age range from the Portsmouth public schools for the unexposed group.

A sample size of 350 exposed children and 175 unexposed children would be similar to the sample sizes used in the Faroes study [Grandjean 2012, 2016] and in a C8 study of 320 exposed children [Stein 2013, 2014b]. However, the sample size of 350 exposed and 175 unexposed would be considerably smaller than most of the C8 children studies and some of the other epidemiological studies of children exposed to PFAS. Therefore, a total of 525 children, 350 exposed and 175 unexposed, should be considered a minimum sample size, and attempts should be made to recruit a higher number of exposed and unexposed children to improve the statistical power of the study.

B. Study Hypotheses

As indicated in the literature review summary, the scientific literature has little information on the health effects of exposures to PFHxS. PFHxS is a key contaminant associated with the use of AFFF for firefighting training and extinguishing flammable liquid fires. The study would be an important contribution in filling this data gap and would generate knowledge relevant to other populations exposed to drinking water contaminated by PFHxS from the use of AFFF. In addition, few studies have been conducted to evaluate possible associations between childhood exposures to PFASs and effects on thyroid function, uric acid and sex hormone levels, delays in reaching puberty, IQ, and immune function. Inconsistent findings have been observed for most of these endpoints, likely in part because of differences in exposures (e.g., drinking water and other sources, such as diet) and PFAS levels of exposure, study population differences (e.g., age differences), and differences in methods. Moreover, few studies have evaluated the same neurobehavioral or immune endpoint. The study would address these issues by using methods and evaluating health effects similar to those used in previous studies of PFAS exposures in children, in particular, methods used in the C8 studies.

Based on the literature review, the following hypotheses could be evaluated:

- 1. Higher serum levels of PFOA, PFOS, or PFHxS are associated with higher total cholesterol, low-density lipoprotein, and triglycerides, and higher prevalence of hypercholesterolemia.
- 2. Higher serum levels of PFOA, PFOS, or PFHxS are associated with differences in thyroid stimulating hormone (TSH), TT4, and TT3, and a higher prevalence of hypothyroidism.
- 3. Higher serum levels of PFOA, PFOS, or PFHxS are associated with a higher level of uric acid and a higher prevalence of hyperuricemia.
- 4. Higher serum levels of PFOA, PFOS, or PFHxS are associated with differences in testosterone, estradiol, and sex hormone-binding globulin (SHBG).
- 5. Higher serum levels of PFOA, PFOS, or PFHxS are associated with delayed puberty.
- 6. Higher serum levels of PFOA, PFOS, or PFHxS are associated with lower IQ.
- 7. Higher serum levels of PFOA, PFOS, or PFHxS are associated with ADHD behaviors and learning problems.
- 8. Higher serum levels of PFOA, PFOS, or PFHxS are associated with a higher prevalences of hypersensitivity-related outcomes (e.g., asthma, rhinitis infectious diseases).
- 9. Higher serum levels of PFOA, PFOS, or PFHxS are associated with lower antibody responses to rubella, mumps, and diphtheria vaccines.

C. Recruitment and Consent

Based on sample size calculations (see Appendix), a minimum of 350 exposed children aged 4–16 years who attended the day-care centers at Pease before June 2014 would need to be recruited. To recruit the children who participated in the blood testing program, NH DHHS would have to send letters to the

parents to ask that their child participate in the study. Additional children who were exposed to the contaminated drinking water while attending the two day-care centers could be recruited via outreach to the two day-care centers at Pease, the Portsmouth public schools, media, and community organizations in the Portsmouth area. The Pease CAP has also offered to assist in recruitment, and CAP involvement will be crucial in achieving high participation rates.

A minimum of 175 children aged 4–16 years, who were unexposed to the PFAS-contaminated drinking water at the Pease Tradeport and whose mother did not work at the Pease Tradeport (or in an occupation that involved PFAS exposure) during the pregnancy and breastfeeding of the child would be recruited from the Portsmouth, NH, public schools. Before enrollment in the study, the child's mother would be interviewed to determine whether the child is eligible for the study. Recruitment would involve outreach to the eight day-care centers in Portsmouth that were located outside the Pease Tradeport, the Portsmouth public schools, media, and community organizations. The Pease CAP has offered to help with the recruitment effort. The total enrollment of Portsmouth's elementary, middle, and high schools is projected to be 2,687 in 2016–17. To encourage participation of exposed and unexposed children, an appropriate incentive would be provided.

The Pease blood testing program's consent form was strictly limited to the use of the participant's blood sample for PFAS analyses <u>only</u>. The participant also consented to complete a brief questionnaire at the time of blood draw concerning demographic information, time at Pease Tradeport, and consumption of drinking water. The consent form did not mention the use of the blood sample for research purposes or the possibility of re-contacting the participant for future studies. Moreover, the amount of blood drawn from the children was only sufficient for the PFAS analyses. Therefore, ATSDR cannot directly contact the participants in the Pease blood testing program to recruit them for a children's study. In addition, these participants must sign a new consent form to participate in a research study.

A parent of each child would be asked to sign a parental permission form requesting a blood sample (about 4 teaspoons or 20 mL) from the child for the analyses of PFASs and the effect biomarkers (i.e., lipids, TSH, uric acid, sex hormones, and immune function parameters). The consent form would also ask that the child be administered the Wechsler Abbreviated Scale of Intelligence (IQ) tests if aged 6 years or older or the Wechsler Preschool and Primary Scale of Intelligence for children younger than 6 years. The consent form would ask permission to access the child's school records, including special education records. The parent would be asked to sign a consent form to complete a questionnaire. Children ages 7 years and older would be asked to give their assent to participate in the study.

D. Questionnaire

The parents of the child participant could be asked to complete the questionnaire. The questionnaire could obtain demographic information, medical history of the parents and child, the child's medications, the dates the child's mother worked at the Pease Tradeport (or in other occupations involving PFAS exposures) and her reproductive history, the dates the child attended daycare at the Pease Tradeport, water consumption of the mother and child while at Pease Tradeport (including use of water for formula, juices, etc.) if applicable, bottled water consumption by the mother and child, length of time the child was breastfed, parental information (e.g., education, primary occupation, maternal age at birth of the participating child), the child's height and weight, and whether the child regularly exercises, currently smokes (and the number of cigarettes/day), or consumes alcohol (and the number of drinks/week).

Specific questions could be included in the questionnaire that address health outcomes of interest based on the final study design. For example, for ADHD, the questionnaire could ask, "Has a doctor or health professional ever told your child that your child has/had ADD or ADHD?" If the answer is "yes," a second question could ask for a list of medications being used for the condition. Parents would also be asked if the child had learning or behavioral problems, and if so, the type of problem and the treatment being used. Questions would be included for the hypersensitivity-related outcomes, asthma, atopic dermatitis (or atopic eczema), and allergies. Information on the child's vaccination history would also be requested from the parents. The parents would also be asked when the female child first began to menstruate.

E. Biomarkers of exposure and effect

The following biomarkers of lipids, thyroid function, kidney function, sex hormones, and immune function could be analyzed in the serum:

- Total cholesterol, low density lipoprotein, high density lipoprotein, total triglycerides
- Thyroxine (T4), T3, thyroid stimulating hormone (TSH)
- Uric acid, creatinine
- Testosterone, estradiol, sex hormone-binding globulin (SHBG), follicle stimulating hormone, insulin-like growth factor
- Immunoglobulin G (IgG), IgA, and IgM; antibodies to measles, mumps, rubella, tetanus, and diphtheria

Approximately 4 teaspoons of blood (20 mL) could be drawn from each participant to be analyzed for the standard panel of PFAS compounds and the effect biomarkers. An attempt would be made to obtain an 8-hour fasting blood sample. The parents could be asked how long the child fasted before the blood draw. The cut points of 50 ng/dL of total testosterone and 20 pg/mL of estradiol would be used to identify sexual maturation in boys and girls, respectively. IgG antibodies for measles, rubella, and diphtheria would be analyzed to determine vaccine responses. Allergen-specific IgE (mold, dust mites, dog, cat, cow's milk, peanut, hen's egg, and birch) could be analyzed. Serum levels of thyroid stimulating hormone (TSH) and total T4 could be analyzed separately and also used to determine clinical and subclinical hypothyroidism. Uric acid, total cholesterol, low-density and high-density lipoprotein, and triglycerides could be analyzed.

For children older than 6 years, the Wechsler Abbreviated Scale of Intelligence could be administered to the child to assess verbal IQ, performance IQ, and full-scale IQ. For children aged 4–6 years, the Wechsler Preschool and Primary Scale of Intelligence would be administered. For each child, school records, including special education records could be reviewed to identify learning problems and behavioral problems. The SDQ could be administered to parents to assess emotional, conduct, and peer relationship problems as well as problems with hyperactivity and inattention.

F. Exposure Assessment

As stated earlier, the analyses by NH DHHS of the data from the blood testing program at Pease indicated that geometric mean PFHxS serum levels were higher for persons who drank ≥4 cups of water per day than for those who drank <4 cups per day. The strongest finding was for serum PFHxS in participants aged 0–19 years and water consumption (β = 0.31, SE=0.15, marginal effect=36.4%).

Geometric mean PFOS and PFOA serum levels were also higher among those who, while at the Tradeport, drank ≥4 cups of water per day than for those who drank <4 cups per day [NH DHHS 2016]. Although these findings are based on a "convenience sample" (or a "volunteer sample," i.e., not a statistically-based sample), it is clear from these results that consumption of PFAS-contaminated drinking water at the Pease Tradeport was a complete exposure pathway.

Study participants could submit blood samples for PFAS and biomarker analyses during 2018. For those who participated in the 2015 blood testing program, these measurements would be used to assess their exposures. For those who did not participate in the 2015 blood testing program but who attended daycare at the Pease Tradeport during January 2008–May 2014, the PFAS serum levels obtained in 2018 could be used to estimate serum levels during 2015 by adjusting for PFAS elimination rates and taking into account background PFAS exposures. For those who consumed drinking water from the Pease Tradeport after the Haven well was taken out of service, the adjustment could also take into account the PFAS levels in the drinking water after May 2014. The 2015 (estimated or measured) PFAS serum levels and 2018 measured PFAS serum levels would be used in the analyses.

No water samples from the Pease Tradeport distribution system for PFAS testing are available before 2014. Using a simple mixing model that takes into account the pumping rates for each of the three wells, the total water demand, and the concentrations of PFAS in the wells during the April and May 2014 sampling, we can estimate historical PFAS levels in the distribution system, assuming that contamination concentrations are approximately uniform throughout the distribution system and assuming that the contamination was present at least from 2008 through May 2014.

To estimate serum levels of PFOA and PFOS over the child's life, the historical estimates of the drinking water contamination could be combined using PBPK modeling with information from the questionnaires on 1) the dates and length of time the child attended daycare at the Tradeport and the child's consumption of drinking water at the daycare and 2) whether the child's mother worked at the Pease Tradeport during pregnancy and during the period of breastfeeding and the length of the period when the child was breastfed. PBPK modeling estimates would also incorporate information from NHANES and from the PFAS serum levels of the unexposed comparison group to estimate background levels of PFAS in serum. For those children whose mothers worked at the Pease Tradeport, estimates of the mother's serum levels during the pregnancy and breastfeeding of the child would be needed. If the mother participated in the 2015 blood testing program at Pease, her measured PFAS serum levels could be used in the modeling. Children's serum levels from the 2015 NH DHHS Pease blood testing program and serum levels obtained for this study would be used to calibrate the PBPK models.

No human PBPK model for PFHxS is currently available. However, correlation coefficients for serum PFHxS and serum PFOS and PFOA were quite high among persons ages 2–14 years who participated in the 2015 testing (Pearson correlation for PFHxS was 0.75, and for PFOS and PFOA was 0.73). Therefore, it might be possible to predict historical serum levels of PFHxS based on historical estimates for serum PFOA and PFOS.

G. Sample Size

The sample size for the Pease children study should include at a minimum 350 exposed children. It should also include a minimum of 175 unexposed children randomly sampled from the Portsmouth public schools with frequency matching to the exposed children on age, sex, and race. This minimum sample size is based on several considerations. First, 370 children ages 1–13 years participated in the

2015 blood testing at Pease. That would be a 75% participation rate, assuming that a minimum of 500 children attended daycare at Pease and would be in that age range in 2015. It should be possible to recruit a similar percentage of the children who attended daycare at Pease. However, children who did not participate in the 2015 blood testing would have to be recruited, as well as a high percentage of those who did participate. Second, some studies conducted of PFAS exposure and children had similar or smaller sample sizes than the 350 exposed and 175 unexposed children at Pease (e.g., Zeng [2015] and Qin [2016] in Taiwan, Grandjean [2012] in the Faroes, Stein [2013] in a C8 study of neurobehavioral effects, Hoffman [2010] in a NHANES study), but had sufficient statistical power to observed findings to achieve statistical significance. Finally, sample size calculations conducted for this feasibility assessment indicated that at least some of the health-related endpoints of interest could be evaluated, with sufficient statistical power (i.e., statistical power ≥80%) to detect effects of exposure that are equal to or greater than those listed in Tables 6a and 6b as well as effects observed in other PFAS studies that occurred at PFAS serum levels similar to those in the Pease children population.

Sample size calculations were conducted using four different combinations for type 1 error (α error or false positive error) and type 2 error (β error, false negative error, or 1 – statistical power):

- 1. Type 1 error = 0.05 (corresponds to a two-tail hypothesis test using a p-value cutoff of 0.05, or a 95% confidence interval, to determine statistical significance) and a type 2 error = 0.05 (corresponding to statistical power of 95%).
- 2. Type 1 error = 0.05 and type 2 error = 0.20 (80% power).
- 3. Type 1 error = 0.10 (corresponds to a one-tail hypothesis test using a p-value cutoff of 0.05, or a 90% confidence interval, to determine statistical significance) and a type 2 error = 0.10 (90% power).
- 4. Type 1 error = 0.10 and type 2 error = 0.20 (80% power).

(Note: Setting the type 1 and type 2 errors to be equal indicates an equal concern for false negatives and false positives and could be justified from a public health perspective.)

Table 6a indicates the minimum effect sizes that can be detected with a sample size of 350 Pease children and 175 unexposed children from the Portsmouth area using the four combinations of type 1 and type 2 errors. Table 6b also includes the minimum effect sizes that can be detected with a sample size of 500 exposed and 250 unexposed. These minimum effect sizes assume a simple comparison between the exposed and unexposed children that is not adjusted for possible confounding risk factors or stratified into smaller exposure groupings (e.g., low, medium, and high exposure).

Another approach to sample size calculations that might be informative was to fix the minimum detectable effects to the effect sizes observed in previous studies for PFAS serum levels similar to those observed in the Pease population, select the type 1 and type 2 error rates, and allow the sample size to "float" instead of the minimum detectable effect. However, this approach is problematic because there are few studies of PFAS exposures and the childhood outcomes being considered for the Pease children study. In some instances, studies evaluating similar PFAS serum levels obtained very different effect sizes for the same outcome. In other instances, a study with a lower PFAS serum level obtained a higher effect size for an outcome than a study with a higher PFAS serum level. Moreover, there are no studies of children exposed to PFAS drinking water contamination as a result of AFFF use. Therefore, there is

much uncertainty about the effect size for each health-related endpoint that would be expected for PFAS serum levels observed among the Pease children.

With these caveats, the following sample size per stratum calculations use the findings from studies of PFAS-exposed children. (Note: a sample size of 500 per stratum means that the study would need 500 exposed and 500 unexposed children. If the goal is to compare an outcome by exposure quartiles, then each quartile would need 500 children. Also, a 2:1 ratio of exposed to unexposed requires a larger total sample size than a 1:1 ratio of exposed to unexposed.) Table 6c provides a summary of the sample size considerations for each health-related endpoint.

Lipids

Mean Total Cholesterol, LDL, HDL, triglycerides: In the Taiwan study of lipids (Zeng 2015), the sample size of 225 children aged 12-15 was sufficient to detect total cholesterol and LDL differences of 11-12 mg/dL for PFOA serum levels similar to Pease. Table 6 indicates that with a sample size of 350 exposed and 175 unexposed, much lower mean differences in total cholesterol could be detected with sufficient statistical power. However, the observed PFOA OR of 1.2 for hypercholesterolemia would have required a sample size of over 1,700 per stratum with a type 1 error of 0.10 and 80% power (using the prevalence of hypercholesterolemia in this study of 28.4%). Using a lower type 1 error and/or higher statistical power would require even larger sample sizes to detect an OR of 1.2 for hypercholesterolemia.

The serum levels of PFOA and PFOS among the children at Pease would put them in the first quartile (i.e., the reference level) if they had been in the C8 study (Frisbee 2010). In the lower PFOA and PFOS quartiles, the ORs for hypercholesterolemia were between 1.2 and 1.3, requiring sample sizes of 800 – 1660 per stratum with type 1 error of 0.10 and 80% power (using the prevalence of hypercholesterolemia in this study of 34.2%). The strongest findings in this study for total cholesterol were observed for the top quintile of PFOS serum levels. When the top quintile PFOS serum level was compared with the reference level, the mean difference in total cholesterol was 8.5 mg/dL and the OR for hypercholesterolemia was 1.6. Both of these findings are within the range that could be detected with sufficient statistical power in a Pease study with 350 exposed and 175 unexposed children. However, the top quintile for PFOS in the C8 study contained serum levels several times higher than serum levels in the top quintile of the Pease children.

A study using NHANES data for 1999–2008 [Geiger 2014] observed a mean difference in total cholesterol of 4.7 mg/dL for the 2nd tertile serum levels of PFOA compared with the reference level. The 2nd tertile serum levels of PFOA in this study correspond to the PFOA serum levels among children at Pease. To calculate a sample size to detect this mean difference, a standard deviation of 28 mg/dL (similar to the standard deviations for total cholesterol in the Taiwan and C8 study) was used. With type 1 error of 0.10 and 80% power, the sample size required to detect a mean difference of 4.7 mg/dL would be 439 per stratum (or with an exposed to unexposed ratio of 2, as suggested for the Pease children study, 660 exposed and 330 unexposed would be required). In the NHANES study, the 2nd tertile PFOS serum levels corresponded to the PFOS serum levels among Pease children. The mean difference in total cholesterol for this tertile was 3.4 mg/dL, which would require 630 per stratum with type 1 error of 0.10 and 80% power.

In the NHANES study, the ORs for hypercholesterolemia corresponding to serum PFOA and PFOS levels among children at Pease were 1.49 and 1.35, respectively. To detect an OR of 1.49 with type 1

error of 0.10 and 80% power would require 358 per stratum (or with an exposed to unexposed ratio of 2, 540 exposed and 270 unexposed).

Kidney function and uric acid

In a study of adolescents (aged 12–19 years) and kidney function using NHANES data for 2003–2010 [Kataria 2015], the top quartile for serum PFOA would correspond to the top quartile for serum PFOA among the Pease children. The mean difference in the estimated glomerular filtration (eGFR) for the top quartile of PFOA compared with the 1st quartile reference level was -6.6 mL/min/1.73 m², which would be in the range detectable, with sufficient statistical power, by the Pease study sample size of 350 exposed and 175 unexposed children.

In this study, the serum uric acid mean difference of 0.21 mg/dL was observed, comparing the top quartile PFOA to the reference level. To detect this difference with a type 1 error of 0.10 and 80% power would require a sample size larger than that projected for the Pease children study, i.e., 398 per stratum (or for an exposed to unexposed ratio of two, 596 exposed and 298 unexposed children).

The serum PFOS levels in the 3rd quartile of the NHANES study would correspond to the top quartile for serum PFOS among the Pease children. The mean difference in eGFR for the 3rd quartile PFOS level compared to the reference level was -7.2 mL/min/1.73 m², which would be in the range detectable with sufficient statistical power by the Pease study sample size of 350 exposed and 175 unexposed children. However, the mean difference in uric acid was 0.05 mg/dL which would require a sample size of more than 5,000 per stratum.

In a Taiwan study of uric acid [Qin 2016], the sample size of 225 children aged 12–15 years was sufficient to obtain a statistically significant OR for hyperuricemia of 1.65 for PFHxS at serum levels much lower than among the Pease children. For PFOA, the OR for hyperuricemia was 2.2 at serum levels much lower than observed among the Pease children. A sample size of 350 exposed and 175 unexposed children would be sufficient to detect this OR with sufficient statistical power.

Attention Deficit/Hyperactivity Disorder (ADHD) and other neurobehavioral endpoints

In a C8 study of ADHD (Stein 2011), the first quartile or reference level for PFOA and PFOS would correspond to the serum PFOA and PFOS levels among the children at Pease. For PFHxS, the serum levels among the children at Pease would correspond to the 3rd quartile level in the C8 study. For the 3rd quartile of PFHxS, the OR for ADHD was 1.43, and with current medications, the OR was 1.55. The prevalence of ADHD was 12.4%, and with current medications, 5.1%. To detect an OR of 1.43 with a prevalence of 12.4 %, the required sample size for a type 1 error of 0.10 and 80% power would be 829 per stratum. To detect an OR of 1.55 with a prevalence of 5.1%, the required sample size for a type 1 error of 0.10 and 80% power would be 1,179 per stratum.

In a study that used NHANES data for 1999–2004 [Hoffman 2010], the serum PFHxS levels were about half the levels among the children at Pease. For serum levels corresponding to the top quintile level among the Pease children, the OR for ADHD was 1.67 (using the regression coefficient in the logistic model). To detect this OR, a sample size of 540 per stratum would be required for type 1 error of 0.10 and 80% power. For PFOA, the serum levels corresponding to the top quintile level among the children

at Pease in the NHANES population would have an OR of 1.82 for ADHD. For this OR, the required sample size would be 390 per stratum (or 596 exposed and 298 unexposed children) for a type 1 error of 0.10 and 80% power.

For neurobehavioral outcomes other than ADHD, some of the neurobehavioral outcome studies (e.g., Stein [2013, 2014b]; Wang [2015], Lien [2016]) were also in the range of the minimum sample size suggested for the Pease children study. IQ differences in the range of 3 to 4 points could be detected with reasonable statistical power with a sample size of 350 exposed and 175 unexposed children.

One study [Liew 2015] evaluated autism spectrum disorder and obtained an OR of 1.3 for serum PFHxS. With a prevalence of about 1.5%, a sample size of several thousand children would be necessary to detect this OR. To detect an OR of 2.0 with sufficient statistical power would require sample sizes of over 1,600 exposed and 1,600 unexposed.

Sex hormones and delayed puberty

In the C8 study of sex hormones [Lopez-Espinosa 2016], the serum levels of PFOA, PFOS, and PFHxS were considerably higher than among the children at Pease. For PFOS, the natural log estradiol percent difference in boys of -4% (per interquartile range of the natural log of PFOS) would require at least 1,154 per stratum for type 1 error of 0.10 and 80% power. The strongest finding in this study was the decrease in testosterone among girls associated with PFOS. The natural log testosterone percent difference in girls was -6.6% per interquartile range of the natural log of PFOS. To detect a percent difference this large with type 1 error of 0.10 and 80% power would require at least 290 per stratum, or 434 exposed and 217 unexposed children.

There was insufficient information to make sample size calculations for the endpoint, delayed puberty. The C8 study that evaluated this endpoint in included thousands of boys and girls [Lopez-Espinosa 2011].

Growth hormone

In the C8 study that evaluated sex hormones, insulin-like growth factor-1 (IGF-1) was also evaluated [Lopez-Espinosa 2016]. The difference in the natural log IGF-1 among boys and girls was -2.5% and -2.1% per interquartile range of the natural log of PFHxS, respectively. To detect these differences with sufficient statistical power, a sample size of 350 exposed and 175 unexposed children would be sufficient.

Thyroid disease and function

A C8 study [Lopez-Espinosa 2012] evaluated thyroid disease among children. The prevalence of participant-reported thyroid disease among children in this study was very low, about 0.6% and an OR of 1.44 was obtained for PFOA serum levels considerably higher than those in the Pease population. To detect this OR with 80% statistical power would require a sample size of over 10,000 exposed children.

In the C8 study of thyroid function [Lopez-Espinosa 2012], the largest percent difference for natural log TSH was 3.1%, and 2.3% for TT₄. These percent changes were for PFOA and PFOS serum levels considerably higher than the serum levels among the children at Pease. To detect a 2.3% change in TT₄ would require a sample size of at least 850 per stratum (type 1 error = 0.10 and 80% power). To detect a 3.1% change in natural log TSH would require a sample size of at least 8,545 per stratum (type 1 error = 0.10 and 80% power).

In the Taiwan study of thyroid function [Lin 2013], the sample size for those aged 12–19 years was 212. The geometric means for serum PFOA and PFOS were lower than the geometric mean serum levels among the children at Pease. For males and females, the natural log TSH declined by 0.5 mIU/L and 0.35 mIU/L respectively, for the >90th percentile serum PFOA compared with the reference level. To detect either of these differences with sufficient statistical power, a sample size of 350 exposed and 175 unexposed children would be sufficient.

Immune function and diseases related to immune function

For immune function, one study [Grandjean 2012] had a similar sample size (N = 532) as the minimum proposed for the Pease children study (i.e., 350 exposed and 175 unexposed children), and two studies had somewhat larger sample sizes that might be achievable at Pease (Stein [2016a], N = 640; and Buser [2016], N = 637). The data reported in these studies were insufficient to conduct sample size calculations.

For asthma, the ORs observed in the NHANES studies [Humblet 2014, Stein 2016a] were in the range of 1.2 – 1.3 and would require much larger sample sizes than can be recruited at Pease to achieve sufficient statistical power. However, a Taiwan study [Dong 2013] obtained ORs for asthma between 3.8 and 4.0 for PFHxS and PFOA serum levels lower than those observed in the Pease children population. A sample size of 350 exposed and 175 unexposed would be sufficient to detect these ORs with sufficient statistical power.

Only one study [Stein 2016a] evaluated rhinitis and observed an OR of 1.35 for serum PFOA. To detect an OR this low with sufficient statistical power would require a sample size larger than could be recruited from the Pease population. However, with sufficient statistical power, ORs in the range of 1.5 – 1.6 could be detected in a study of the Pease population with a sample size of 500 exposed and 250 unexposed children. These ORs would fall within the 95% CI for the finding in this study.

Other health-related endpoints

A NHANES study [Geiger 2014b] evaluated PFOS and PFOA serum levels and hypertension and obtained ORs < 1.0. Since there is no evidence so far of an association between PFAS serum levels and hypertension in children, this endpoint is not considered further.

A study conducted in the Faroes [Karlsen 2016] evaluated serum levels of PFOA, PFOS and PFHxS and overweight/obesity in children. At age 5 years, the ORs for overweight/obesity and the third tertile serum levels of PFOA, PFOS and PFHxS were 1.88, 0.94, and 1.22. The serum levels of the PFAS chemicals were considerably lower than at Pease. An OR of 1.62 could be detected with 80% statistical power with a sample size of 350 exposed and 175 unexposed children.

Childhood cancers

For childhood cancers such as leukemias, the incidence and prevalence is very low, requiring large sample sizes. For example, the probability of getting a leukemia at \leq 15 years is 0.08% or 8 per 10,000. For ages \leq 20 years, the probability is 0.09% or 9 per 10,000. At ages \leq 14 years, the incidence rate for leukemias is 5.5 per 100,000 person-years. A study that attempted to evaluate leukemias or other childhood cancers would have to be multi-site or national.

H. Conclusion

Very little is known about the health effects from exposure to PFHxS, a PFAS that was considerably elevated in the serum of children tested at Pease. More information is available on the health effects of PFOS exposure, which was also elevated in the serum of children at Pease. However, there are still major data gaps and inconsistencies in the findings concerning the health effects of PFOS exposure, particularly effects on immune, thyroid and kidney function, neurobehavioral endpoints, sex hormones, and age at puberty. Based on sample size calculations, a study of children at Pease could have sufficient statistical power to evaluate several health-related endpoints. The study could also meet the criteria of public health significance and scientific importance, and could address some of the health concerns voiced by the Pease CAP and the previous CAB.

The study population can be enumerated and selection bias can be minimized if recruitment is carefully done to avoid selection bias (i.e., selection that is associated with exposure and disease status). A sample of Portsmouth public school students would be an appropriate comparison group for the Pease children. There is a complete exposure pathway and a well-defined exposed population. The health-related endpoints under consideration have been evaluated in at least one epidemiological study of PFAS exposures to children, and these endpoints can be measured accurately. Information on potential confounding factors can be obtained via questionnaire. The issue of reverse causation and confounding from the use of measured serum PFAS levels can be avoided by predicting serum levels using PBPK modeling. Therefore, a children's study at Pease could provide meaningful and credible results.

A key issue is whether a study limited to the children exposed at the Pease Tradeport would have sufficient statistical power and precision for some of the endpoints under consideration. A minimum sample size of 350 exposed Pease children and 175 unexposed children from the Portsmouth area would be sufficient for several outcomes of interest. For example, Table 6 indicates that a sample size of 350 exposed and 175 unexposed children is sufficient to detect effects of reasonable size for most of the endpoints listed in the table. In addition, some of the immune and neurobehavioral studies that had sufficient statistical power to obtain effect estimates that achieved statistical significance had sample sizes within the range suggested as a minimum for the Pease children study.

When the effect sizes seen in previous PFAS studies are considered, the suggested minimum sample size for the Pease children study could be sufficient for several endpoints, such as mean differences in lipids, eGFR, and IGF-1. For other outcomes, such as uric acid mean difference, the sex hormones testosterone and estradiol, and thyroid function, the sample size of a study limited to the Pease children population might not be sufficient. Based on sample size calculations assuming 350 Pease children and 175 unexposed children, and assuming a simple comparison of exposed versus unexposed, health endpoints are grouped below into three categories: 1) feasible to study, 2) possible to study (but might require a

larger sample size, e.g. 500 exposed and 250 unexposed), and 3) not feasible to study using the Pease children population, unless additional populations exposed to PFAS-contaminated drinking water are included in the study.

Health endpoints feasible to study in children at Pease

- Mean difference in lipids (total cholesterol, LDL, HDL, triglycerides)
- Mean difference in estimated glomerular filtration rate (eGFR), a measure of kidney function
- Insulin-like growth factor-1 (IGF-1, a measure of growth hormone deficiency)
- Overweight/Obesity

<u>Health endpoints that might be possible to study in children at Pease</u> (although a larger sample size may be needed)

- Mean difference in uric acid, a measure of kidney function
- Elevated total cholesterol (hypercholesterolemia)
- Elevated uric acid (hyperuricemia)
- IQ/neurobehavioral
- Thyroid function
- Sex hormones
- Asthma and atopic dermatitis (immune function)
- Rhinitis (stuffy, runny nose)
- Antibody responses to rubella, mumps, and diphtheria vaccines

<u>Health endpoints not feasible to study using the Pease children population</u> (to address these health endpoints, populations from other sites with PFAS-contaminated drinking water would need to be included, along with the Pease children population)

- Attention deficit/hyperactivity disorder (ADHD)
- Autism spectrum disorder
- Delayed puberty
- Thyroid disease
- Childhood cancers

To evaluate exposure response relationships, more than two strata are necessary. For some of the candidate outcomes that are listed above as feasible to study or possible to study, the Pease children population that can be recruited to participate will not be large enough to be split into exposure tertiles or quartiles and still have sufficient statistical power for comparisons between each of the exposure strata and a reference (unexposed) stratum.

Data analyses similar to those used in the C8 studies could be used. The methods include linear regression of continuous (untransformed and natural log transformed) effect biomarkers on continuous (untransformed and natural log transformed) PFAS serum levels and categorized PFAS serum levels, and logistic regression of categorized effect biomarkers (e.g., hypercholesterolemia) or disease prevalence on continuous (untransformed and natural log transformed) and categorical PFAS serum

levels. Restricted cubic splines for linear and logistic regression would be conducted to obtain flexible, smoothed exposure-response curves. Measured PFAS serum levels would be evaluated. In addition, for PFOS and PFOA (and possibly PFHxS, if an historical reconstruction modeling method becomes available), estimated cumulative serum levels and estimated serum levels during critical vulnerability periods (e.g., in utero exposure) could be evaluated.

In summary, a study limited to the Pease children population will likely have a sufficient sample size for some of the candidate endpoints <u>if</u> the comparisons are simply between an exposed and unexposed group. For some of the candidate endpoints, the sample size will be insufficient for even a simple comparison between an exposed an unexposed group. Moreover, for many of the candidate endpoints, the Pease children population will be of insufficient size to split into tertiles or quartiles to evaluate exposure—response trends. Therefore, the inclusion of other sites with PFAS-contaminated drinking water could be considered.

Feasibility of an epidemiological study of adults at the Pease Tradeport

Compared with NHANES data, PFHxS serum levels were elevated among adults who participated in the 2015 NH DHHS blood testing program. However, the literature review indicated that very few studies have been conducted that evaluated PFHxS exposures and adult health effects. PFOS serum levels were also elevated among the adults who participated in the NH DHHS blood testing program. Although considerably more studies found evaluated PFOS exposures and adult health effects, there remain data gaps and inconsistencies in the findings for liver function, kidney function and kidney disease, thyroid disease and thyroid function, autoimmune diseases and immune function, osteoporosis/osteoarthritis, endometriosis, and most cancers.

The public health significance of conducting a study of adults at Pease is that the study will be relevant to other adult populations exposed to drinking water primarily contaminated with PFOS and PFHxS. A study might also provide an opportunity for early medical intervention for certain health endpoints that might be associated with PFAS exposure but not evaluated in routine physical exams, such as alterations in thyroid, liver, and kidney function. A study of adults at Pease would have scientific importance because it potentially could help to fill critical data gaps mentioned above concerning the health effects of PFHxS and PFOS exposures. Based on animal studies, there is biological plausibility that PFAS exposures could result in alterations of immune function and might have endocrine-disruptive properties that could lead to alterations in thyroid function. However, few epidemiological studies have evaluated PFHxS or PFOS exposures and these health endpoints. Finally, a study of adults at Pease has the potential to provide meaningful and credible results (from the perspective of statistical power) for some of the adverse outcomes of interest and would be responsive to community concerns. However, a study limited to Pease adults would likely not be sufficiently large to associate exposures and some adverse health outcomes (e.g., rare diseases such as specific cancers and specific chronic diseases).

A. Study hypotheses

Based on the literature review, the following hypotheses could be evaluated:

1. Higher serum levels of PFOA, PFOS, or PFHxS are associated with higher total cholesterol, low-density lipoprotein and triglycerides, and a higher prevalence of hypercholesterolemia.

- 2. Higher serum levels of PFOA, PFOS, or PFHxS are associated with higher prevalences of coronary artery disease and hypertension.
- 3. Higher serum levels of PFOA, PFOS, or PFHxS are associated with differences in thyroid stimulating hormone (TSH), TT4, and TT3, and a higher prevalence of hypothyroidism.
- 4. Higher serum levels of PFOA, PFOS, or PFHxS are associated with a higher level of uric acid and a higher prevalence of hyperuricemia.
- 5. Higher serum levels of PFOA, PFOS, or PFHxS are associated with a lower estimated glomerular filtration rate (eGFR) and a higher prevalence of kidney disease.
- 6. Higher serum levels of PFOA, PFOS, or PFHxS are associated with higher levels of liver function biomarkers alanine transaminase (ALT), γ-glutamyltransferase (GGT), and direct bilirubin and a higher prevalence of liver disease.
- 7. Higher serum levels of PFOA, PFOS, or PFHxS are associated with higher prevalences of osteoarthritis and osteoporosis.
- 8. Higher serum levels of PFOA, PFOS, or PFHxS are associated with a higher prevalence of endometriosis.
- 9. Higher serum levels of PFOA, PFOS, or PFHxS are associated with higher prevalences of autoimmune diseases such as ulcerative colitis, rheumatoid arthritis, lupus, and multiple sclerosis.
- 10. Higher serum levels of PFOA, PFOS, or PFHxS are associated with differences in serum levels of IgA, IgE, IgG, IgM, C reactive protein (CRP), and antinuclear antibodies (ANA) and alterations in specific cytokines.

A study of adults could include the collection of new blood samples to analyze PFAS serum levels. The blood samples would also be analyzed for lipids and biomarkers of kidney, liver, thyroid, and immune function. A questionnaire could be used to ascertain kidney disease, liver disease, cardiovascular disease, hypertension, thyroid disease, autoimmune diseases, osteoporosis, osteoarthritis, pregnancy-induced hypertension, and endometriosis. Diseases ascertained via questionnaire would be confirmed using medical records

B. Study population

According to the census, Portsmouth has 21,530 residents. About 67.5 % are adults aged 19-64 years and another 15.9% are aged 65 years and older. This would mean that there are about 14,500 adults aged 18-64 years and about 3,400 aged 65 years and over. Although the actual number is unknown, some of the workers at the Pease Tradeport live in New Hampshire towns other than Portsmouth or in the bordering states of Massachusetts and Maine. The Pease Tradeport has a workforce of >9,000 persons. In the 2015 blood testing program at Pease, 1,182 adults aged ≥ 18 years participated. Table 5 provides PFAS serum data for the 1,190 participants in the 2015 Pease blood testing program who will be age ≥ 18 years in 2018.

C. Recruitment and consent

As stated previously, the NH DHHS Pease blood testing program's consent form was strictly limited to use of the participant's blood sample for PFAS analyses <u>only</u>. The participant also consented to complete a brief questionnaire at the time of blood draw concerning demographics, time at Pease Tradeport, whether the worker was a firefighter, and consumption of drinking water. The consent form did not mention the use of the blood sample for research purposes or the possibility of re-contacting the participant for future studies. Therefore, the blood samples were not stored for future use, and ATSDR cannot directly contact the participants in the Pease blood testing program to recruit them for a study. Adults would need to sign a new consent form to participate.

The consent form would request a blood sample (about 35 mL or 1.2 ounces) from the adult for the analyses of PFASs and the effect biomarkers. (Note: 35 mL was the maximum amount of blood obtained from adults in the C8 studies.) The consent form could also ask the participant to complete a questionnaire covering demographics, water consumption, dates and length of time working at Pease, occupational history, lifestyle and health behaviors, diseases diagnosed by a physician or other health provider, and provider contact information.

To recruit adult study participants, NH DHHS would have to contact those who participated in the 2015 blood testing program. Another approach is to work with the Tenants Association at Pease (TAP) and the Pease International Development Authority (PDA) to contact firms on their mailing lists. TAP sends newsletters and email notices to subscribing firms at the Tradeport. The PDA list, with mailing addresses and email addresses of all firms at the Pease Tradeport, was provided to ATSDR to help recruit members to the Pease CAP. This list could be used to conduct outreach to recruit adult study participants. Other methods of outreach include contacting community groups and the media.

D. Biomarkers of effect

The following biomarkers would be analyzed in the serum:

- Total cholesterol, low density lipoprotein, high density lipoprotein, total triglycerides
- Thyroxine (T4), T3, thyroid stimulating hormone (TSH)
- Uric acid, creatinine
- Alanine transaminase (ALT), γ-glutamyltransferase (GGT) and direct bilirubin
- Immunoglobulin G (IgG), IgA, IgE and IgM; C reactive protein, and antinuclear antibodies (ANA), and alterations in specific cytokines.

E. Exposure assessment

Exposure assessment could be based on the serum PFAS levels obtained in the study supplemented by the serum PFAS levels for those who participated in the 2015 NH DHHS Pease blood testing program. Using historical estimates of the PFAS contaminant levels in the drinking water at the Pease Tradeport (based on water modeling methods), PBPK modeling can be used to estimate historical serum levels of PFOA and PFOS, combining information from the questionnaire on water consumption and dates and length of time employed at Pease Tradeport, and information on background PFAS serum levels from NHANES and from a comparison group unexposed to PFAS-contaminated drinking water or occupationally exposed to PFAS or AFFF. Serum levels from the 2015 NH DHHS Pease blood testing

program and serum levels obtained for this study would be used to calibrate the PBPK models. If feasible, historical estimates of serum PFHxS can be based on historical estimates for serum PFOA and PFOS, because serum levels of PFHxS and PFOS were highly correlated among the Pease adults who participated in the 2015 blood testing program (Pearson correlation coefficient = 0.73).

F. Sample size considerations

A key problem for an adult study at Pease will be identifying an appropriate comparison population of workers from the Portsmouth area with similar occupations as the Pease workforce and who were not exposed to PFAS-contaminated drinking water or occupationally exposed to PFAS or AFFF. Another key problem will be recruiting a sufficient number of participants to achieve reasonable statistical power and precision of effect estimates.

Studies conducted of the adult C8 population included tens of thousands of participants. For example, studies of thyroid disease [Winquist 2014a], cardiovascular disease and lipids [Winquist 2014b], kidney disease [Dhingra 2016], and liver function [Darrow 2016] included 28,541 community members and 3,713 workers at the DuPont plant. Smaller studies using NHANES data (e.g., Wen [2013], Webster [2016], Shankar [2011], Gleason [2015], and Lin [2010]) had sample sizes of 1,181–4,333 adults.

Table 7a indicates the minimum detectable effects for a study that included 1,500 participants per stratum. For a simple comparison between exposed and unexposed, this would require a total of 3,000 participants, i.e., 1,500 exposed and 1,500 unexposed. If the study population were divided into quartiles of PFAS serum levels, with the first quartile being the reference exposure level, then this would result in a total sample size of 6,000 persons (i.e., 4,500 exposed and 1,500 unexposed). Four combinations of type 1 error (α error) and type 2 error (β error) are used in the table. A type 1 error of 0.05 corresponds to a two-tailed hypothesis test using a p-value cutoff of 0.05 to determine statistical significance, or using a 95% confidence interval. A type 1 error of 0.10 corresponds to a one-tail hypothesis test using a p-value cutoff of 0.05 to determine statistical significance, or using a 90% confidence interval. A type 2 errors of 0.05, 0.10, and 0.20 correspond to statistical power of 95%, 90% and 80%, respectively.

Another possible approach to sample size calculations that might be informative would be to fix the minimum detectable effects to the effect sizes observed in previous studies for similar levels of exposure, select the type 1 and type 2 error rates, and allow the sample size to "float" instead of the minimum detectable effect. However, this approach is problematic because there are few studies of PFAS exposures and the adult outcomes being considered for the Pease adult study. In some instances, studies evaluating similar PFAS serum levels obtained very different effect sizes for the same outcome. In other instances, a study with a lower PFAS serum level obtained a higher effect size for an outcome than a study with a higher PFAS serum level. Moreover, there are no studies of adults exposed to PFAS drinking water contamination as a result of AFFF use. Therefore, there is much uncertainty about the effect size for each health-related endpoint that would be expected for PFAS serum levels observed among the Pease adults. With these caveats, the following sample size per stratum calculations use the findings from studies of PFAS-exposed adults. Table 7b provides a summary of the sample size considerations for each health-related endpoint.

Lipids

In the lipid study conducted of the C8 adult population [Steenland 2009], PFOS serum levels corresponding to the PFOS serum levels among adults who participated in the Pease blood testing program would result in a 3–4 mg/dL change in total cholesterol and in LDL. Table 7a indicates that detecting a difference of about 4 mg/dL in total cholesterol would require a sample size of about 1,500 per stratum. To detect a difference of 3 mg/dL would require a larger sample size. For LDL, a sample size of 1,500 per stratum would be sufficient for mean differences in the 3–4 mg/dL range.

The predicted increase in total cholesterol at the highest decile for PFOA and PFOS in the C8 study was 11–12 mg/dL. To detect a difference of 11 mg/dL, a sample size in the range of 200–300 per stratum would probably be sufficient. However, the highest decile for PFOA and PFOS in the C8 population is considerably higher than the serum levels observed for the adult participants in the Pease 2015 blood testing.

In a C8 study [Steenland 2009] and a Canadian study [Fisher 2013], ORs in the range of 1.35 - 1.6 were observed for hypercholesterolemia. Although PFAS serum levels were higher in the C8 population than the Pease population, the PFAS serum levels in the Canadian study were lower than in the Pease population. Table 7a indicates that ORs in this range for hypercholesterolemia can be detected with sufficient statistical power with a sample size of 1,500 per stratum.

Kidney disease/function, and uric acid

In the C8 study of chronic kidney disease [Dhingra 2016], the highest hazard ratio (HR) was observed for the lowest quintile of exposure (compared with the reference level) and was equal to 1.36. To detect this HR, given the low prevalence of the disease (approximately 1.4%). would require a sample size of at least 8,600 per stratum.

In the C8 study of uric acid [Steenland 2010], serum PFOS levels that correspond to those observed among the adult participants in the Pease blood testing program resulted in a difference of 0.14 mg/dL. To detect this difference would require a sample size in the range of 1,600–2,100 per stratum.

The largest differences in uric acid observed in this study was 0.28 mg/dL for PFOA serum levels ≥188.7 ng/mL and 0.22 mg/dL for PFOS serum levels ≥40.5 ng/mL. These serum levels are considerably higher than those observed for the adults at Pease. Based on sample size calculations, a uric acid difference of 0.28 mg/dL could be detected with reasonable statistical power and a sample size in the range of 500–600 per stratum. Table 7a indicates that much lower differences in uric acid could be detected with reasonable statistical power using a sample size of 1,500 per stratum.

In the C8 study, the OR for hyperuricemia for PFOA serum levels similar to those at Pease equaled 1.02. For the top quintile of serum PFOA in the C8 population, the OR was 1.47. Based on sample size calculations, a sample size in the range of 450–600 would be sufficient to detect an OR of 1.47 with reasonable statistical power. However, the top quintile serum PFOA level in the C8 study was considerably higher than observed in the Pease population.

In a study using NHANES data [Shankar 2011], a change in uric acid of 0.40 mg/dL was observed for serum PFOA levels similar to those observed for Pease. Based on sample size calculations, this difference could be detected with reasonable statistical power using a sample size of about 300 per

stratum. For hyperuricemia, an OR of 1.90 was observed for serum PFOA levels similar to Pease. Based on sample size calculations, an OR of 1.90 can be detected with reasonable statistical power using a sample size of about 240 per stratum.

Liver function

For liver function, to detect the very subtle changes observed in the C8 studies [Gallo 2012; Darrow 2016] would require a sample size as large as the C8 study itself. The same is true for liver disease. In the Darrow 2016 study, the highest OR observed was 1.19 for the 2nd quintile of serum PFOA. The 2nd quintile of serum PFOA in the C8 study is higher than the serum levels at Pease. To detect an OR of 1.19 would require a sample size of at least 20,000 per stratum.

A study using NHANES data [Gleason 2015] was able to detect associations with uric acid and liver function biomarkers at serum PFAS levels similar to those observed at Pease and with a total sample size of 4,333 persons. This study evaluated quartiles of serum PFAS, so each stratum had a sample size of about 1,083 persons. Another study that used NHANES data [Lin 2010] also was able to detect associations with liver function biomarkers with a total sample size of 2,216 persons. This study also evaluated quartiles, so each stratum had a sample size of about 554 persons.

Cardiovascular disease

The C8 study that evaluated coronary artery disease did not find an elevation in risk [Winquist 2014b]. However, a study that used NHANES data [Shankar 2012] obtained an OR of 2.01 for cardiovascular disease for the 4th quartile PFOA serum levels. These PFOA serum levels, ≥6 ng/mL, would correspond to the 5th quintile of PFOA serum levels among Pease adults. The prevalence of cardiovascular disease in this study was 13%. To detect an OR of 2.01, a sample size of about 250/stratum would probably be sufficient.

Hypertension

One study evaluated hypertension in a community population and observed an OR <1.0 [Winquist 2014b]. The prevalence of hypertension in this study was about 38%. With a sample size of 1,500 per stratum and a prevalence of 38%, ORs between 1.21 and 1.31 could be detected with sufficient statistical power.

Thyroid disease/function

For thyroid disease, the C8 study evaluated self-reported disease and self-reported disease that was confirmed by medical records [Winquist 2014a]. For serum PFOA levels similar to those at Pease, the hazard ratios were in the range of 1.2–1.3. For all self-reported thyroid disease (prevalence = 11.3%), a sample size of about 2,100 per stratum would probably be sufficient to detect a hazard ratio of 1.3. The prevalence for confirmed disease was 6.5%, so that a sample size of about 3,500 per stratum would probably be necessary to detect an HR of 1.3.

A study that used NHANES data evaluated thyroid disease [Melzer 2010]. For confirmed thyroid disease (prevalence = 2.4% in this study), the ORs were slightly above 1.1 for PFOS and PFOA serum levels similar to those at Pease. To detect this OR would require a sample size equivalent to the C8

population. The highest OR observed was 1.89 among men in the top quartile of PFOS and PFOA. To detect this odds ratio, a sample size of about 1,400 per stratum would probably be sufficient.

The C8 study that evaluated thyroid function biomarkers [Knox 2011] observed very subtle changes that would require a study of equivalent size (52,296) to detect associations with sufficient statistical power. On the other hand, a study that used NHANES data [Wen 2013] to evaluate thyroid function observed larger changes that could be detected with a total sample size of <1,200 (or <300 per quartile stratum).

Immune function and autoimmune diseases

Only one published study [Stein 2016b] evaluated serum immune biomarkers at baseline (i.e., cross-sectionally) and PFAS serum levels. The study evaluated de-identified archived blood samples from 75 adults aged 21-49. Given the very small sample size, this should be considered a pilot study. The PFHxS serum levels in this study were considerably lower than in the Pease adult population and a few positive findings were observed but the confidence intervals for these findings were extremely wide indicating little precision and a high degree of uncertainty in the effect estimates. Given the strong animal evidence of effects on the immune system from PFAS exposures [NTP 2016], a cross-sectional evaluation of PFAS serum levels and immune biomarkers in a Pease adult study could provide important information on the effects of PFAS exposures on immune function in humans.

The prevalences of ulcerative colitis, rheumatoid arthritis, lupus, and multiple sclerosis in a C8 study [Steenland 2013] were $\leq 1.2\%$. As indicated in Table 7a, $ORs \leq 2.0$ cannot be detected with sufficient statistical power for these endpoints with a sample size of 1,500 per stratum. For lupus and multiple sclerosis, ORs < 3.5 cannot be detected with sufficient statistical power with a sample size of 1,500 per stratum.

Osteoarthritis and Osteoporosis

Two studies evaluated osteoarthritis. In a C8 study [Innes 2011], an OR of about 1.4 was observed for serum PFOA levels considerably higher than those at Pease. However, in an NHANES study [Uhl 2013], an OR of 1.5 was observed for serum PFOA levels similar to those at Pease. Table 7a indicates that ORs in the range of 1.4 - 1.6 can be detected with sufficient statistical power with a sample size of 1.500 per stratum.

An NHANES study evaluated osteoporosis in women [Khalil 2016] and obtained an OR > 10 for serum PFHxS levels lower than those at Pease. With 750 women per stratum, an OR of 1.58 can be detected with sufficient statistical power.

Endometriosis

An NHANES study [Campbell 2016] obtained ORs of 1.47 and 2.86 for serum PFHxS and PFOA, respectively. The serum levels for these two PFAS were similar to those in the Pease population. Table 7a indicates that with a sample size of 750 per stratum, ORs in the range of 1.55 – 1.85 can be detected with sufficient statistical power.

Pregnancy-induced hypertension

Several C8 studies evaluated pregnancy-induced hypertension. One study observed an OR of 1.6 for serum PFOS. However, the PFOS serum levels in the C8 study were higher than those at Pease. Table 7a indicates that ORs in the range of 1.6 - 1.9 can be detected with sufficient statistical power for a sample size of 750 pregnancies per stratum.

Cancer incidence

For kidney cancer, Table 7a indicates that ORs <3.8 cannot be detected with sufficient statistical power with a sample size of 1,500 per stratum. Even for a cancer with a much higher prevalence than kidney cancer, e.g., prostate cancer, ORs < 2.0 cannot be detected with sufficient statistical power with a sample size of 750 men per stratum.

F. Conclusion

A sample size of about 1,500 per stratum (or a total sample size of 6,000 if quartiles are evaluated) would have sufficient statistical power to detect several of the health-related endpoints, as indicated by Tables 7a and 7b. For some endpoints, such as mean difference in uric acid, hyperuricemia, and cardiovascular disease, smaller sample sizes of about 500 per stratum might be sufficient. For other endpoints, such as ulcerative colitis, rheumatoid arthritis, and chronic kidney and liver disease, sample sizes larger than 1,500 per stratum would be necessary. Based on the sample size calculations that assume a sample size of 1,500 adults employed at the Pease Tradeport and 1,500 adults from the Portsmouth area who were never employed at the Pease Tradeport, and assuming a simple comparison of exposed versus unexposed, health endpoints are grouped below into three categories: 1) feasible to study, 2) possible to study (but might require a larger sample size from the Pease population), and 3) not feasible to study using the Pease adult population unless additional populations exposed to PFAS-contaminated drinking water are included in the study.

Health endpoints feasible to study in adults at Pease

- Mean difference in lipids (total cholesterol, LDL, HDL, triglycerides)
- Elevated total cholesterol (hypercholesterolemia)
- Mean difference in uric acid, a measure of kidney function
- Elevated uric acid (hyperuricemia)
- Thyroid disease (unconfirmed)
- Cardiovascular disease
- Hypertension
- Osteoarthritis and osteoporosis
- Mean differences in serum immunoglobin (IgA, IgE, IgG, IgM), and C-reactive protein (an indicator of inflammation); increase in antinuclear antibodies (an indicator of autoimmune reaction); alterations in specific cytokines

<u>Health endpoints that may be possible to study in adults at Pease</u> (although a larger sample size may be needed)

- Liver function
- Thyroid disease (confirmed)
- Thyroid function
- Endometriosis
- Pregnancy-induced hypertension

<u>Health endpoints not feasible to study using the Pease adult population</u> (in order to address these health endpoints, populations from other sites with PFAS-contaminated drinking water would need to be included along with the Pease adult population)

- Liver disease
- Kidney disease
- Ulcerative colitis
- Rheumatoid arthritis
- Lupus
- Multiple sclerosis
- Kidney cancer (and other adult cancers)

To evaluate exposure—response trends, the study participants would need to be split into tertiles or quartiles based on their serum PFAS levels. For some of the candidate health endpoints that are listed above as feasible to study or possible to study, the Pease adult population that can be recruited to participate will not be large enough to be split into exposure tertiles or quartiles and still have sufficient statistical power for comparisons between each of the exposure strata and a reference (unexposed) stratum. For example, if the study population is to be divided into quartiles, and assuming that a sample size of 1,500 per stratum would be sufficient for many of the endpoints of interest, then it would be necessary to recruit 4,500 adults (aged ≥18 years at the start of the study) from the Pease workforce and a representative group (i.e., employed in similar occupations as the Pease workforce) of 1,500 adults from the Portsmouth area who were not exposed at Pease.

Data analyses similar to those used in the C8 studies would be used. The methods include linear regression of continuous (untransformed and natural log-transformed) effect biomarkers on continuous (untransformed and natural log-transformed) PFAS serum levels and categorized PFAS serum levels; and logistic regression of categorized effect biomarkers (e.g., hypercholesterolemia) or disease prevalence on continuous (untransformed and natural log-transformed) and categorical PFAS serum levels. Restricted cubic splines for linear and logistic regression would be conducted to obtain flexible, smoothed exposure-response curves. Measured PFAS serum levels would be evaluated. In addition, for PFOS and PFOA (and possibly PFHxS if an historical reconstruction modeling method becomes available), estimated cumulative serum levels would be evaluated.

In summary, a study limited to the Pease adult population could likely have a sufficient sample size for some of the candidate endpoints **if** the comparisons are simply between an exposed and unexposed group. Recruitment of at least 1,500 adults from Pease should be feasible, given that the 2015 blood testing program at Pease was able to recruit at least 1,182 adults aged >18 years who worked at Pease.

However, a study limited to the Pease adult population might not have a sufficient sample size to evaluate exposure—response relationships. Moreover, a study limited to the Pease worker population might not have sufficient variability in serum PFAS levels to evaluate exposure—response trends effectively. Sufficient variability in PFAS serum levels might be achieved by including other populations with residential exposures to PFAS-contaminated drinking water.

Feasibility of an epidemiological study of former military service and civilian workers at the former Pease Air Force base

Drinking water contamination at a military base involves potential residential exposures to those living and training at the base and potential exposures to those working at the base. At the former Pease Air Force Base, starting in the 1970s, AFFF foam was used for fire training and to extinguish flammable liquid fires. The PFAS contamination in the Haven well water supply likely occurred sometime during the period from the start of AFFF usage and the closing of the base and would have resulted in exposures to those living and working at the base.

To evaluate the incidence and mortality of specific cancers, a large population of adults would need to be followed for a sufficient number of years to account for the long induction periods of most cancers and to have sufficient statistical power. For example, the Camp Lejeune mortality study of U.S. Marines and Navy personnel followed a cohort of 154,932 from 1979 to 2008 for a total of over 4 million person-years [Bove 2014]. To evaluate cancer incidence for the Camp Lejeune cohort, ATSDR will conduct follow-up using state and federal cancer registries for the period 1996–2016 (1996 is the earliest date that >90% of the state registries were in operation), for a total of over 3 million person-years. For the civilian worker cohort at Camp Lejeune, 8,085 workers will be followed over the period 1996–2016 for cancer incidence, for a total of 121,875 person-years. This is similar in size to a study of cancer incidence among workers at a PFAS manufacturing plant [Raleigh 2014]. A recent study of firefighters followed a pooled cohort of 29,993 from San Francisco, Chicago, or Philadelphia from 1985 through 2009, for a total of 403,152 person-years [Daniels 2014]. A C8 study of cancer incidence that relied on self-reported cancers that were confirmed by medical records and cancer registry review included 32,254 who contributed over 1 million person-years of follow-up [Barry 2013].

In October 1989, 3,465 military personnel were assigned to Pease Air Force Base, accompanied by 4,746 dependents. The Air Force estimates that 537 civilian employees were employed on base at that time [USAF 1990]. From 1970 to 1990, an average of 3,000 personnel and their families were assigned to the base at any one time. Before 1970, the base supported a maximum of 5,000 personnel [USAF 1994]. One important consideration about including Pease service personnel and civilian workers in a cancer incidence and mortality study is that drinking water at the base was also contaminated by TCE from the Haven well during some of the years the base operated. Service personnel and civilian workers stationed at the base before 1986 should not be included because of this contamination. Because the base closed by 1991, the number of service personnel and civilian workers at Pease AFB that could be included in a study would be insufficient to evaluate cancers with sufficient statistical power.

Because of the relatively small numbers of personnel assigned to Pease Air Force Base, we conclude that it is not feasible to conduct a study of cancer incidence and mortality that is limited to the Pease military service personnel and civilian worker cohorts stationed at the base from 1986

onward. For a study to be feasible, it would require a larger population size, for example, by including service personnel and civilian workers from other military bases with PFAS-contaminated drinking water as a result of the use of AFFF. Exposures to other drinking water contaminants, such as TCE, other chlorinated organic chemicals, and benzene, must also be taken into account when considering candidate military bases and defining the cohorts.

Cohorts of service personnel and civilian workers can be identified at military bases from personnel data maintained at the Defense Manpower Data Center. Personnel data are available from 1971, although information on military unit, which is needed to determine the base where the individual was stationed, does not begin until the second quarter of 1975. For civilian workers, data are available starting in the last quarter of 1972, with data missing for the first quarter of 1973. The data contain the location of the workplace (codes for state, city, and ZIP code). The Defense Manpower Data Center data contain Social Security number, name, date of birth, and sex to facilitate follow-up.

Military service personnel constitute a highly mobile population after their tours of duty are completed. For a mortality study, this is not a problem, because the NDI is available to obtain information on causes of death. However, there is no national cancer registry to ascertain cancer incidence. Therefore, a study of military service personnel and civilian workers would require gaining the participation of all or most of the state cancer registries and the Department of Veterans Affairs Central Cancer Registry (VACCR). The Camp Lejeune Cancer Incidence Study is one model for such a study. This study is attempting to recruit at least two-thirds of the state cancer registries and VACCR to cover >90% of the Camp Lejeune and Camp Pendleton cohorts. The study will send the personal identifiers for each cohort member to each registry for matching with the registry's data. For any matches that occur, the registry will send to ATSDR the cancer information that is linked to personal identifier (e.g., Social Security number or a unique identification number linked to the Social Security number). This will allow assessment of exposures and other covariates and cancers at the individual level.

The most appropriate military sites for inclusion would be those with water systems that are not complex so that simple mixing models can be used to estimate PFAS-contaminant levels throughout the distribution system. In addition, candidate sites should have information on the history of AFFF use at the base including major incidents such as spills, fires, etc.

Other study designs and health-related endpoints

1. Adverse birth outcomes

To evaluate adverse birth outcomes such as SGA, preterm birth, and specific congenital malformations with sufficient statistical power, several thousand births should be studied. For example, to detect an OR of 1.5 for SGA (5th percentile) with 80% power would require 1,775 births per stratum. For SGA (10th percentile) and preterm birth, with 80% power, an OR of 1.5 can be detected with a sample size of about 960–990 births per stratum. For rare birth defects, such as neural tube defects, to detect an OR of 2.5 with 80% power would require a sample size of about 22,000 births per stratum. For oral clefts, to detect an OR of 2.0 would require about 15,000 births per stratum.

Birth weight, SGA and preterm birth can be evaluated using birth certificate data. For birth defects, a population-based registry must be used to identify cases.

An adverse birth outcome study is not feasible at Pease because there were too few births to mothers who worked at the Tradeport during their pregnancy. The most appropriate candidate populations for a study of adverse birth outcomes would be one or more large municipalities with residential exposures to PFAS-contaminated drinking water where a simple mixing model could be used to estimate contaminant levels throughout the distribution system, i.e., a system that is not complex but instead has relatively uniform contaminant levels throughout the distribution system.

2. Registry

Creating a registry of exposed children and adults at the Pease Tradeport involves following the health status over a period of time and is similar to an epidemiological, longitudinal study of an exposed cohort. The difference is that an epidemiological study would usually include a comparison, unexposed cohort. A registry, like a longitudinal epidemiological study, can be resource-intensive. A decision would also have to be made concerning the length of the follow-up. As in any longitudinal effort, individuals will drop out over time, resulting in interpretation difficulties (e.g., selection bias resulting from loss to follow-up). In any event, before a registry or longitudinal study can be contemplated, an initial cross-sectional study must first be conducted, similar to the children's study and adult study discussed above.

3. Multi-site studies

The results of sample size calculations indicated that the exposed populations at the Pease Tradeport and the former Pease Air Force Base were of insufficient size for some of the health-related endpoints of interest to the community. Moreover, Pease CAP members have expressed interest in linking the Pease communities with other communities that have been exposed to PFAS-contaminated drinking water. A national database exists that can be used to identify other communities with PFAS-contaminated drinking water. Data on PFAS contamination of public drinking water supplies are available for large systems (serving >10,000 retail customers) and a small sample of small systems (n = 800 or 0.5% of a total of 144,165 systems serving <10,000 retail customers) via the Third Unregulated Contaminant Monitoring Rule (UCMR-3) database maintained by the EPA [US EPA 2016b].

UCMR-3 monitoring for PFAS is required at the entry point to the distribution system for each well and at any interconnection that is in operation. Water utilities had to sample twice during a 12-month period from 2013–2015 with sampling events occurring 5–7 months apart. The UCMR dataset contains sampling data from January 2, 2013 through March 1, 2016. Table A1 in the Appendix lists the utilities ranked by the maximum level of combined PFOS and PFHxS detected in the system. The highest level was detected in the system serving the Mariana Islands. Among the U.S. water systems, the top 10 systems for combined PFOS and PFHxS were Artesian Water Company in Delaware; Security Water System in Colorado Springs, CO; Horsham and Warminster systems in Pennsylvania; Oatman Water Company in Arizona; Issaquah Water System in Washington; Hyannis Water System in Massachusetts; Suffolk County Water Authority in New York; Warrington Township Water in Pennsylvania; and United Water in Pennsylvania, which serves various municipalities.

Although the UCMR database can be used to identify potential sites for further consideration for health studies, it has several limitations. First, most small systems are not included in the database. Second, the data represent levels of contamination at the entry points to the distribution system of the water utility

(e.g., contaminant levels in a supply well) and generally do not represent the levels of contamination reaching particular residences served by the utility. To estimate the population receiving contaminated drinking water and the levels of PFAS in their drinking water, the UCMR data must be supplemented with information on the configuration and operation of the utility's system. For a system that mixes all its sources of water before to entering the distribution system, a simple mixing model can be used to estimate the contaminant levels in the drinking water serving the residences by taking into account the contaminant levels in each source and the contribution of each source to the total supply. This is the situation at the Pease Tradeport, where water from each of the supply wells is mixed at the treatment plant before entering the distribution system. However, many utilities have more complex systems in which each of the supply wells (or surface water sources) primarily serve particular areas of the distribution system. For these systems, additional information is needed (for example, on the operation of the supply wells, tank levels, and the water demand in each area of the distribution system), and complex modeling methods must be used.

Conclusions

The ability of a study of the Pease population to provide useful information will depend to a great extent on the success of recruiting sufficient number of study participants. The feasibility assessment concluded that it is possible to evaluate some health-related endpoints if a sufficient number of children and adults from the Pease population participate. Other health-related endpoints would require larger numbers of exposed individuals and would require the inclusion of populations from other sites who were exposed to PFAS-contaminated drinking water. The feasibility assessment concluded that a third study design, a mortality and cancer incidence study of former military service and civilian worker personnel, would not be feasible solely with the population at Pease.

The feasibility assessment is still a draft. It will be finalized once the Pease Community Assistance Panel (CAP) and the larger Pease Tradeport community have the opportunity to review and make comments on the assessment. ATSDR will then revise the assessment based on the comments received. The feasibility of successfully evaluating particular health-related endpoints (or effect biomarkers) could change depending on final study design and goals.

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Tables

Table 1. Serum levels of selected PFAS in $\mu g/L$, children aged <12 years in Pease and comparison

populations.

	Pease Tradeport*			$\mathbf{T}\mathbf{X}^{\dagger}$		$\mathbf{C}\mathbf{A}^{\ddagger}$			
PFAS	Median	Geometric mean	95% CI	Max.	Median	Max.	Median	Geometric mean	Max.
PFOA	3.63	3.43	3.23 - 3.64	12.00	2.85	13.50	4.50	4.46	19.50
PFOS	8.27	8.11	7.59 - 8.67	30.80	4.10	93.90	6.15	6.28	26.70
PFHxS	4.24	3.83	3.48 - 4.22	31.70	1.20	31.20	1.25	1.30	9.80
PFNA	0.90	0.92	0.86 - 0.98	5.20	1.20	55.80	1.70	1.84	11.20

^{*} Pease: N=366, aged <12 years; sampling occurred in **2015**.

Table 2. Serum levels of selected PFAS in $\mu g/L$, aged ≥ 12 years in Pease and NHANES comparison values.

	Pease Tradeport*				NHANES 2013-2014 [†]			
PFAS	Median	Geometric Mean	95% CI	Max.	Median	Geometric Mean	95% CI	95 th percentile
PFOA	3.10	2.99	2.87 - 3.11	32.00	2.07	1.94	1.76 - 2.14	5.57
PFOS	9.17	8.74	8.37 - 9.13	95.60	5.20	4.99	4.50 - 5.52	18.50
PFHxS	4.16	4.21	3.98 - 4.46	116.00	1.40	1.35	1.20 - 1.60	5.60
PFNA	0.70	0.68	0.65 - 0.70	4.90	0.70	0.68	0.61 - 0.72	2.00

^{*}N=1,212 ages \geq 12 years, sampled in 2015

[†] TX reference group: N=300, age ≤12 years (Schecter et al 2012), sampling occurred in **2009**.

[‡]CA reference group: N=68, ages 2-8 years (Wu et al 2015), sampling occurred in **2007-2009**.

 $^{^{\}dagger}$ N=2,168 ages ≥12 years.

Table 3. Summary of the PFAS literature on adults.

Table 5. Summary of the FFAS liter	PFOS	PFHxS	PFOA
Cancer			
Prostate	+	+	+
Bladder	+	*	+
Colorectal	+	*	I
Breast	I	I	+
Pancreatic	I	*	+
Testicular	*	*	+
Kidney	*	*	+
Thyroid	*	*	+
Liver	*	*	+
Leukemia	*	*	+
non-Hodgkin lymphoma	*	*	+
Multiple myeloma	*	*	*
Ovarian	*	*	+
Other diseases			
Kidney disease/kidney function	*	*	+
Hyperuricemia	+	I	+
Liver disease/liver function	+	+	+
Cardiovascular Disease,	+	+	+
hypertension, hypercholesterolemia			
Thyroid disease/function	+	+	+
Autoimmune diseases	*	*	+
Osteoarthritis, osteoporosis and	+	+	+
bone mineral density			
Immune response	+	+	+
Reproductive outcomes	+	+	+

[&]quot;+" One or more studies suggesting increased risk of an adverse outcome (e.g., OR or $RR \ge 1.20$)

[&]quot;*" no studies were conducted (for liver cancer and PFOS, and multiple myeloma and PFOA, there were too few deaths (≤2) to evaluate).

[&]quot;I" inconclusive – the findings have not suggested an increased risk (e.g., an OR or RR <1.20)

Table 4. Summary of the PFAS literature on children.

	PFOS	PFHxS	PFOA
Adverse birth outcomes	+	+	+
Lipids	+	I	+
Thyroid function	+	*	+
Thyroid disease	I	*	+
Uric acid	+	+	+
Sex hormones	+	+	+
Delay in reaching puberty	+	I	+
Neurobehavioral outcomes	+	+	+
Immune function	+	+	+
Hypertension	I	*	I
Adiposity/BMI/Overweight	+	+	+

[&]quot;+" One or more studies suggesting increased risk of an adverse outcome (e.g., OR or $RR \ge 1.20$)

Note: adverse birth outcomes are not included in this table because these outcomes are not feasible to study at Pease. Although the number of children potentially exposed to the PFAS-contaminated drinking water while attending daycare at the Pease Tradeport can be estimated, there is a lack of information on the number of children potentially exposed in utero to the PFAS-contaminated drinking water because their mothers were employed at the Pease Tradeport during the pregnancy. To evaluate adverse birth outcomes with sufficient statistical power would require the inclusion of several hundreds of exposed births.

[&]quot;*" no studies were conducted.

[&]quot;I" inconclusive – the findings have not suggested an increased risk (e.g., an OR or RR <1.20)

Table 5. Pease serum levels in $\mu g/L$ for PFOS, PFOA and PFHxS, based on ages in 2018.

Population age (2018)		PFOS	PFOA	PFHxS
4-16 (n = 370)	Mean	9.74	3.89	5.54
during 2018	SD	6.05	2.14	4.65
	Median	8.16	3.47	4.20
	Geometric mean	8.05	3.34	3.80
	Maximum	30.80	12.00	31.70
	Top quartile	12.90	5.01	7.80
	Top quintile	14.20	5.50	8.80
\geq 18 (n = 1,190)	Mean	11.59	3.76	7.05
during 2018	SD	9.63	2.74	8.75
	Median	9.30	3.12	4.21
	Geometric mean	8.82	3.02	4.26
	Maximum	95.60	32.00	116.00
	Top quartile	14.20	4.65	8.60
	Top quintile	16.10	5.20	10.00
≥18, not a firefighter	Mean	10.95	3.75	6.52
(n = 1,092)	SD	8.32	2.78	7.44
	Median	8.97	3.10	4.08
	Geometric mean	8.53	2.99	4.04
	Maximum	78.00	32.00	61.40
	Top quartile	13.70	4.62	8.10
	Top quintile	15.50	5.10	9.45
Firefighter (n = 98)	Mean	18.70	3.93	12.95
	SD	17.41	2.14	16.64
	Median	11.75	3.40	8.14
	Geometric mean	12.80	3.37	7.74
	Maximum	95.60	12.10	116.00
	Top quartile	23.10	5.26	14.70
	Top quintile	28.64	5.90	17.46

Table 6a. Minimum detectable effects for a Pease children study with 350 exposed and 175 unexposed. *

Endpoint	α and $\beta = .05$	$\alpha = .05, \beta = .20$	α and $\beta = .10$	$\alpha = .10, \beta = .20$
Total cholesterol	9.8 mg/dL	7.6 mg/dL	8.0 mg/dL	6.8 mg/dL
(mean difference)				
Hypercholesterolemia	OR = 2.00	OR = 1.73	OR = 1.77	OR =1 .63
Hyperuricemia	OR = 2.30	OR = 1.96	OR = 2.00	OR = 1.83
Uric acid (mean	0.40 mg/dL	0.31 mg/dL	0.33 mg/dL	0.28 mg/dL
difference)				
eGFR (mean	8.0	6.2	6.5	5.5
difference)#				
ADHD [¶]	OR = 2.47	OR = 2.09	OR = 2.13	OR = 1.94
ADHD + meds¶	OR = 3.50	OR = 2.80	OR = 2.89	OR = 2.52
Atopic dermatitis	OR = 2.49	OR = 2.10	OR = 2.15	OR = 1.95
Asthma	OR = 2.56	OR = 2.16	OR = 2.21	OR = 2.00
Rhinitis	OR = 2.08	OR = 1.79	OR = 1.83	OR = 1.69
Hypertension	OR = 2.12	OR = 1.80	OR = 1.85	OR = 1.69
Overweight/Obese	OR = 2.00	OR = 1.72	OR = 1.76	OR = 1.62

Table 6b. Minimum detectable effects for a Pease children study with 500 exposed and 250 unexposed. *

Endpoint	α and $\beta = .05$	$\alpha = .05, \beta = .20$	α and $\beta = .10$	$\alpha = .10, \beta = .20$
Total cholesterol	8.2 mg/dL	6.4 mg/dL	6.7 mg/dL	5.7 mg/dL
(mean difference)				
Hypercholesterolemia	OR = 1.78	OR = 1.57	OR = 1.60	OR =1 .50
Hyperuricemia	OR = 2.04	OR = 1.75	OR = 1.79	OR = 1.65
Uric acid (mean	0.34 mg/dL	0.26 mg/dL	0.27 mg/dL	0.23 mg/dL
difference)				
eGFR (mean	6.7	5.2	5.4	4.6
difference)#				
ADHD [¶]	OR = 2.18	OR = 1.85	OR = 1.90	OR = 1.73
ADHD + meds¶	OR = 2.98	OR = 2.40	OR = 2.48	OR = 2.19
Atopic dermatitis	OR = 2.20	OR = 1.86	OR = 1.91	OR = 1.74
Asthma	OR = 2.26	OR = 1.91	OR = 1.96	OR = 1.78
Rhinitis	OR = 1.85	OR = 1.62	OR = 1.65	OR = 1.54
Hypertension	OR = 1.88	OR = 1.64	OR = 1.68	OR = 1.56
Overweight/Obese	OR = 1.79	OR = 1.58	OR = 1.61	OR = 1.50

^{*} Some health-related endpoints are not included in the table because there was insufficient information to calculate minimum detectable effects. For sex hormones, insulin-like growth factor – 1, and thyroid function, see the appendix for a description of the assumptions used in the sample size calculations and the resulting calculations.

[#] mL/min/1.73 m²

The prevalence of an ADHD diagnosis reported by a study participant in the C8 study (Stein 2011) was 12.4%. In this study, the prevalence of an ADHD diagnosis reported by a study participant who also reported currently using a medication commonly used to treat ADHD was 5.1%.

Table 6c. Summary of information used to categorize the feasibility of studying health-related endpoints for a Pease children study.

Health-related Endpoint	Minimum Detectable Effect Size: 350 exposed, 175 unexposed	Other Sample Size Considerations	Conclusion
Lipids (total cholesterol)	6.8 mg/dL	A Taiwan study (Zeng 2015) obtained mean differences of 11-12 mg/dL for total cholesterol and low density lipoprotein at PFOA serum levels similar to Pease.	Feasible to study at Pease
Estimated glomerular filtration rate (eGFR)	5.5 mL/min/1.73 m ²	A NHANES study (Kataria 2015) observed a mean difference of 6.6 mL/min/1.73 m ² for PFOA serum levels similar to those at Pease. For PFOS, the mean difference was 7.2 mL/min/1.73 m ²	Feasible to study at Pease
Insulin-like growth hormone-1 (IGF-1)	See appendix for sample size calculations and assumptions required for the calculations.	A C8 study (Lopez-Espinosa 2016) observed a reduction of IGF-1 for PFHxS serum levels similar to those at Pease that could be detected with sufficient power by a sample size of 350 exposed and 175 unexposed.	Feasible to study at Pease.
Overweight/Obesity	OR=1.62	A Faroes study (Karlsen 2016) observed and OR of 1.88 for PFOA serum levels below those at Pease. This OR could be detected with a sample size of 350 exposed and 175 unexposed children.	Feasible to study at Pease.
Hypercholesterolemia	OR=1.63	A NHANES study (Geiger 2014) obtained ORs of 1.49 and 1.35 for serum PFOA and PFOS levels similar to those at Pease. To detect an OR of 1.49 with 80% power requires a minimum of 540 exposed and 270 unexposed	Possible to study at Pease although a sample size of at least 500 exposed and 250 unexposed would be necessary (see table 6b).
Uric acid	0.28 mg/dL	A NHANES study (Kataria 2015) obtained a mean difference of 0.21 mg/dL for PFOA serum levels similar to Pease. However, for PFOS, the mean difference was 0.05 mg/dL.	Possible to study at Pease although a larger sample size than 500 exposed and 250 unexposed would be necessary.

Health-related Endpoint	Minimum Detectable Effect Size	Other Sample Size Considerations	Conclusion
Hyperuricemia	OR=1.83	A Taiwan study (Qin 2016) obtained an OR of 1.65 for PFHxS serum levels much lower than at Pease. For PFOA serum levels lower than at Pease, an OR of 2.2 was obtained.	Possible to study at Pease although a sample size of at least 500 exposed and 250 exposed may be necessary to evaluate the effect of serum PFHxS. (For serum PFOA, the Pease sample size of 350 exposed and 175 unexposed may be sufficient)
IQ	3 point mean difference	A Taiwan study (Wang 2015) obtained IQ mean differences of ≤2 points for PFOS serum levels higher than at Pease. A C8 study (Stein 2013) did not find a decrease in IQ with PFOA exposure and did not evaluate PFOS or PFHxS.	Possible to study at Pease although a sample size larger than 500 exposed and 250 unexposed would be necessary.
Neurobehavioral	Could not be calculated due to insufficient information	Some studies had sample sizes achievable at Pease while others had much larger sample sizes. The effects observed were not large (e.g., an OR for learning problems was 1.2 for PFHxS and lower for the other PFAS, and ORs for hyperactivity and coordination problems were <1.5 for each of the PFAS). The few studies that have been conducted evaluated different neurobehavioral tests.	Similar conclusion as for IQ: Possible to study at Pease although a sample size larger than 500 exposed and 250 unexposed would be necessary.
Sex hormones	See appendix for sample size calculations and assumptions required for the calculations.	At PFOS serum levels much higher than at Pease, a C8 study (Lopez-Espinosa 2016) observed reductions in estradiol that would require a sample size of over a thousand of exposed to achieve sufficient statistical power. However, the observed reductions in testosterone would require a sample size of between 500 and 1,000 exposed.	Possible to study at Pease although a sample size larger than 500 exposed and 250 unexposed would be necessary.

Health-related Endpoint	Minimum Detectable Effect Size	Other Sample Size Considerations	Conclusion
Thyroid function	See appendix for sample size calculations and assumptions required for the calculations.	A C8 study (Lopez-Espinosa 2012) observed small differences for PFOS and PFOA serum levels considerably higher than at Pease. To detect these differences would require a sample size of over a thousand exposed. On the other hand, a Taiwan study (Lin 2013) observed differences that could be detected with sufficient power with a sample size of 350 exposed and 175 unexposed.	Possible to study at Pease.
Atopic dermatitis	OR=1.95	A Taiwan study (Wang 2011) obtained an OR of 2.19 for PFOS serum levels similar to Pease. However, the study evaluated children aged 2 years. No other PFAS study evaluated atopic dermatitis	Possible to study at Pease.
Asthma	OR=2.00	Two NHANES studies (Humblet 2014, Stein 2016) observed ORs between 1.2 and 1.3 which would require a sample size of over 2,000 exposed. However, a Taiwan study (Dong 2013) obtained ORs between 3.8 and 4.0 for PFHxS and PFOA serum levels lower than at Pease.	Possible to study at Pease.
Rhinitis	OR=1.69	A NHANES study (Stein 2016a) evaluated rhinitis and obtained an OR of 1.35 for serum PFOA similar to those at Pease. To detect this OR would require over a thousand exposed. However, ORs between 1.5 and 1.6 could be detected with sufficient statistical power with a sample size of 500 exposed and 250 unexposed. These are ORs that are reasonable to detect and fall within the 95% CI for the finding in the NHANES study.	Possible to study at Pease
Antibody response to childhood vaccines	Could not be calculated due to insufficient information	Three studies that have been conducted of these endpoints had sample sizes that could be achievable at Pease. Only two studies (Granum 2013, Stein 2016) have evaluated the same endpoint – rubella.	Possible to study at Pease although a sample size larger than 500 exposed and 250 exposed may be necessary.

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Health-related Endpoint	Minimum Detectable Effect Size	Other Sample Size Considerations	Conclusion
Attention deficit/hyperactivity disorder (ADHD)	ORs: 1.9 – 2.5	A C8 study (Stein 2011) obtained an OR of 1.55 (ADHD + meds) for PFHxS serum levels similar to Pease. A NHANES study (Hoffman 2010) observed an OR of 1.67 for PFHxS serum levels similar to Pease.	Not feasible to study using the Pease population alone (for ADHD confirmed by current medications)
Autism spectrum disorder (ASD)	ORs > 4.0	One study (Liew 2015) obtained an OR of 1.3 for serum PFHxS levels lower than at Pease. To detect this OR would require >10,000 exposed.	Not feasible to study using the Pease population alone.
Delayed puberty	Could not be calculated due to insufficient information	Only one study evaluated delayed puberty among children. This was a C8 study (Lopez-Espinosa 2011) that evaluated several thousand children. It is likely that sample sizes much larger than at Pease would be necessary.	Not feasible to study using the Pease population alone.
Thyroid disease	OR > 8.0	A C8 study (Lopez-Espinosa 2012) obtained an OR of 1.44 for PFOA serum levels considerably higher than those in the Pease population. To detect this OR with 80% statistical power would require a sample size of over 10,000 exposed children.	Not feasible to study using the Pease population alone.
Childhood cancers		No PFAS study has evaluated childhood cancers. Given the incidence and prevalence of cancers such as leukemia, a sample size of many thousands of exposed would be necessary.	Not feasible to study using the Pease population alone.

The minimum detectable effect size is based on a sample size of 350 children exposed and 175 children unexposed, and specifying statistical power of 80% (or a type 2 or " β " error of .20) and a type 1 (" α ") error of .10 (see table 6a). This minimum detectable effect size is compared to the adverse effect sizes observed in other PFAS studies. Where possible, the focus is on adverse effect sizes in the PFAS studies observed for PFAS serum levels similar to those among the Pease children. An endpoint is considered feasible to study at Pease if an adverse effect size observed in PFAS study can be detected with sufficient statistical power (i.e., statistical power of \geq 80%) by a sample size achievable at Pease, i.e., a sample size of 350 exposed children at Pease and 175 children unexposed to the PFAS-contaminated drinking water at Pease. If only one PFAS study has been conducted on a health-related endpoint, then the endpoint was considered feasible to study at Pease if an odds ratio of <2.0 could be detected with statistical power of 80%.

Note: The studies mentioned in the column of the table labeled "Other Sample Size Considerations" are included only to give a sense of the adverse effect sizes that might occur in a Pease study. Due to the paucity of studies for each health-related endpoint, there is considerable uncertainty concerning the effect sizes that might be expected to occur in a Pease study.

OR: Odds ratio. The odds ratio roughly approximates the risk ratio. The risk ratio is the proportion of the exposed population with a disease divided by the proportion of the unexposed population with a disease.

Note: Hypertension is not included in this table because there is no evidence so far of an association between PFAS serum levels and hypertension in children. Adverse birth outcomes are not included in this table because these outcomes are not feasible to study at Pease. Although the number of children potentially exposed to the PFAS-contaminated drinking water while attending daycare at the Pease Tradeport can be estimated, there is a lack of information on the number of children potentially exposed in utero to the PFAS-contaminated drinking water because their mothers were employed at the Pease Tradeport during the pregnancy. To evaluate adverse birth outcomes with sufficient statistical power would require the inclusion of several hundreds of exposed births.

Note: The health-related endpoints listed in this table satisfy the criteria of scientific importance and public health significance as discussed on page 8 of the text.

Table 7a. Minimum detectable effects for an adult epidemiological study, 1,500 per stratum.*

Endpoint	α and $\beta = .05$	$\alpha = .05, \beta = .20$	α and $\beta = .10$	$\alpha = .10, \beta = .20$
Chronic kidney disease	OR=2.54	OR=2.14	OR=2.20	OR=2.00
Thyroid disease, unconfirmed	OR=1.48	OR=1.36	OR=1.38	OR=1.32
Thyroid disease, confirmed	OR=1.63	OR=1.48	OR=1.50	OR=1.42
Total cholesterol (mean	5.5 mg/dL	4.3 mg/dL	4.5 mg/dL	3.8 mg/dL
difference)				
LDL (mean difference)	4.5 mg/dL	3.5 mg/dL	3.7 mg/dL	3.1 mg/dL
Hypercholesterolemia	OR=1.42	OR=1.32	OR=1.34	OR=1.28
Uric acid (mean difference)	0.21 mg/dL	0.17 mg/dL	0.18 mg/dL	0.15 mg/dL
Hyperuricemia	OR=1.35	OR=1.27	OR=1.28	OR=1.24
Elevated ALT (>45 IU/L, men;	OR=1.49	OR=1.37	OR=1.39	OR=1.33
>34 IU/L, women)				
Elevated GGT (>55 IU/L, men;	OR=1.44	OR=1.33	OR=1.35	OR=1.29
>38 IU/L, women)				
Elevated direct bilirubin	OR=2.80	OR=2.34	OR=2.40	OR=2.16
(>0.03 mg/dL)				
ALT (mean difference)	2.65 IU/L	2.06 IU/L	2.15 IU/L	1.83 IU/L
GGT (mean difference)	5.92 IU/L	4.60 IU/L	4.80 IU/L	4.09 IU/L
Direct bilirubin (mean	0.079 mg/dL	0.060 mg/dL	0.064 mg/dL	0.055 mg/dL
difference)				
Liver disease	OR=2.24	OR=1.92	OR=1.97	OR=1.80
Cardiovascular disease	OR=1.45	OR=1.34	OR=1.36	OR=1.30
Hypertension	OR=1.31	OR=1.24	OR=1.25	OR=1.21
Ulcerative colitis	OR=4.13	OR=3.24	OR=3.38	OR=2.94
Rheumatoid arthritis	OR=2.70	OR=2.25	OR=2.32	OR=2.10
Lupus	OR=6.87	OR=4.97	OR=5.24	OR=4.33
Multiple Sclerosis	OR=5.30	OR=3.97	OR=4.15	OR=3.50
Osteoporosis	OR=1.73	OR=1.55	OR=1.58	OR=1.48
Osteoarthritis	OR=1.58	OR=1.44	OR=1.46	OR=1.39
Endometriosis	OR=1.92	OR=1.69	OR=1.73	OR=1.61
(750 per stratum)				
Pregnancy-induced hypertension	OR=1.84	OR=1.63	OR=1.66	OR=1.55
(750 per stratum)				
Kidney cancer	OR=5.60	OR=4.27	OR=4.45	OR=3.80

^{*} Some health-related endpoints are not included in the table because there was insufficient information to calculate minimum detectable effects. For thyroid function, see the appendix for a description of the assumptions used in the sample size calculations and the resulting calculations.

Table 7b. Summary of information used to categorize the feasibility of studying health-related endpoints for a Pease adult study.

Health-related Endpoint	Minimum Detectable Effect Size: 1,500 exposed and 1,500 unexposed	Other Sample Size Considerations	Conclusion
Lipids (total cholesterol)	3.8 mg/dL	A C8 study (Steenland 2009) observed a 3 – 4 mg/dL change in total cholesterol and LDL for PFOS serum levels similar to those at Pease.	Feasible to study at Pease
Hypercholesterolemia	OR=1.28	A Canadian study (Fisher 2013) obtained an OR of 1.57 for PFHxS serum levels similar to those at Pease.	Feasible to study at Pease
Uric acid	0.15 mg/dL	A NHANES study (Shankar 2011) observed a mean difference of 0.40 mg/dL for serum PFOA levels similar to those at Pease.	Feasible to study at Pease
Hyperuricemia	OR=1.24	A NHANES study (Shankar 2011) obtained an OR of 1.90 for serum PFOA levels similar to those at Pease.	Feasible to study at Pease
Thyroid disease (unconfirmed)	OR=1.32	A C8 study (Winquist 2014a), hazard ratios ≤1.3 were obtained for PFOA serum levels similar to those at Pease. (Only PFOA was evaluated in this study.)	Feasible to study at Pease
Cardiovascular disease	OR=1.30	A NHANES study (Shankar 2012) obtained an OR of 2.01 for PFOA serum levels similar to those at Pease.	Feasible to study at Pease
Hypertension	OR=1.21	Only one community study (a C8 study, Winquist 2014b), evaluated hypertension and obtained an OR < 1.0 for serum PFOA (the only PFAS evaluated). However, the sample size achievable at Pease is capable of detecting very low ORs with sufficient statistical power.	Feasible to study at Pease
Osteoarthritis	OR=1.39	A NHANES study (Uhl 2013) obtained an OR of 1.5 for serum PFOA levels similar to those at Pease.	Feasible to study at Pease
Osteoporosis	OR=1.48	A NHANES study (Khalil 2016) obtained an OR > 10 among women, for serum PFHxS levels lower than those at Pease.	Feasible to study at Pease

Health-related Endpoint	Minimum Detectable Effect Size	Other Sample Size Considerations	Conclusion
Serum Immune Biomarkers	Could not be calculated due to insufficient information	Only one published study (Stein 2016b) has been conducted that evaluated serum immune biomarkers at baseline (i.e., cross-sectionally). This study had a sample size of 75 adults. A cross-sectional evaluation of PFAS serum levels and immune biomarkers in a Pease adult study could provide important information on the effects of PFAS exposures on immune function in humans.	Feasible to study at Pease
Liver function: Elevated ALT Elevated GGT Elevated direct bilirubin	OR=1.33 OR=1.29 OR=2.16	A NHANES study (Gleason 2015) evaluated PFAS serum levels similar to those at Pease. For elevated ALT, ORs between 1.2 and 1.5 were obtained. For elevated GGT, ORs between 1.0 and 1.3 were obtained. For elevated direct bilirubin, ORs between 1.1 and 1.7 were obtained.	Possible to study at Pease, but may require a larger sample size than 1,500 exposed and 1,500 unexposed to evaluate PFOS and PFHxS serum levels and ALT and GGT. Direct bilirubin is probably not feasible to study using the Pease population alone.
Thyroid disease (confirmed)	OR=1.42	A C8 study (Winquist 2014a), hazard ratios ≤1.3 were obtained for PFOA serum levels similar to those at Pease. (Only PFOA was evaluated in this study.)	Possible to study at Pease, but will require a larger sample size than 1,500 exposed and 1,500 unexposed.
Thyroid function	See appendix for sample size calculations and assumptions required for the calculations.	A C8 study (Knox 2011) observed very subtle changes that would require a study of equivalent size (52,296) to detect associations with sufficient statistical power. On the other hand, a NHANES study (Wen 2013) observed larger changes (at PFAS serum levels similar to those at Pease) that could be detected with a sample size achievable at Pease.	Possible to study at Pease.

Health-related Endpoint	Minimum Detectable Effect Size	Other Sample Size Considerations	Conclusion
Endometriosis	OR=1.61 (750 exposed & 750 unexposed)	A NHANES study (Campbell 2016) obtained ORs of 1.47 and 2.86 for serum PFHxS and PFOA, respectively. The serum levels for these two PFAS were similar to those in the Pease population.	Possible to study at Pease if sufficient numbers of women can be recruited.
Pregnancy-induced hypertension	OR=1.55 (750 exposed pregnancies and 750 unexposed pregnancies	A C8 study (Stein 2009, Darrow 2013) obtained an OR of 1.6 for serum PFOS levels higher than at Pease.	Possible to study at Pease but may require a larger sample size than 1,500 exposed and 1,500 unexposed in order to achieve a sufficient number of pregnancies.
Liver disease	OR=1.80	A C8 study (Darrow 2016) and a NHANES study (Melzer 2010) observed no elevation in liver disease. However, the C8 study evaluated only PFOA and the NHANES study evaluated PFOA and PFOS but not PFHxS.	Not feasible to study using the Pease population alone.
Kidney disease	OR=2.00	A C8 study (Dhingra 2016a) evaluated only PFOA and obtained ORs of 1.26 and 1.36 for the retrospective and prospective analyses, respectively, at the second quintile PFOA serum level. (Smaller ORs were observed at higher PFOA serum levels.)	Not feasible to study using the Pease population alone.
Ulcerative colitis	OR=2.94	A C8 study (Steenland 2013) observed RRs between 2.8 and 3.1 at the highest serum PFOA levels, considerably higher than those at Pease. At lower PFOA serum levels, the RRs were <2.2	Not feasible to study using the Pease population alone.
Rheumatoid arthritis	OR=2.10	A C8 study (Steenland 2013) observed RRs between 1.3 and 1.7 for serum PFOA.	Not feasible to study using the Pease population alone.
Lupus	OR=4.33	A C8 study (Steenland 2013) observed RRs <1.3 for serum PFOA.	Not feasible to study using the Pease population alone.
Multiple sclerosis	OR=3.50	A C8 study (Steenland 2013) observed RRs between 1.1 and 1.6 for serum PFOA	Not feasible to study using the Pease population alone.

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Health-related	Minimum	Other Sample Size Considerations	Conclusion
Endpoint	Detectable Effect		
	Size		
Kidney cancer	OR=3.80 for kidney	A C8 study of a community population (Vieira 2013)	Not feasible to study using the
	cancer	observed an RR of 1.70 for those served by the Little	Pease population alone. (Due to the
		Hocking water system.	very low background prevalences
			of other adult cancers, it is not
			feasible to study cancers using the
			Pease population alone.)

The minimum detectable effect size is based on a sample size of 1,500 adults exposed and 1,500 adults unexposed, and specifying statistical power of 80% (or a type 2 or " β " error of .20) and a type 1 (" α ") error of .10 (see table 6a). This minimum detectable effect size is compared to the adverse effect sizes observed in other PFAS studies. Where possible, the focus is on adverse effect sizes in the PFAS studies observed for PFAS serum levels similar to those among the Pease adults. An endpoint is considered feasible to study at Pease if an adverse effect size observed in PFAS study can be detected with sufficient statistical power (i.e., statistical power of \geq 80%) by a sample size of 1,500 exposed and 1,500 unexposed. If only one PFAS study has been conducted on a health-related endpoint, then the endpoint was considered feasible to study at Pease if an odds ratio of <2.0 could be detected with statistical power of 80%.

Note: the studies mentioned in the column of the table labeled "Other Sample Size Considerations" are included only to give a sense of the adverse effect sizes that might occur in a Pease study. Due to the paucity of studies for each health-related endpoint, there is considerable uncertainty concerning the effect sizes that might be expected to occur in a Pease study.

OR: odds ratio. The odds ratio roughly approximates the risk ratio (RR). The risk ratio is the proportion of the exposed population with a disease divided by the proportion of the unexposed population with a disease.

A hazard ratio can be interpreted in the same way as a risk ratio.

Note: The health-related endpoints listed in this table satisfy the criteria of scientific importance and public health significance as discussed on page 8 of the text.

Appendix

Literature review

The literature review focused on the epidemiological results for PFOA, PFOS and PFHxS since these were the major contaminants detected in the Haven Well during the April and May 2014 sampling as well as the elevated PFAS in the serum of those tested in the NH DHHS Pease testing program. The purpose of the literature review was to identify the health-related endpoints that have been evaluated in at least one epidemiological study, and to assess the extent of the epidemiological research on the health effects of PFHxS and PFOS. The findings of the studies included in the literature review were also used to inform sample size calculations.

Literature searches using PubMed were conducted to identify epidemiological studies that evaluated measured or estimated serum levels of PFOS, PFOA and PFHxS. The key words used in the search were perfluorooctane sulfonate, PFOS, perfluorooctanoic acid, PFOA, perfluorohexane sulfonate, PFHxS, perfluoroalkyl substances, PFAS, perfluorinated compounds, PFC, and perfluorinated chemicals. The PubMed search identified epidemiological studies through October 31, 2016.

Cancers

The C8 science panel in 2012 reviewed the literature on PFAS and cancers and concluded that there was a "probable link" between exposure to PFOA and testicular and kidney cancer (http://www.c8sciencepanel.org/pdfs/Probable_Link_C8_Cancer_16April2012_v2.pdf). No other cancers were considered to have a probable link with PFOA exposure. The panel noted that PFOA caused liver, testicular and pancreatic tumors (adenomas) in rodent studies.

A review of the literature by DuPont researchers noted that PFOS causes liver adenomas in rodent studies (Kennedy and Symons 2015) but concluded that the evidence of associations between community drinking water exposures to PFOA and kidney and testicular cancers was "limited". The review also concluded that studies of populations exposed to low levels of PFOA and PFOS had equivocal results with no consistent associations across studies. Studies of workers exposed to higher levels of PFOS and PFOA were also viewed as lacking consistent associations. Their overall conclusion was: "Based on the evidence reported to date, the prospect for developing a carcinogenic outcome following exposure to PFOA and PFOS is remote."

ATSDR's literature search identified fifteen studies and the results from these studies are summarized in Table A2. Based on its assessment of the epidemiological literature, ATSDR concluded that for most cancers there was either limited information or no information concerning associations with PFAS exposures. In particular, very few studies have evaluated PFHxS exposures and cancers. Although more information is available for PFOS exposure and cancers, the information is still very limited. Clearly more research is needed to investigate whether PFHxS and PFOS exposures are associated with increased risks of specific cancers. More information is available on PFOA exposure and cancers because of the studies conducted of the C8 population (workers and community members) and of workers at the 3M Cottage Grove plant. However, the available information is still too limited to determine whether a causal association exists between PFOA exposure and specific cancers. While cancer research at Pease is not feasible, additional research on the effects of PFOA exposure on specific cancers has the potential to provide the evidence necessary to assess whether PFOA is a cause of one or more specific cancers.

Other adult diseases (Table A3)

The C8 science panel reviewed the literature for adult non-cancer diseases and found probable links for PFOA and ulcerative colitis (an autoimmune disease), hypercholesterolemia (high cholesterol), thyroid disease (hyperthyroidism in females, hypothyroidism in males), and pregnancy-induced hypertension (PIH). The panel concluded that the evidence was not sufficient for a probable link between PFOA and other autoimmune diseases (e.g., lupus, rheumatoid arthritis and Crohn's disease), stroke, hypertension, coronary artery disease, diabetes, chronic kidney or liver disease, asthma, chronic obstructive pulmonary disease, osteoarthritis or Parkinson disease (http://www.c8sciencepanel.org/prob_link.html). Another review of the literature noted that PFOA was linked to uric acid levels and that PFAS exposure was associated with elevated liver enzymes, osteoarthritis, kidney disease and immunotoxicity in some studies, but the findings across studies were inconsistent (Khalil 2015). The following literature review of other adult diseases focuses on studies of populations (e.g., studies utilizing NHANES data) and highly exposed communities (e.g., the C8 population). In these studies, the exposure assessment is based on serum PFAS levels, either measured or predicted based on physiologically-based pharmacokinetic (PBPK) models. We use the same classification scheme for the other adult diseases as was used above for cancers.

1. Kidney function/kidney disease

Two studies of the C8 population were conducted to evaluate kidney disease or kidney function. The first study of chronic kidney disease used both a retrospective and prospective longitudinal approach (Dhingra 2016a). In the retrospective approach, there were 397 confirmed cases arising from a cohort of 32,254. Of the 397 cases, 187 were non-diabetic. In the prospective approach, the cohort was restricted to those who were kidney disease-free at the baseline interview (2005-2006) and evaluated 212 confirmed cases (106 non-diabetic) that occurred after the baseline interview. Analyses were also conducted restricting the cohort and the kidney disease cases to those without diabetes. The study evaluated yearly modeled PFOA serum concentrations and cumulative exposure. In the full cohort, the estimated hazard ratio (HR) in the top quintile of cumulative serum PFOA in the retrospective analysis was 1.24 (95% CI: 0.88, 1.75), and the trend was not monotonic. In the prospective analysis, the HR in the top quintile was 1.12 (95% CI: 0.72, 1.75) and the trend was also not monotonic. Similar findings were obtained for the non-diabetic population. When exposures were lagged, the HRs were reduced.

In a second C8 study, the phenomenon of "reverse causation" was assessed in a cross-sectional evaluation of impaired kidney function (estimated glomerular filtration rate, eGFR) and earlier menopause (Dhingra 2016b). Self-reported menopause was evaluated among 9,192 women aged 30-65 and kidney function among 29,499 adults. Although there was a non-monotonic negative trend for eGFR across measured serum PFOA quintiles ($\beta \pm S.E.=-0.98 \pm 0.27$ for the top quintile), neither modeled serum PFOA nor modeled cumulative exposure showed associations with eGFR. This suggested that the finding for eGFR was due to reverse causation, i.e., that reduced kidney function as measured by eGFR caused the increased measured serum PFOA. This result would occur because reduced kidney function slowed the excretion of PFOA. The study also found a significant increasing trend of reported early menopause with increasing measured PFOA category, but when using modeled serum or modeled cumulative exposure PFOA instead of measured, this trend disappeared. Again, this suggested reverse causation, i.e., that early menopause caused the increased PFOA serum levels. The study found that measured serum PFOA levels increased on average 4% per year for the first seven years after menopause and then stopped increasing, he authors emphasized that caution is necessary when using exposure biomarkers in cross-sectional studies.

A study using NHANES data for the 2003-2010 cycles evaluated estimated GFR and serum uric acid among adolescents aged 12-19 years (Kataria 2015). PFOA and PFOS serum levels were associated with lower eGFR and higher serum uric acid. However, given the findings in the C8 study, these associations may, at least in part, be due to the phenomenon of reverse causation.

In summary, there is little clear evidence that PFAS affects kidney function or increases the risk of kidney disease. However, because so few studies have been conducted, there is insufficient information to determine whether PFAS exposures affect the kidney. Therefore additional research is needed. In order to avoid the issue of reverse causation, future cross-sectional studies should not rely solely on serum PFAS levels to assess exposures.

Three studies have evaluated uric acid in adults and PFAS exposures. In a C8 cross-sectional study of 54,951 adults, both PFOA and PFOS serum levels were associated with increased uric acid (Steenland 2010). Elevated uric acid (hyperuricemia) was also evaluated, and the OR in the top quintile was 1.47 (95% CI: 1.37, 1.58) with a monotonic trend. PFOS also was associated with hyperuricemia, but to a lesser extent than PFOA. A second cross-sectional study used NHANES data for 1999-2000 and 2003-2006 (serum uric acid was not included in the 2001-2002 NHANES cycle) and found that PFOA and PFOS were associated with hyperuricemia (Shankar 2011). Again, PFOA had the stronger relationship with a fourth quartile OR of 1.97 (95% CI: 1.44, 2.70) and a monotonic trend compared to a fourth quartile OR of 1.48 (95% CI: 0.99, 2.22) for PFOS and a non-monotonic trend. A third cross-sectional study evaluated NHANES data for the 2007-2010 cycles and found that both PFOA and PFOS serum levels were associated with increasing serum uric acid (Gleason 2015). However, the study found that only serum PFOA had a monotonic increased risk of hyperuricemia.

Because these studies relied on serum PFAS levels to assess exposure and were cross-sectional, it is possible that the phenomenon of reverse causation may explain at least part of these findings, i.e., that a reduction in kidney function or chronic kidney disease causes hyperuricemia as well as increased PFAS serum levels due to a reduction in excretion. Nevertheless, the studies consistently found an increased risk of hyperuricemia associated with serum levels of PFOA, and to a lesser extent PFOS. Further research is needed that supplements serum PFAS measurements with modeled serum PFAS estimates to account for possible reverse causation.

2. Liver disease/liver function

Two C8 studies and one study that evaluated NHANES data have assessed PFAS exposures and liver function. The first C8 cross-sectional study included 47,092 adults and measured alanine transaminase (ALT), γ -glutamyltransferase (GGT) and direct bilirubin (Gallo 2012). Both PFOA and PFOS serum levels were associated with increased ALT in linear models and with high ALT in logistic models. The trends were not monotonic. The second C8 study evaluated liver disease among 32,254 adults including 3,713 DuPont workers and liver biomarkers among 30,723 adults including 1,892 DuPont workers (Darrow 2016). The liver biomarker part of the study was cross-sectional. The study avoided the issue of reverse causation by using modeled estimates of yearly serum levels of PFOA. Estimated cumulative exposure and the estimated PFOA serum level during 2005-2006 were evaluated. PFOA was associated with increasing levels of ALT and with abnormal ALT. PFOA was not associated with liver disease. An earlier study conducted in the C8 study area by Emmett 2006 was limited by a small sample size (371 residents including 20 children ages 2-10 and 29 individuals aged 11-20). Another study using NHANES data for the 1999-2006 cycles found no association with liver disease, with 4th quartile ORs for PFOA and PFOS of 0.61 and 0.95, respectively (Melzer 2010).

A study using NHANES data for 2007–2010 found associations between serum PFAS and liver function parameters (Gleason 2015). PFHxS was associated with increased ALT, PFOS was associated with increased total bilirubin, PFOA was associated with increased ALT, GGT and total bilirubin, and PFNA was associated with increased ALT. An earlier study using NHANES data for 1999-2004 found positive associations between PFOA and ALT and natural log GGT and PFHxS and PFNA and total bilirubin (Lin 2010). The association between PFOA and liver enzymes was more evident in obese subjects, as well as subjects with insulin resistance and/or metabolic syndromes.

In summary, the two studies that evaluated PFAS exposures and liver disease found no association. Consistent associations across three studies were found for PFOA and increased ALT. PFOS was also associated with increased ALT in a C8 study. The C8 study that evaluated modeled estimated PFOA serum levels avoided the issue of reverse causation and observed an association between PFOA and increased ALT. Even though the number of studies is small, PFOA has consistently been associated with increased ALT. Because few studies have been conducted, further research is needed to evaluate PFAS exposures and liver function, supplementing serum PFAS measurements with modeled serum PFAS estimates to account for the issue of reverse causation.

3. Coronary Artery Disease, hypertension, hypercholesterolemia (high cholesterol)

One C8 study evaluated incident coronary artery disease and hypertension and modeled PFOA serum levels (Winquist and Steenland 2014b). There was no association with hypertension or coronary artery disease − in both analyses, the HRs were higher for the lower quintiles than the higher quintiles, no HR was higher than 1.26, and most HRs were ≤1.1. For coronary artery disease, the HR for the top quintile of cumulative exposure to PFOA was 1.07 (95% CI: 0.93, 1.23) with a non-monotonic trend. For hypertension, the HR for the top quintile of cumulative exposure to PFOA was 0.98 (95% CI: 0.91, 1.06).

A case-control study of coronary heart disease was conducted in Sweden with 253 cases and 253 matched controls (Mattsson 2015). The adjusted ORs for serum PFOS for the 3rd and 4th quartile were 1.30 (95% CI: 0.74, 2.26) and 1.07 (95% CI: 0.60, 1.92), respectively. The strongest finding in this study was for perfluoroheptanoic acid (PFHpA) with adjusted ORs for the 3rd and 4th quartile of 2.58 (95% CI: 1.39, 4.78) and 1.73 (95% CI: 0.94, 3.16). All of the other PFAS had adjusted ORs in the 4th quartile <1.0.

A study that evaluated NHANES data for 1999-2006 found a slight increase in heart disease for PFOA (4th quartile OR=1.08, 95% CI: 0.70, 1.69) and no association with PFOS (4th quartile OR=0.91) (Melzer 2010). Another study that evaluated NHANES data (1999-2003) found PFOA associated with cardiovascular disease (Shankar 2012). For the top quartile of PFOA, the OR for cardiovascular disease was 2.01 (95% CI: 1.12, 3.60) with a monotonic trend. Elevated ORs were observed for PFOA and both coronary heart disease and stroke but the trend was not monotonic, with 4th quartile ORs of 2.24 (95% CI: 1.02, 4.94) and 4.26 (95% CI: 1.84, 9.89), respectively.

Several studies have evaluated hypercholesterolemia or serum lipids and have found consistent positive associations with PFAS. In the C8 cross-sectional study of 46,294 adults, both PFOS and PFOA were associated with increasing total cholesterol and LDL (Steenland 2009). The predicted increase in total cholesterol from the lowest to highest decile of PFOS and PFOA was 11–12 mg/dL. For hypercholesterolemia, the OR for the top quartile of PFOA was 1.38 (95% CI: 1.28, 1.50) with a

monotonic trend, while for PFOS, the OR was 1.51 (95% CI: 1.40, 1.64), with a monotonic trend. PFOA was also associated with triglycerides. In a longitudinal study of 560 adults from the C8 population who were followed for 4.4 years, individuals with the greater declines in serum PFOA and PFOS had greater LDL decreases (Fitz-Simon 2013). For an individual whose serum PFOA fell by half, the predicted fall in LDL was 3.6% (95% CI: 1.5%, 5.7%). A stronger finding was observed for PFOS: a decline in serum PFOS by half was predicted to decrease LDL by 5% (95% CI: 2.5%, 7.4%). Incident hypercholesterolemia was found to be associated with PFOA in a C8 study, especially among men aged 40-60. The HR for the top quintile of cumulative exposure was 1.44 (95% CI: 1.28, 1.62) although the trend was not monotonic (Winquist and Steenland 2014b).

A study of cholesterol using NHANES data for the 2003-2004 cycle found positive associations for PFOS, PFOA and PFNA but a negative association with PFHxS (Nelson 2010). The highest quartile of PFOS exposure had total cholesterol levels 13.4 mg/dL (95% CI: 3.8–23.0) higher than in the lowest quartile. For PFOA, PFNA, and PFHxS, effect estimates were 9.8 (95% CI, –0.2 to 19.7), 13.9 (95% CI, 1.9–25.9), and –7.0 (95% CI, –13.2 to –0.8), respectively. A study using 2007-2009 cross-sectional data from the Canadian Health Measures Survey found a positive association between PFHxS and hypercholesterolemia (Fisher 2013). For the top quartile of PFHxS, the weighted OR was 1.57 (95% CI: 0.93, 2.64) with a monotonic trend. The finding for PFHxS contradicted the NHANES study which found that PFHxS was associated with a decline in cholesterol. In the Canadian study, the top quartile for PFOA had an OR of 1.50 (95% CI: 0.86, 2.62) with a non-monotonic trend.

In a small study conducted in China, 133 individuals were evaluated for PFAS exposure and high serum lipid levels (Fu 2014). The study did not evaluate PFHxS, but did evaluate PFOA, PFOS, PFNA, perfluorodecanoic acid (PFDA) and perfluoroundecanoic acid (PFUdA). For high total cholesterol, there was a monotonic trend for PFDA with the top quartile OR of 3.84 (95% CI: 0.87, 16.95). Nonmonotonic trends for high total cholesterol were observed for PFOS (4th quartile OR=2.27, 95% CI: 0.47, 10.92) and PFUdA (4th quartile OR=3.70, 95% CI: 0.76, 18.03). For high LDL, monotonic trends were observed for PFOS (top quartile OR=2.27, 95% CI: 0.50, 10.37) and PFUdA (top quartile OR=4.16, 95% CI: 0.96, 18.00). The wide confidence intervals in this study were due to the small sample size.

A study of 891 pregnant women in Norway found a monotonic trend for PFOA and increasing total cholesterol with a regression coefficient of 2.58 (95% CI: -4.32, 9.47) per natural log PFOA (ng/ml) (Starling 2014). Non-monotonic trends were also observed for total cholesterol and PFOS and PFHxS with regression coefficients (per natural log-ng/ml) of 8.96 (95% CI: 1.70, 16.22) and 3.00 (95% CI: -1.75, 7.76), respectively. Monotonic trends were observed for PFOS, PFHxS, and PFUdA and HDL, but no monotonic trends were observed for PFAS and LDL. The strongest finding for LDL was for PFOS with a regression coefficient of 6.48 (95% CI: -0.07, 13.03). Smaller effects were observed between LDL and PFOA and PFHxS with regression coefficients of 2.25 (95% CI: -3.97, 8.48) and 1.92 (95% CI: -2.50, 6.33), respectively. No associations were observed for triglycerides.

In summary, because of the small number of studies and conflicting findings, more research is needed to evaluate whether PFAS exposures affect the risk of cardiovascular disease or hypertension. Several studies have evaluated PFOA and PFOS and lipids and the findings consistently indicate that an association exists with increased lipids.

4. Thyroid function/disease

A C8 study that evaluated thyroid disease in retrospective and prospective analyses included 32,254 adults aged ≥20 years (28,541 community members and 3,713 workers) who completed baseline questionnaires in 2005-2006 and follow-up questionnaires during 2008-2010 and 2010-2011 (Winquist and Steenland, 2014a). About 2/3 of the thyroid diseases were hypothyroidism. In the retrospective analysis, the HR for the top quintile of cumulative exposure to PFOA and functional thyroid disease was 1.28 (95% CI: 1.06, 1.53) with a non-monotonic trend. The finding was much stronger in females with a 5th quintile HR of 1.37 (95% CI: 1.11, 1.68) and a monotonic trend compared to the results in males (5th quintile HR=1.05, 95% CI: 0.66, 1.66, and no trend). PFOA was more strongly associated with hypothyroidism with a HR for the top quintile of 1.40 (95% CI: 1.12, 1.75) and a non-monotonic trend. Elevated HRs were observed for both males and females and hypothyroidism, but there was a monotonic trend for females. Elevated HRs were observed for females and hyperthyroidism, but the trend was not monotonic. HRs were not elevated for males and hyperthyroidism. Similar findings were observed when the community cohort was evaluated separately. However there was a monotonic trend for females and hyperthyroidism, and the HRs for males and hypothyroidism were higher than for females although the trend was not monotonic.

In the prospective analysis, the HR for the top quintile of cumulative PFOA exposure and functional thyroid disease was 1.12 (95% CI: 0.82, 1.52) with a non-monotonic trend. There was no association for females, but there was a monotonic trend for males with an HR for the top quintile of 1.85 (95% CI: 0.93, 3.68). The highest HRs were for hyperthyroidism among males and females although based on relatively small numbers, particularly among males, and the trends were not monotonic. For hypothyroidism, there was no association among females, but there was a monotonic trend for males with an HR for the top quintile of 2.02 (95% CI: 0.87, 4.65). Similar findings were observed when the community cohort was evaluated separately. The authors concluded that there was an association between PFOA exposure and thyroid disease, especially for hyperthyroidism among women in the retrospective analyses and for hypothyroidism among men in the prospective analyses.

A study that evaluated NHANES data for 1999-2006 found associations between PFOA and PFOS and reported ever had thyroid disease and reported current thyroid disease with medication (Melzer 2010). None of the exposure-response trends were monotonic. For the 4th quartile PFOA, the ORs for ever had thyroid disease were 1.68 (95% CI: 1.14, 2.49) for women and 1.50 (95% CI: 0.66, 3.39) for men. For current thyroid disease with medication, the ORs for 4th quartile PFOA were 2.24 (95% CI: 1.38, 3.65) for women and 2.12 (95% CI: 0.93, 4.82) for men. For the 4th quartile PFOS, the ORs for ever had thyroid disease were 1.15 (95% CI: 0.78, 1.70) for women and 1.78 (95% CI: 0.58, 5.52) for men. For current thyroid disease with medication, the ORs for 4th quartile PFOA were 1.27 (95% CI: 0.82, 1.97) for women and 2.68 (95% CI: 1.03, 6.98) for men.

In summary, although only two studies have evaluated thyroid disease in adults and PFAS exposure, both had positive findings. In particular, the C8 study found elevated HRs for thyroid diseases in the prospective analyses. However, there also appears to be effect modification by gender. Because of the few studies that have evaluated PFAS exposure and thyroid disease, more research is necessary, in particular, studies designed to evaluate effect modification by gender.

Thyroid function biomarkers were evaluated in a C8 cross-sectional study that included 52,296 adults with a year or more exposure to PFOA (Knox 2011). The biomarkers evaluated were thyroxine (T4), T3 uptake, and thyroid stimulating hormone (TSH). Both PFOA and PFOS were associated with an

elevation in serum thyroxine and a reduction in T3 uptake. Interactions between gender and PFOS were observed for T3 uptake and thyroxine and between PFOA and gender for T3 uptake.

Three studies evaluated NHANES data and thyroid function. The first study evaluated NHANES data for 2007-2008 and found that TSH and TT3 levels increased with PFOA and TT4 levels increased with PFHxS (Jain 2013). This study is not included in Table A3 because no confidence intervals were presented, the age range included adolescents (i.e., ages 12-18), and these NHANES data were evaluated by a second study that evaluated NHANES data for 2007-2010 (Wen 2013). The latter study found mixed results by gender (Wen 2013). PFHxS was associated with an increase in TT4 for women (β = 0.26, 95% CI: 0.11, 0.41) but not for men (β = -0.03, 95% CI: -0.18, 0.64). On the other hand PFHxS was associated with a decline in T4 among men (β = -0.016, 95% CI: -0.029, -0.003) but not for women (β = 0.003, 95% CI: -0.024, 0.030). PFHxS was associated with an increase in TT3 for women β = 4.07, 95% CI: 2.23, 5.92) but not for men (β = -0.08, 95% CI: -1.70, 1.56). PFOA was associated with TT3 among women (β = 6.63, 95% CI: 0.55, 12.72) but not for men (β = 0.78, 95% CI: -3.05, 4.60).

The third NHANES study evaluated NHANES data for 2007-2008 but focused on potential susceptible subgroups with thyroid "stressors", i.e., with low iodine status, with high thyroid peroxidase antibody (TPOAb) or with both (Webster 2016). The key findings were that all 4 PFAS evaluated (PFOA, PFOS, PFHxS and PFNA) were associated with increased fT3, increased fT3/fT4, increased TSH, and increased TT3 in the group with joint exposure to high TPOAb and low iodine. PFOS and PFHxS were also associated with decreased fT4 in the group with high TPOAb and low iodine. The findings were considerably weaker for those with normal iodine and TPOAb status and those with either low iodine alone or high TPOAb alone.

A group of 87 men and women residing in the upper Hudson River area of New York who were originally recruited for a study of PCB exposure were evaluated for PFOA and PFOS exposure and thyroid function (Shrestha 2015). Natural log PFOS was positively associated with fT4 (β =0.054, 95% CI: 0.002, 0.106) and T4 (β =0.766, 95% CI: 0.327, 1.205), which corresponded to 4% and 9% increases in fT4 and T4 per interquartile range difference in PFOS. A positive association also was observed for log PFOS and TSH (β =0.129, 95% CI: -0.023, 0.281). An association was also observed for natural log PFOA and T4 (β =0.380, 95% CI: -0.070, 0.830). When both PFOS and PFOA were included in the models, the association between PFOS and T4 persisted, but the association between PFOS and fT4 was attenuated.

A cohort of 633 individuals aged >12 years from Siheung, Korea was evaluated for PFAS exposure and thyroid function (Ji 2012). Slight declines in T4 (e.g., for PFOS, β = -0.021, 95% CI: -0.048, 0.005) and slight increases in TSH (e.g., for PFNA, β = 0.110, 95% CI: -0.035, 0.255) were observed. The strongest findings were for perfluorotridecanoic acid (PFTrDA) and decreased T4 and increased TSH.

In summary, based on findings from these studies, there appears to be gender differences in the effects of PFAS exposures on thyroid function. This is not surprising given the evidence of effect modification by gender for the associations between PFOS and PFOA and thyroid diseases. In general, TSH and TT4 increased with PFOA, PFOS and PFHxS exposure. TT3 also appeared to increase but not in the C8 study. A susceptible population with low iodine status and high TPOAb status was identified in one study. Although several studies have been conducted of PFAS exposure and adult alterations in thyroid function, there are inconsistencies in the findings for TT3 and TT4. Additional research is needed to resolve these inconsistencies. In particular, studies should be designed to evaluate effect modification by gender as well as by vulnerable subpopulations such as those who have thyroid "stressors".

Several studies evaluated thyroid function among pregnant women primarily because of the potential effect on the fetus of maternal thyroid hormone disruption. A study in Taiwan of 285 women in their third trimester observed the strongest findings for PFNA, perfluorododecanoic acid (PFDoDA), and perfluoroundecanoic acid (PFUnDA) and decreased maternal free T4 and TT4. The effects of PFHxS on free T4 and TT4 were considerably weaker than these three PFAS but much stronger than PFOA or PFOS (Wang 2014). In a Canadian study of 152 women, those with normal TPOAb had little effects from PFAS exposure on fT4, TT4 or TSH (Webster 2014). However, women with high TPOAb had increases of 46% to 69% in TSH for interquartile range increases of PFOA, PFOS, and PFNA. (Interquartile range increase in PFHxS was associated with only a 2% increase in TSH in these women.) All four PFAS evaluated were associated with a 3% – 7% decrease in fT4 among women with high TPOAb. No associations were found for TT4.

In a study of 375 women in Norway, 4th quartile PFOS levels were associated with a 0.35 (95% CI: 0.21, 0.50) mean difference in TSH, corresponding to a 24% higher mean concentration of TSH (Berg 2015). A multipollutant assessment of persistent organic pollutants (POPs) including PCBs, DDT, hexachlorobenzene, and PFAS was conducted with a similar group of Norway women, and PFOS was again found to be associated with increased TSH with a monotonic trend controlling for other POPs (Berg 2017). A study of 392 women from Hokkaido, Japan found a negative correlation between PFOS and TSH and a smaller positive correlation with fT4 (Kato 2016). There was little correlation between PFOA and TSH or fT4.

In general, the studies of maternal thyroid function and PFAS exposure suggested associations between PFAS and increased TSH. For the other thyroid biomarkers, the results were mixed.

5. Autoimmune diseases

A C8 study evaluated autoimmune diseases retrospectively and prospectively in a cohort of 32,254 adults including 3,713 workers (Steenland 2013). Self-reported autoimmune diseases were confirmed via medical records. For the retrospective analyses, the follow-up was from 1952 through the interviews conducted in 2008-2011. For the 4th quartile cumulative PFOA exposure, the RRs for ulcerative colitis for unlagged and 10-year lagged exposures were 2.86 (95% CI: 1.65, 4.96) and 3.05 (1.56, 5.96), respectively. The trends for ulcerative colitis were monotonic. For multiple sclerosis, the 4th quartile exposure RRs, unlagged and 10-year lagged, were 1.26 (0.65, 2.42) and 1.32 (0.61, 2.84), respectively, but the trends were not monotonic. For rheumatoid arthritis, an elevated RR for 4th quartile PFOA exposure was observed only for the 10-year lagged exposure (RR=1.35, 95% CI: 0.87, 2.11), but the trend was not monotonic. In the prospective analyses, autoimmune cases that occurred between the baseline 2005-2006 interviews and the 2008-2011 follow-up interviews were too few to evaluate multiple sclerosis, lupus, type 1 diabetes or Crohn's disease.

For ulcerative colitis, the RRs for the 4th quartile cumulative PFOA exposure, unlagged and 10-year lagged, were 1.62 (95% CI: 0.57, 4.61) and 1.51 (95% CI: 0.43, 4.30). The wide confidence intervals were due to the small number (n=30) of cases that occurred during the follow-up period. The trends were not monotonic. The RRs for rheumatoid arthritis for the 4th quartile PFOA cumulative exposure were <1.0.

On the basis of this study, the C8 Science Panel decided that there was a probable link between PFOA exposure and ulcerative colitis. Of note, the RRs were elevated in both the retrospective and prospective analyses. A study of ulcerative colitis conducted of the 3,713 workers at the DuPont West Virginia plant

found a RR of 6.57 (95% CI: 1.47, 29.4) for the top quartile of cumulative PFOA exposure and a 10-year exposure lag. The trend was monotonic and the findings were based on 28 total cases. For rheumatoid arthritis and no exposure lag, the RR for the top quartile was 4.45 (95% CI: 0.99, 19.9) with a monotonic trend based on 23 total cases (Steenland 2015, see Table A2).

The C8 studies indicate an association between PFOA exposure and ulcerative colitis. For the other autoimmune diseases, the information is inadequate to determine whether PFOA is associated with increased risk. For the other PFAS, there is a lack of information on the risk for autoimmune diseases. Research is needed to determine whether PFAS exposures increase the risk of autoimmune diseases, especially since there is toxicological evidence that PFAS exposures affect the immune system (NTP 2016).

6. Osteoarthritis, osteoporosis and bone mineral density

Two studies evaluated osteoarthritis. In a C8 study, 49,432 adults were included and 3,731 reported a physician diagnosed case of osteoarthritis in the baseline survey (Innes 2011). For the top quartile of serum PFOA levels, the OR for osteoarthritis was 1.42 (95% CI: 1.26, 1.59) with a monotonic trend. No association (ORs < 1.00) was observed for serum PFOS. Higher ORs were observed for PFOA and osteoarthritis among those aged <55 years and those who were not obese. In the NHANES study, data for 2003-2008 were evaluated (Uhl 2013). For PFOA, the OR for the top quartile was 1.55 (95% CI: 0.99, 2.43) with a non-monotonic trend. The elevated ORs were entirely due to the effect in females, since the ORs for males were <1.00. For the top quartile of serum PFOS, the OR was 1.77 (95% CI: 1.05, 2.96) with a non-monotonic trend. Elevated ORs were observed in both males and females.

Two studies evaluated bone mineral density using NHANES data. The first study used NHANES data for 2005-2008 to evaluate serum PFOS and PFOA and lumbar spine and hip bone mineral density among men, women in menopause and premenopausal women (Lin 2014). Self-reported fractures were also evaluated. A unit increase in natural log serum PFOS was associated with a decrease in total lumbar spine bone mineral density by 0.022 g/cm² (95% CI: -0.038, -0.007) in women not in menopause, but the trend was not monotonic and the decline was only observed among those with PFOS serum levels >75th percentile. No associations were observed for PFOA, and no associations were observed for total hip bone mineral density. For all types of self-reported fractures, the ORs for women in menopause were 1.53 (95% CI: 0.63, 3.74) for PFOA and 1.59 (95% CI: 0.88, 2.86) for PFOS. No associations were found for all types of fractures among men or premenopausal women. Premenopausal women had elevated ORs for PFOA and hip fracture (OR=1.59, 95% CI: 0.57, 4.46) and spine fracture (OR=1.83, 95% CI: 0.59, 5.61). Men had an elevated OR for PFOA and spine fracture (OR=1.54, 95% CI: 0.85, 2.79).

The second NHANES study used data for 2009-2010 to evaluate bone mineral density and osteoporosis (Khalil 2016). Both sexes had declines in femur bone mineral density for each of the PFAS evaluated (PFOA, PFOS, PFHxS and PFNA). The strongest association was for PFOS among postmenopausal women (natural log-PFOS β = -0.033, 95% CI: -0.049, -0.015). For femur neck mineral density, declines were seen for PFOS, PFHxS and PFNA in both sexes and for PFOA among premenopausal women. The strongest association was for PFOS among postmenopausal women (natural log-PFOS β = -0.033, 95% CI: -0.049, -0.017). For lumbar spine bone mineral density, declines were observed for PFOA and PFOS among men and postmenopausal women, PFNA and both sexes, and PFHxS and postmenopausal women. The strongest association was for PFNA among postmenopausal women (natural log-PFNA β = -0.043, 95% CI: -0.073, -0.013). For osteoporosis, the 4th quartile ORs for PFOA, PFOS, PFHxS and

PFNA were 2.59 (95% CI: 1.01, 6.67), 1.07 (95% CI: 0.36, 3.19), 13.20 (95% CI: 2.72, 64.15) and 3.23 (95% CI: 1.44, 7.21), respectively. None of the trends were monotonic except possibly for PFOA where the ORs for the 2nd and 3rd quartiles were essentially the same and could be considered monotonic.

In summary, some positive findings were observed in the two studies that evaluated bone mineral density and the two studies that evaluated osteoarthritis. Only one study evaluated osteoporosis and positive findings were observed as well. Because only a few studies have been conducted, additional research is needed.

7. Immune Response

Four studies have evaluated immune response or immune biomarkers. A report by the C8 Science Panel on a cross-sectional study that has not yet been published evaluated immune biomarkers such as IgG, IgM, IgA, IgE, total antinuclear antibodies (ANA) and C reactive protein (CRP) among the C8 adult population (C8 Science Panel 2009). The study included 56,315 adults. The Panel reported that "several statistically significant associations between levels of immunoglobulins and C8 were found: For IgA the pattern of association indicated a significant decreasing trend with increasing PFOA; this was also apparent for IgE but only in females. For IgG there was not a consistent trend with PFOA. ANA shows a positive significant relationship with increasing PFOA. CRP showed a strong downward trend with increasing PFOA,"

A second C8 study evaluated influenza vaccine response in 403 adults who did not have influenza within the last 3 months and who provided pre- and post-vaccination blood samples to determine virusspecific antibody titers (Looker 2014). Associations between PFOA and PFOS and self-reported influenza and colds in the past 12 months as reported on questionnaires (n = 755) were also assessed. Elevated PFOA serum concentrations (4th vs 1st quartiles of exposure) were associated with reduced antibody titer rise which may correlate with an increased risk of not attaining the antibody threshold considered to offer long-term protection. Small negative associations (regression coefficients for the geometric mean antibody titer ranging from -0.03 to -0.22) were observed comparing the highest and lowest quartiles of PFOA and antibody titer rise and ratios for influenza B and Influenza A/H3N2 and antibody titer rise for Influenza A/H1N1; there was a monotonic exposure response relationship for H1N1. For PFOS, small negative associations were observed comparing the highest and lowest quartiles and antibody titer rise and ratios for Influenza A/H3N2 (-0.04 and -0.03, respectively). People exposed to the highest quartile PFOA and PFOS were less likely to seroconvert following vaccinations for Influenza B (ORs = 0.71 and 0.87, respectively). Additionally, people exposed to the highest quartile of PFOS were less likely to seroconvert following vaccinations for Influenza A/H1N1 (OR = 0.94) and people exposed to the highest quartile of PFOA were less likely to seroconvert following vaccinations for Influenza A H3N2 (OR = 0.62). OR for cold or flu ranged from 1.09-1.20 for people exposed to the highest quartile of PFOS.

An exploratory study measured serum-PFAS concentrations in 12 adults whose antibody responses were followed for 30 days after a booster vaccination with diphtheria and tetanus (Kielsen 2016). Participants were healthy volunteers from a hospital in Denmark. Diphtheria antibody concentrations post-vaccination were decreased by 8.2 to 18.2% for a doubling of exposures to several PFASs, including PFOA, PFOS, and PFHxS. Tetanus antibody concentrations were decreased by 4.4 to 10.8% for a doubling of exposures to several PFASs, including PFOS and PFHxS (but not PFOA). The authors note that "serum PFAS concentrations showed significant negative associations with the rate of increase in

the antibody responses." The authors also noted that this effect was particularly strong for the longer-chain PFAS such as PFDA and PFNA.

Another exploratory study included 78 adults and evaluated the immune response to vaccination with FluMist intranasal live attenuated influenza vaccine and PFAS exposure (Stein 2016b). Between 9% and 25% of the adults seroconverted after vaccination. PFAS exposure was associated with seroconversion but the small numbers that had seroconverted resulted in extremely wide confidence intervals. The strongest association observed between PFAS and immune marker response was for PFHxS and lower mean interferon-γ (IFN-γ) and tumor necrosis factor-α (TNF-α) levels. The second and third tertile regression coefficients for PFHxS and IFN-γ were -40 (95% CI: -76, -3.7), and -40 (95% CI: -84, 2.69). For TNF-α, the second and third tertile regression coefficients were -5.3 (95% CI: -9.2, -1.3) and -4.8 (955 CI: -9.4, -0.10). However the authors concluded that the findings did not support an association between PFAS exposure and reduced immune response to FluMist vaccination, although the study had severe limitations including small sample size and the limited antibody response to FluMist.

Although there were positive findings in the four studies that have been conducted, each study had serious limitations. Therefore additional research is needed with improved study designs and sufficient statistical power to evaluate PFAS exposures and immune response.

8. Reproductive outcomes

Table A4 provides details on the studies of subfertility and infertility. Six studies evaluated subfertility (time to pregnancy [TTP]) and/or infertility. Positive associations were found for subfertility/delayed TTP for PFOA in three studies (Bach 2015a, Fei 2009, Whitworth 2012a) and PFOS in four studies (Bach 2015a, Fei 2009, Jorgenson 2014, Whitworth 2012a), and slightly positive associations were found for PFOA, PFHxS, and PFOS in one study (Velez 2015) and for PFNA in one study (Jorgenson 2014). Positive associations were found for infertility and PFOA in two studies (Fei 2009, Velez 2015), for PFOS in two studies (Fei 2009, Jorgenson 2014), and for PFHxS and PFNA in one study each (Jorgenson 2014, Velez 2015), and slightly positive associations were found for PFOS and PFOA in one study each (Jorgenson 2014, Velez 2015). An additional study assessed adult male semen quality, testicular volume, and reproductive hormone levels in men who were exposed in utero to PFOS and PFOA and found positive associations with these chemicals and outcomes (Vested 2013).

A systematic review assessed fertility by evaluating studies on reproductive hormones (10 studies in men and 3 in women) and TTP (2 studies in men and 8 in women) as well as nine studies of semen characteristics (Bach 2016a). In men, there were inconsistent results across PFAS studies of semen volume, sperm concentration, total sperm count, motility, and morphology; levels of testosterone, free androgen index/free testosterone, estradiol, SHBG, LH, FSH, and inhibin B; and TTP. In women, studies showed mostly positive associations for PFOS and PFOA and infertility and fecundability in parous women, but were inconsistent for PFAS and reproductive hormones. The review concluded that there was not strong evidence for an association between PFAS exposures and reproductive outcomes in men or women. Another review of the epidemiological literature on reproductive outcomes came to a similar conclusion (Khalil 2015).

Table A5 provides details on the studies of pre-eclampsia and pregnancy-induced hypertension. Five studies evaluated pre-eclampsia. Positive associations were found for pre-eclampsia and PFOA (Savitz 2012a) and PFOS (Stein 2009) and slightly positive associations were found with PFOA in two studies

(Avanasi 2015, Nolan 2010) and PFOS in one study (Starling 2014). Three studies evaluated pregnancy-induced hypertension (PIH). Positive associations were found for PIH and PFOA in three studies (Darrow 2013, Nolan 2010, Savitz 2012b) and PFOS in one study (Darrow 2013). The C8 Science Panel concluded that there was a probable link between PFOS exposure and pregnancy-induced hypertension.

Two studies evaluated PFAS exposure and endometriosis. The first study included a sample of 495 women aged 18-44 scheduled for laparoscopy/laparotomy in clinics located in Salt Lake City and San Francisco (Buck Louis 2012). The second sample was a population-based sample of 131 women matched to the first sample on age and residence within a 50-mile radius of the participating clinics. Forty-one percent (n=190) of the women scheduled for laparoscopy ("operative sample") had newly diagnosed endometriosis and 11% (n=14) of the population-based sample had endometriosis. The ORs for PFOS, PFOA, PFHxS and PFNA among the operative sample were 1.25 (95% CI: 0.87, 1.80), 1.62 (95% CI: 0.99, 2.66), 0.85 (0.42, 1.73), and 1.99 (95% CI: 0.91, 4.33). Among those cases with stage 3 or stage 4 disease, the ORs for PFOS, PFOA, PFHxS and PFNA were 1.50 (95% CI: 0.82, 2.74), 1.86 (95% CI: 0.81, 4.24), 1.24 (0.47, 3.31), and 0.99 (95% CI: 0.27, 3.65).

The second study evaluated NHANES data for 2003-2006 (Campbell 2016). Seven percent of the sample of 753 women reported physician-diagnosed endometriosis (n=54). Fourth quartile ORs for PFOA, PFOS, PFHxS and PFNA were 2.86 (95% CI: 0.63, 12.91), 3.48 (95% CI: 1.00, 12.00), 1.47 (95% CI: 0.40, 5.41), and 3.24 (95% CI: 0.81, 12.91). None of the trends were monotonic.

Both studies of PFAS exposures and endometriosis had positive findings. However, since only two studies have been conducted, more research is needed to determine whether PFAS exposures increase the risk of endometriosis.

Summary of the Literature Review for adult diseases

For most adult diseases there is little or no information on the effects of exposures to PFHxS. Although there is more information for PFOS, it is still inadequate to determine whether exposures increase the risk for most of the adult diseases. PFOA has the most information, primarily because of the C8 studies. Still additional research is needed to determine whether PFOA exposures increase the risk of several adult cancers and non-cancers including colorectal cancer, multiple myeloma, kidney function/kidney diseases, liver function, autoimmune diseases other than ulcerative colitis, and immune function.

Health Effects of PFAS in Children

Tables A6 and A7 provides details of studies of adverse birth outcomes and congenital malformations. Table A8 provides details of studies of other adverse outcomes in children aged ≥ 2 years.

1. Adverse birth outcomes

Table A6 provides details on the studies that evaluated adverse birth outcomes. Twenty studies evaluated birth weight and the results are presented in Table A6. Additionally, two meta-analysis of birthweight found an overall decrease in birthweight associated with PFOA and PFOS (Verner 2015, Bach 2015b). Ten studies evaluated preterm birth. Five studies evaluated small for gestational age (SGA). Ten studies evaluated birth length, seven evaluated head circumference, one evaluated

abdominal circumference, and eight evaluated gestational age as a continuous variable. As evident from Table A6, virtually all of the studies of these adverse birth outcomes evaluated PFOA and PFOS but only a minority evaluated PFHxS.

Table A7 presents the results of five studies that evaluated birth defects. Positive associations with PFOA and club foot, heart defect, and circulatory defect were found in one study with very small numbers of cases (Nolan 2010). One study found a positive association between PFOS and PFOA (slight) and cryptorchidism (Jensen 2014) while another study found no associations with cryptorchidism and hypospadias (Toft 2016). One study relying on maternally-reported cases found positive associations with PFOA and defects of the brain, limb, eye, and heart (Stein 2014a).

A study on congenital hypothyroidism found a "large" difference in PFOA concentrations between cases and controls and mean concentrations of PFOA and PFNA in cases were "significantly higher" than in controls, but the study was based on small numbers (Kim 2016).

One study evaluated cerebral palsy (Liew 2014). Using case-cohort sampling of the Danish National Birth Cohort during 1996-2002, the study evaluated 156 cases and 550 controls. Maternal serum PFAS were associated with cerebral palsy for boys, in particular PFOS (RR per ln unit increase=1.7, 95% CI: 1.0, 2.8) and PFOA (RR=2.1, 95% CI: 1.2, 3.6). However risks were not elevated in girls except for PFHxS (RR=1.1, 95% CI: 0.6, 1.9).

Five studies evaluated miscarriage and two studies evaluated stillbirth. Positive associations were found for miscarriage and PFOS in two studies (Jensen 2015, Darrow 2014); PFNA, PFDA, and PFHxS in one study (Jensen 2015); and PFOA (slight) in one study (Darrow 2014). No associations were found for stillbirth.

Because of inconsistencies in the findings across studies, more research is needed to evaluate the effect of PFAS exposures on birth weight, SGA, head circumference and other fetal growth parameters, reduced gestational age, and preterm birth. Few studies have been conducted of PFAS exposures and miscarriage, stillbirth, birth defects, congenital hypothyroidism or cerebral palsy, therefore additional research is necessary with improved study designs and sufficient statistical power.

2. Lipids

Table A8 presents the results of the five studies, including a C8 study, that evaluated lipids (Frisbee 2010, Nelson 2010, Geiger 2014a, Maisonet 2015a, Zeng 2015). All five studies found increases in lipids with increasing exposures to PFOS and/or PFOA. In one NHANES study (Nelson 2010), PFHxS was associated with increased lipids. Overall, the findings of increased lipids from exposures to PFAS has been consistent.

3. Thyroid function

There have been several studies conducted of infants and most have observed that prenatal PFAS exposures disrupted thyroid function. Only two studies have been conducted of older children, and their results are presented in Table A8. The C8 study found increases in a thyroid hormone (TSH [thyroid stimulating hormone]) with increasing serum levels of PFOS and PFOA (Lopez-Espinosa 2012). PFHxS was not evaluated. PFOA, but not PFOS, was associated with an increased risk of thyroid disease.

In a Taiwan study (Lin 2013), findings were inconsistent for PFOS and PFOA and thyroid hormones when boys and girls were evaluated separately (effect was stronger in males). PFHxS was not evaluated.

Because of the few studies that evaluated PFAS exposures and thyroid function among children older than infants, more research is needed.

4. Uric acid

Table A8 presents the results of studies that evaluated uric acid. A study in Taiwan found elevated uric acid levels associated with PFOA, PFHxS, and PFOS (Qin 2016). Two studies that evaluated NHANES data found elevations in uric acid and in the risk of hyperuricemia for PFOA and PFOS (Geiger 2013, Kataria 2015). More research is needed to follow up these findings. Because these studies were cross-sectional, there is concern about the possibility of reverse causation (e.g., impaired kidney function could cause both elevated uric acid and a reduction in the elimination of PFAS via the kidney resulting in higher serum PFAS levels). Future studies should attempt to predict serum PFAS levels and/or evaluate this outcome longitudinally (prospectively).

5. Sex hormones

Four studies evaluated sex hormones and are presented in Table A8. The C8 study found declines for testosterone in boys and girls with increasing serum levels of PFOA and PFOS and decline in boys with increasing serum levels of PFHxS (Lopez-Espinosa 2016). In the larger of the two Taiwan studies, declines in testosterone levels were observed in both sexes with increasing serum levels of PFOA, in boys only with increasing serum levels of PFOS, and in girls only with increasing serum levels of PFHxS (Zhou 2016). On the other hand, a study of girls in the UK observed increases rather than decreases in testosterone with increasing serum levels of PFOS, PFOA and PFHxS (Maisonet 2015b).

Overall, there is some evidence that PFAS exposure may decrease testosterone levels, but the findings have not been consistent across the few studies that have been conducted. More research is needed to determine whether and how PFAS exposures affect sex hormone levels.

6. Delay in reaching puberty

Three studies evaluated delays in reaching puberty and are presented in Table A8. In the C8 study, both PFOA and PFOS were associated with delays in puberty (Lopez-Espinosa 2011). PFHxS was not evaluated. In a study conducted in Denmark, PFOS and PFOA were also associated with delay in reaching puberty (Kristensen 2013). However, a study conducted in the UK found that PFOA was associated with an earlier age at puberty while PFOS was associated with delayed puberty, and the results were conflicting for PFHxS (Christensen 2011). More research is needed to evaluate whether PFAS exposure can cause delays in reaching puberty.

7. Neurobehavioral outcomes

Neurobehavioral outcomes are presented in Table A8. Two studies evaluated IQ. The C8 study found only slight differences in IQ and the results were not consistent for PFOA, the only PFAS evaluated (Stein 2013). A study conducted in Taiwan found deficits in IQ for PFOS, but not for PFOA or PFHxS (Wang 2015).

There is some evidence from the C8 study (Stein 2011) and studies conducted in Denmark (Liew 2015), Sweden (Ode 2014), and using NHANES data (Hoffman 2010) that PFAS may be associated with ADHD. The C8 study found no association for PFOA, but elevated risks for PFOS and PFHxS with odds ratios (ORs) of 1.3 and 1.6, respectively (Stein 2011).

The C8 study found a slight increase in risk (OR=1.2) for higher exposures to PFHxS and learning problems but no associations for PFOS or PFOA (Stein 2011). A recent study that evaluated measures of executive function of "clinical relevance" found elevated risks especially for PFOS and PFHxS (Vuong 2016). However, in general, the effects observed have not been large for neurobehavioral outcomes. Evaluating the evidence for associations between PFAS exposures and IQ, ADHD, and other neurobehavioral outcomes is hampered by different methods for ascertaining ADHD, different methods for testing IQ, lack of consistency in the other neurobehavioral outcomes evaluated, and the small number of studies that have been conducted. Therefore additional research is necessary to determine whether PFAS exposures are associated with adverse neurobehavioral outcomes in children such as IQ, depression, deficits in executive function, ADHD, and developmental delay,

8. Immune function

Few studies have been conducted to evaluate immune function and PFAS exposure. These studies are presented in Table A8. Three studies have evaluated whether PFAS exposures suppress the antibody response to specific vaccines, but only two of these studies evaluated the same vaccine, i.e., rubella. Both of these studies found deficits in serum rubella antibody response (Granum 2013, Stein 2016). The studies in the Faroes have evaluated tetanus and diphtheria longitudinally and found deficits in antibody to these vaccines (Grandjean 2012, 2016).

Asthma was evaluated in three studies, with strong risks found in a Taiwan case-control study (Dong 2013) but considerably weaker risks found in two NHANES studies (Humblet 2014, Stein 2016). Other outcomes such as atopic dermatitis and infectious diseases such as gastroenteritis and the common cold were not evaluated in more than one study.

A systematic review of the evidence for immunotoxicity associated with exposures to PFOA and PFOS conducted by NTP concluded that these exposures alter immune function in humans but that the epidemiological evidence was too limited to conduct meta-analyses. Issues include the heterogeneity of the studies and the small number of studies that evaluated the same outcome. NTP concluded that more research is needed to evaluate the same vaccines and hypersensitivity-related outcomes in children across different populations using similar research methods.

9. Hypertension and adiposity

Studies of hypertension and adiposity are presented in Table A8. One study has evaluated hypertension in children using NHANES data and found no elevation in risk (Geiger 2014b). Three studies have evaluated adiposity in children, adolescents and/or young adults. In one study, an association between PFAS and measures of adiposity was found only in girls (Mora 2016). A second study (Karlsen 2016) found slightly elevated risks for being overweight except for PFOA among children aged 5 years where a stronger risk was observed (OR = 1.88, 95% CI: 1.05, 3.35). In a third study, PFOS was found to be associated with measures of adiposity (Domazet 2016). Additional research is needed to determine whether PFAS exposures increase the risk of hypertension or adiposity in children.

Summary of the Literature Review for childhood diseases

For most adverse outcomes in children evaluated in this assessment, the information was inadequate to determine whether children exposed to PFAS were at increased risk. In particular, very few studies have been conducted of PFHxS exposures, the PFAS that was considerably elevated in the serum of the children tested at Pease. Additional research is needed for PFAS exposures and adverse birth outcomes; thyroid, liver, kidney and immune function; uric acid; sex hormones; delays in reaching puberty; ADHD and other neurobehavioral outcomes; hypertension; and adiposity.

Description of sample size calculations

Sample size calculations were conducted using OpenEpi Version 3.03. (Dean AG, Sullivan KM, Soe MM. OpenEpi: Open Source Epidemiologic Statistics for Public Health, www.OpenEpi.com, updated 2014/09/22). For some health-related endpoints, calculations could not be conducted because of a lack of information in the studies on the parameters needed to make the calculations.

Sample size calculation for mean difference:

$$N_1 =$$
(variance of group 1 + variance of group 2 / (N_2/N_1)) $(Z_{1-\alpha/2} + Z_{1-\beta})^2$ (Mean difference)²

Where N_2/N_1 is the ratio of the two sample sizes. Then N_2 is simply this ratio multiplied by N_1 . For a type 1 error (or α error) of .05, the $Z_{1-\alpha/2}$ value is 1.96. This calculation is for a two-tailed hypothesis test and equivalent to using a 95% confidence interval to determine statistical significance. For a one-tail test with α =.05, the $Z_{1-\alpha/2}$ in the above equation is replaced by $Z_{1-\alpha}$ and its value is 1.65, equivalent to using a 90% confidence interval to determine statistical significance. The $Z_{1-\beta}$ in the above equation is the $Z_{1-\beta}$ value for the selected power. For 80% power, $Z_{1-\beta}$ = 0.84, for 90% power, $Z_{1-\beta}$ = 1.28, and for 95% power, $Z_{1-\beta}$ = 1.65. (See Rosner B. Fundamentals of Biostatistics, 7th Edition, equation 8.27, p. 302).

The sample size calculations for odds ratios, risk ratios, etc. are as follows:

The sample size formula without the correction factor by Fleiss is:

$$n_{1} = \frac{\left[Z_{m2}\sqrt{(r+1)\overline{pq}} + Z_{1-\beta}\sqrt{rp_{1}q_{1} + p_{2}q_{2}}\right]^{2}}{r(p_{1} - p_{2})^{2}}$$

$$n_{2} = r n_{1}$$

For the Fleiss method *with* the correction factor, take the sample size from the uncorrected sample size formula and place into the following formula:

$$n_{kc} = \frac{n_1}{4} \left[1 + \sqrt{1 + \frac{2(r+1)}{n_1 r |p_2 - p_1|}} \right]$$

$$n_{2rr} = r n_{1rr}$$

When the input is provided as an odds ratio (OR) rather than the proportion of exposed with disease, the proportion of exposed with disease is calculated as:

$$p_1 = \frac{p_2OR}{1 + p_2(OR - 1)}$$

When the input is provided as a risk (or prevalence) ratio (RR) rather than the proportion of exposed with disease, the proportion of exposed with disease is calculated as:

$$p_1 = p_2 RR$$

Fleiss JL. Statistical Methods for Rates and Proportions. John Wiley & Sons, 1981.

The following provides information on the parameters (e.g., standard deviation, disease prevalence) used in the sample size calculations provided in Table 6 for 350 Pease children and 175 unexposed children. (Sample size calculations for odds ratios or risk ratios used the formula with the correction factor.)

Lipids

In the C8 study (Frisbee 2010), the mean total cholesterol level in the study population was 160.7 mg/dL and the standard deviation (SD) was 29.3. The sample size calculations assumed the same SD in the Pease children and the unexposed group. For hypercholesterolemia (total cholesterol \geq 170 mg/dL), the prevalence in the C8 study was 34.2%.

Uric Acid

In the NHANES study (Geiger 2013), the mean uric acid level in the study population was 5.07 mg/dL with a SD of 1.19. The sample size calculations assumed the same SD in the Pease children and the unexposed group. The prevalence of hyperuricemia (uric acid ≥ 6 mg/dL) in the NHANES study was 16%.

Kidney Function

The mean estimated glomerular filtration rate (eGFR) in the C8 study of children and adolescents (Watkins 2013) was 133 mL/min/1.73 m² with a SD of 23.9. The sample size calculations assumed the same SD in the Pease children and the unexposed group.

Attention Deficit/Hyperactivity Disorder (ADHD)

In the C8 study (Stein 2011), the prevalence of participant-reported ADHD was 12.4% and the prevalence for participant-reported + used medications for ADHD was 5.1%. Sample size calculations used the 12.4% prevalence. (Using the 5.1% prevalence would require much larger sample sizes.)

Hypersensitivity-related Outcomes

From an NHANES study (Stein 2016), the prevalences of current asthma and rhinitis among those aged 12-19 were 10.9% and 25.6%, respectively. For atopic dermatitis, the prevalence for children and adolescents (ages 5-17) is about 12% based on data from the National Health Interview Survey.

The following sample size calculations were conducted using the minimum detectable effect levels seen in the C8 and other studies that corresponded to similar serum levels of PFAS as observed among the Pease children and adults.

Sex hormones and Insulin-like growth factor – 1 (IGF-1)

C8 study of children (Lopez-Espinosa 2016)

a. Estradiol

For PFOS, there was a -4% difference in the natural log estradiol among boys (per interquartile range of the natural log of PFOS). Among boys, the median estradiol level was 10 pg/mL, with an interquartile range (IQR) of <LOD, 15 where the LOD (estradiol detection limit) was 7 pg/mL. For the sample size calculation of mean difference, the standard deviation (SD) was assumed to be equal for the exposed and unexposed groups and equal to 8.5. (Assuming LOD/2 was the lower limit of the IQR, the range = 15 - 3.5 = 11.5. Assuming a normal distribution for estradiol, dividing 11.5 by 1.35 converts the IQR to a standard deviation, which equaled 8.5)¹. To obtain the mean difference, the median estradiol level (10 pg/mL) was assumed to be the reference level (i.e., the level among unexposed). The natural log of the median equals 2.302. A 4% decrease equals 2.21. Exponentiating 2.21 equals 9.12. The mean difference is then 10 - 9.12 = 0.88.

Assuming a 95% CI and 80% power, the sample size = 1,465/group; for a ratio of 2 (exposed/unexposed), the sample sizes = 2,198 and 1,099.

Assuming a 95% CI and 95% power, the sample size = 2,425/group; for a ratio of 2, the sample sizes = 3,638 and 1,819.

Assuming a 90% CI and 80% power = the sample size = 1,154/group; for a ratio of 2, the sample sizes = 1,732 and 866.

b. Testosterone

For PFOS, there was a -6.6% difference in the natural log testosterone among girls (per interquartile range of the natural log of PFOS). Among girls, the median testosterone level was 15 ng/dL with an IQR of <LOD, 21 and the LOD of 10 ng/dL. For the sample size calculation of mean difference, the standard deviation was assumed to be equal for the exposed and unexposed groups and equal to 11.85. (Assuming LOD/2 was the lower limit of the IQR, the range = 21 - 5 = 16. Assuming a normal distribution, dividing 16 by 1.35 converts the IQR to a standard deviation, which equaled 11.85)¹. To obtain the mean difference, the median testosterone level (15 ng/dL) was assumed to be the reference level (i.e., the level among the unexposed). The natural log of the median equals 2.71. A 6.6% decrease equals 2.53. Exponentiating 2.53 equals 12.55. The mean difference is then 15 - 12.55 = 2.45.

Assuming a 95% CI and 80% power, the sample size = 368/group; for a ratio of 2, the sample sizes = 552 and 276.

Assuming a 95% CI and 95% power, the sample size = 608/group; for a ratio of 2, the sample sizes = 912 and 456.

c. IGF-1

For PFHxS, there was a -2.5% difference in the natural log IGF-1 among boys (per interquartile range of the natural log of PFHxS). Among boys, the median IGF-1 level was 147 ng/mL with an IQR of 116, 187. For the sample size calculation of mean difference, the standard deviation was assumed to be equal for the exposed and unexposed groups and equal to 52.6. (The IQR range was 187 - 116 = 71. Assuming a normal distribution, dividing 71 by 1.35 converts the IQR to a standard deviation, which equaled 52.6). To obtain the mean difference, the median IGF-1 level (147 ng/mL) was assumed to be the reference level (i.e., the level among the unexposed). The natural log of the median equals 4.99. A 2.5% decrease equals 4.865. Exponentiating 4.865 equals 129.7. The mean difference is then 147 - 129.7 = 17.3.

Assuming a 95% CI and 80% power, the sample size = 146/group; for a ratio of 2, the sample sizes = 218 and 109.

Assuming a 95% CI and 95% power, the sample size = 241/group; for a ratio of 2, the sample sizes = 362 and 181.

For PFOS, there was a -5.9% difference in the natural log IGF-1 among boys (per interquartile range of the natural log of PFOS). This would require considerably smaller sample sizes for IGF-1 than those for PFHxS.

Thyroid function – Children/Adolescents

- 1. C8 study of children (Lopez-Espinosa 2012)
 - a. Thyroid –stimulating hormone (TSH)

Fourth quartile PFOS was associated with a 3.1% change in the natural log TSH. The median TSH level was 1.83 μ IU/mL and the IQR was 1.31, 2.55. For the sample size calculation of mean difference, the standard deviation was assumed to be equal for the exposed and unexposed groups and equal to 0.92. (The IQR range was 2.55 – 1.31 = 1.24. Assuming a normal distribution, dividing 1.24 by 1.35 converts the IQR to a standard deviation, which equaled 0.92.) To obtain the mean difference, the median TSH level (1.83 μ IU/mL) was assumed to be the reference level (i.e., the level among the unexposed). The natural log of the median equals 0.604. A 3.1% increase equals 0.623. Exponentiating 0.623 equals 1.865. The mean difference is 1.865 – 1.83 = 0.035.

Assuming a 95% CI and 80% power, the sample size=10,846/group.

b. Total thyroxine (TT₄)

Fourth quartile PFOS was associated with a 2.3% change in TT₄. The median TT₄ level was 7.4 µg/dL and the IQR was 6.5, 8.4. For the sample size calculation of mean difference, the standard deviation was assumed to be equal for the exposed and unexposed groups and equal to 1.41. (The IQR range was 8.4 –

6.5 = 1.9. Assuming a normal distribution, dividing 1.9 by 1.35 converts the IQR to a standard deviation, which equaled 1.41.)¹ To obtain the mean difference, the median TT₄ level (7.4 μ g/dL) was assumed to be the reference level (i.e., the level among the unexposed). An increase of 2.3% in TT₄ produces a mean difference of 0.17.

Assuming a 95% CI and 80% power, the sample size = 1,080/group; for a ratio of 2, the sample sizes = 1,620 and 810.

Assuming a 95% CI and 95% power, the sample size = 1,788/group; for a ratio of 2, the sample sizes = 2,682 and 1,341.

2. Taiwan study of children. (Lin 2013)

a. For males aged 12-19, there was a mean difference in the log TSH of -.50 mIU/L for PFOA levels in the 90th percentile (>9.71 ng/ml) compared to the reference level of PFOA exposure. The standard error for the reference group was 0.26 with N=32 in this group; and the standard error for the 90th percentile group was 0.33 with N=6. The standard deviations for the reference and 90th percentile groups were therefore 1.47 and 0.81, respectively.

Assuming a 95% CI and 80% power, the sample size = 89/group; for a ratio of 2, the sample sizes = 158 and 79.

Assuming a 95% CI and 95% power, the sample size = 147/group; for a ratio of 2, the sample sizes = 260 and 130.

b. For females aged 12-19, there was a mean difference in the log TSH of -.35 mIU/L for PFOA levels in the 90th percentile (>9.71 ng/ml) compared to the reference level of PFOA exposure. The standard error for the reference group was 0.18 with N=71 and the standard error for the 90th percentile group was 0.24 with N=14. The standard deviations for the reference and 90th percentile groups were therefore 1.52 and 0.90, respectively.

Assuming a 95% CI and 80% power, the sample size = 200/group; for a ratio of 2, the sample sizes = 348 and 174.

Assuming a 95% CI and 95% power, the sample size = 331/group; for a ratio of 2, the sample sizes = 578 and 289.

Liver Function – Adults

In the C8 study (Darrow 2016), the mean alanine aminotransferase (ALT) level was 26 IU/L and the standard deviation was 19. The linear regression coefficient for the natural log ALT in the fifth quintile level of cumulative natural log PFOA was 0.058. Assuming that the reference group had an ALT level

equal to the mean, the natural log of the mean ALT would be 3.26. Therefore the natural log of ALT for the fifth quintile cumulative log PFOA would be 3.32. Exponentiating 3.32 equals 27.6. The mean difference in the untransformed ALT is then 1.6.

Assuming a 95% CI and 80% power, the sample size = 2,214/group.

Assuming a 95% CI and 95% power, the sample size = 3,665/group.

In the C8 study (Gallo 2012), the linear regression on the natural log of ALT resulted in a regression coefficient for the natural log PFOS of 0.029. The top quintile of PFOS level in the Pease adult population was about 15 $\,$ ng/mL. The natural log of 15 is 2.71; multiplying by 0.029 results in a natural log ALT increase of 0.08. From the graph in the article, the reference level of ALT is about 21.3 $\,$ IU/L. The natural log of 21.3 is 3.06. Adding 0.08 to 3.06 equals 3.14, and exponentiating 3.14 equals 23.1. Therefore the mean difference is 23.1 - 21.3 which equals 1.8.

The ALT standard deviation for the entire population was 20.1, and it was assumed that this was the standard deviation for each quintile PFOS.

Assuming a 95% CI and 80% power, the sample size = 1,958/group.

Assuming a 95% CI and 95% power, the sample size = 3,241/group.

Thyroid Function – Adults

In a study done by Shrestha 2015, the sample size was 87 adults aged 55-74. Mean and SD for TSH was 2.58 μ IU/mL and 1.47, respectively. The linear regression of the natural log TSH resulted in a coefficient for the natural log PFOS of 0.129. Using a PFOS level of 15 ng/mL, the natural log of 15 is 2.71; multiplied by 0.129 equals 0.35. The reference level TSH was assumed to be the median TSH of 2.15 μ IU/mL. The natural log of 2.15 is 0.77; adding 0.35 equals 1.12. Exponentiating 1.12 equals 3.06. The mean difference is then 3.06 – 2.15 = 0.91. The standard deviation of 1.47 was used for each group.

Assuming a 95% CI and 80% power, the sample size = 41/group.

Assuming a 95% CI and 95% power, the sample size = 68/group.

a. TSH

In Ji 2012, the sample size was 633, ≥ 12 years of age and the median TSH level was $1.37 \,\mu$ IU/mL with an IQR of 0.90, 2.01. The standard deviation was estimated as the IQR range divided by 1.35: (2.01 - .90)/1.35 = 0.82. This standard deviation was assumed for each group. For TSH, the linear regression coefficients for PFOS and PFHxS were 0.062 and 0.013, respectively. Using a PFOS level of 15 ng/mL and a PFHxS level of 9 ng/mL, the mean difference for PFOS and PFHxS are 0.93 and 0.12, respectively.

Assuming a 95% CI and 80% power, the sample size = 13/group for PFOS

Assuming a 95% CI and 95% power, the sample size = 21/group for PFOS

Assuming a 95% CI and 80% power, the sample size = 733/group for PFHxS

Assuming a 95% CI and 95% power, the sample size = 1,214/group for PFHxS

b. TT₄ (total thyroxine)

In Ji 2012, the sample size was 633, ≥ 12 years of age and the median TT_4 level was $7.4 \mu g/dL$ and the IQR was 6.7, 8.1. The standard deviation was estimated: (8.1 - 6.7)/1.35 = 1.04. This standard deviation was assumed for each group. For TT_4 , the linear regression coefficients for PFOS and PFHxS were - 0.021 and -0.007, respectively. Using a PFOS level of 15 ng/mL and a PFHxS level of 9 ng/mL, the mean difference for PFOS and PFHxS are -0.32 and -0.06, respectively.

Assuming a 95% CI and 80% power, the sample size = 166/group for PFOS

Assuming a 95% CI and 95% power, the sample size = 275/group for PFOS

Assuming a 95% CI and 80% power, the sample size = 4,716/group for PFHxS

Assuming a 95% CI and 95% power, the sample size = 7,809/group for PFHxS

Footnotes

1. The use of the formula, IQR range divided by 1.35, to obtain the standard deviation assumes a normal distribution. However, given the outcome has been log-transformed presumably to achieve a normal distribution, the untransformed outcome is unlikely to have a normal distribution. Therefore, using this formula when the outcome does not have a normal distribution may underestimate the SD by as much as 20% according to simulations conducted in Wan X. 2014. A higher SD would increase the sample size requirement.

Other sites with PFAS-contaminated drinking water from the UCMR-3

Table A1 shows the maximum combined levels of PFHxS and PFOS in any sample taken from each utility. Only utilities with detectable levels of either PFHxS or PFOS are listed. The data are from the UCMR-3 database as of July 2016 (US EPA 2016b). The ten utilities with the highest PFOS/PFHxS levels in a sample are the Commonwealth Utilities Corporation serving the Mariana Islands, the Artesian Water Company serving portions of the state of Delaware, the Security Water and Sanitation Districts serving Colorado Springs, the Horsham Water & Sewer (PA), the Warminster Municipal Authority (PA), the Oatman Water Company (AZ), the Issaquah Water System (WA), the Hyannis Water System (MA), the Suffolk County Water Authority (NY) and the Warrington Township Water & Sewer (PA). Three of the top 10 utilities are located near each other in the vicinity of Philadelphia, PA: Horsham, Warminster, and Warrington. ATSDR is currently considering whether it is feasible to include children and adults from these towns in studies that would also evaluate the Pease populations.

Willow Grove Naval Air Station/Air Reserve Station (a.k.a. Naval Air Station Joint Reserve Base and Air Force Reserve Station), Montgomery County, Pennsylvania

The Naval Air Station Joint Reserve Base (NASJRB) and Air Reserve Station (ARS) at Willow Grove ("Willow Grove") are two separate, but co-located military facilities in Montgomery County, Pennsylvania. The Navy acquired site in 1942 and began jet training there in 1949; the air force base began operations in 1958. In 2001, the Willow Grove bases employed 1,571 active-duty individuals, 993 members of the National Guard, 3,500 members of the Reserves, and 778 civilians with approximately 1,700 staff on-station daily. About 230 people resided on the bases year-round: less than 30 people resided in single family dwellings and less than 200 resided in barracks. Additionally, there were five officer family units, 200 enlisted family units, and 250 unaccompanied enlisted units as well as a daycare center on base for 96 children. The Willow Grove Branch Medical Clinic was also located there and provided primary care, medical support, preventive medicine, and occupational health services to 20,000 active duty, reserve, retired personnel, and their family members (ATSDR 2002a). Willow Gove became an Air National Guard Base in September 2011. The surplus land with the runways was turned over to Horsham Township for redevelopment.

AFFF used on the Willow Grove bases resulted in PFAS contamination of two nearby water systems – the Warrington Township Water and Sewer Department (WTWSD) which served the eastern portion of Warrington and the Horsham Water and Sewer Authority (HWSA).

In late October 2014, three of eight wells in the southern portion of the WTWSD were above the EPA Provisional Health Advisory Level (PHAL) for PFOS and were taken out of service. PFOS levels were the following: Well 1 (0.21 μ g/L), Well 2 (1.6 μ g/L), and Well 6 (1.3 μ g/L). Although the wells pump directly into the distribution system, wells 1, 2, and 6 are blended together at a tank and enter the distribution system at one point. These wells constituted about 30% of the WTWSD supply. Well 3, in the northeast area of the eastern section, and well 9, which is centrally located in the eastern section, had very low levels of contamination.

Using currently available water distribution system information, ATSDR determined that for "present-day" conditions, the northern part of the eastern section of the WTWDS system generally received water that did not contain PFOA and PFOS. If any customers in the northern part of the system received water containing PFOA and PFOS, it was at levels below the EPA Lifetime Health Advisory (LTHA). The central part of the eastern section of the system may have received water containing PFOA

and PFOS concentrations above the EPA LTHA. The southeastern part of the eastern section of the system received water containing PFOA and PFOS concentrations up to 10 times the EPA LTHA. More detailed analyses of the water-distribution system need to be conducted to estimate historical PFAS concentrations at specific housing areas. These analyses would involve looking at the water-distribution system operating conditions, historical monthly well pumping records, and customer consumption information in more detail.

The western section of the Warrington system is supplied by water purchased from North Wales Water Authority and is not contaminated with PFAS. However, there is an interconnection between the eastern and western sections of the system which is used when there is a need in the eastern section.

Warrington Township Water and Sewer Department (WTWSD) UCMR 2014-2015 data*

Well	PFOS (μg/L)	PFHxS (µg/L)	PFOA (μg/L)	PFNA (µg/L)
Wells 1, 2, 6	0.67	0.24	0.12	-
Well 3	0.06	0.04	0.02	-
Well 9	0.09	0.06	0.03	-

^{*}Wells 1, 2, 3, and 6 were sample 11/11/2014; Well 9 was sampled 5/11/2015

The HWSA is served by 15 wells as well as interconnections with other nearby water utilities. The water system is separated into two pressure zones, "high" and "low," with the wells in each zone pumping to fill storage tanks. The high zone has two storage tanks supplied by three wells and two interconnections. The low zone has three storage tanks served by 11 wells and an interconnection with Aqua Pennsylvania Southeastern Division. (Note: the Aqua system had $0.009 \,\mu\text{g/L}$ of PFOS and $0.005 \,\mu\text{g/L}$ PFOA during UCMR-3 sampling in 4/16. There are now samples from 7/16 which measured $0.0068 \,\mu\text{g/L}$ for PFOS and $0.0065 \,\mu\text{g/L}$ for PFOA.). June 2014 drinking water sample results indicated that PFAS contamination was solely in the low pressure zone which serves the majority of the service area. Prior to 1996 the system did not have pressure zones which means customers located in the current high pressure zone may have received water from wells in the low pressure zone. Generally, demand is met using water from the storage tanks. There are three elevated tanks, and each tank generally supplies a certain area of the system. Each tank will have different PFAS concentrations depending on which wells are supplying water to them. However, it is possible that a property in close proximity to a well which has a demand at the same time the well is pumping will have a higher percentage of water from the nearby well than other areas.

In June 2014, HWSA wells were tested for PFAS as part of the UCMR-3. Two wells, well #26 and well #40, had levels of PFOS greater than the EPA PHAL of 0.2 μ g/l, and well #26 also exceeded the EPA HAL of 0.4 μ g/l for PFOA. The PFAS contamination levels from the UCMR-3 for the Horsham supply wells are shown in the table below. Both wells #26 and #40 were removed from service in July 2014. According to the 2014 consumer confidence report for the HWSA, the average level of PFOS reported was 0.06 ppb, the average level of PFH_xS was 0.037 ppb, and PFOA was not detected. The two contaminated wells generally supplied about 25% of the water for the system; however, there were times that the two contaminated wells supplied as much as 35% of the water for the system.

In May 2016 subsequent to the EPA announcement of its lifetime health advisory for PFOA/PFOS, wells 10, 17, and 21 were immediately taken out of service. One of these three wells was shut down to comply with the EPA's new LTHA. The other two wells, which tested below the LTHA,

were shut down as a precaution. The other nine wells that now supply public drinking water across the township have tested below the EPA lifetime health advisory levels.

ATSDR used currently available water-distribution system information to determine that for "present-day" conditions, some areas in the southern and southeastern part of the low pressure zone received water containing PFOA and PFOS concentrations up to 9 times the EPA LTHA. The northeastern part of the low pressure zone received water containing PFOA and PFOS concentrations less than the EPA LTHA. More detailed analyses of the system need to be conducted to estimate historical PFAS concentrations at specific housing areas. These analyses would involve looking at the water-distribution system operating conditions, historical monthly well pumping records, and customer consumption information in more detail.

In addition to the five total public wells that HWSA shut down, the Navy and the EPA identified approximately 40 additional private wells in Horsham that are at or above the EPA guidance of 70 parts per trillion (ppt). The Navy is providing bottle water to these private well owners.

Horsham Water and Sewer Authority (HWSA) UCMR 2014 data*

	<i>,</i> ,	,		
Well	PFOS (µg/L)	PFHxS (µg/L)	PFOA (μg/L)	PFNA (µg/L)
Well 10	0.05	0.04	0.03	-
Well 17	0.10	0.05	0.03	-
Well 21	0.14	0.08	-	-
Well 26	0.70	0.39	0.29	-
Well 40	1.00	0.59	0.06	-

^{*}Wells 10 and 17 were sampled 12/9/2014; Wells 21, 26, and 40 were sampled 6/24/2014

Other drinking water contaminants

Supply wells on base contained volatile organic compounds (VOC) and metals. Maximum detected levels in supply wells from sampling conducted in 1979-1984 were 91 ppb for PCE and 300 ppb for TCE. After contamination was detected, the well with the highest levels of contamination was used mainly for fire protection. Additionally, the Navy installed an air stripper to treat groundwater prior to distribution, and monitoring of treated water between 1996 and 1998 found no contaminants above EPA's Maximum Contaminant Levels (MCLs) (ATSDR 2002a). According to the EPA, over 800 employees at the two facilities may have drank or come into contact with treated water from the Navy supply wells (https://cumulis.epa.gov/supercpad/cursites/csitinfo.cfm?id=0303820). VOC contamination in off-site wells has not been attributed to the base, and the local water authorities (HWSA and WTWSD) treat the water for VOCs before distribution (ATSDR 2002a).

Naval Air Warfare Center (a/k/a Naval Air Development Center), Warminster Township, Bucks County, Pennsylvania

The former Naval Air Warfare Center (NAWC) is located in Warminster Township. The base operated from 1944 until its closure in September 1996. In 1994, approximately 1,850 civilians and 1,000 military personnel were stationed or employed on base. At its peak, the base employed 2,800 civilians, 200 military personnel, and up to 300 daily contractors (ATSDR 2002b).

Approximately 800 to 1,000 military personnel and their families stationed at nearby Willow Grove Naval Air Station lived in two on-base housing areas at NAWC while as many as six families may have resided in officer housing. Between 450-550 enlisted personnel and their families lived at the Shenandoah Woods housing complex. Site 5, a former landfill, was located in Shenandoah Woods. Quarters A and B, located within Area C, provided housing for the base's commanding officer and second-in-command (ATSDR 2002b).

Four out of eighteen of the Warminster Municipal Authority (WMA) public water supply wells are in close proximity to the former NAWC site. The WMA provides water to approximately 40,000 people. The water supplied to the customers is from water supply wells in the WMA system and may be purchased from the North Wales Water Authority (NWWA) as well as the Upper Southampton Municipal Authority on an emergency basis. WMA's water supply wells are connected individually to the distribution network and are subsequently blended within the distribution system in tanks and standpipes. Therefore, customers located geographically closest to a given water supply well will likely receive more water from that well than users located further away (ATSDR 2016).

AFFF was used for decades at the base for firefighting training activities. PFAS were first tested for in groundwater as emerging contaminants in preparation for the CERCLA 2012 Five Year Review for this site. In summer 2013, PFOS levels above the EPA PHAL were first discovered in groundwater on the former Navy property. As part of the EPA's UCMR-3, sampling for six PFAS in the WMA first occurred in November 2013. UCMR-3 monitoring for PFAS is required at the entry point to the distribution system for each well and at any interconnection in operation. Accordingly, WMA conducted sampling in November 2013 and May 2014 for all wells and conducted sampling in November 2013 and February, May, and August 2014 for the interconnection with NWWA (ATSDR 2016).

Samples taken in the WMA system detected levels of PFOS, PFOA, PFHxS and/or PFHpA. The source of the contamination was the use of AFFF at NAWC. In November 2013, three WMA public water wells had levels at or above EPA's PHAL for PFOS. In this sampling event, 17 samples covering 17 wells in the WMA and one sample of the NWWA interconnection were taken and analyzed for PFAS. One of the 17 WMA samples represents the combined water extracted from WMA Wells 43 and 44. Water from these two wells is combined for treatment and samples are taken after treatment at the entry point to the distribution system. PFOS was detected in 6 public wells and PFOA was detected in 8 public wells. PFOS was detected in Well 26 at 0.791 µg/L, more than three times the 0.2 µg/L PFOS PHAL value. Wells 10 and 13 had PFOS concentrations of 0.193 and 0.16 µg/L that can be rounded to 0.2 µg/L. None of the PFOA detections exceeded the PFOA PHAL in the WMA wells. Well 26 had the highest detections for PFOA and PFOS. In summer 2014, PFOS was detected in four public wells. The highest concentrations were in Well 26 at 1.09 µg/L, more than five times the 0.2 µg/L PFOS PHAL value, and in Well 10 at 0.176 μg/L. PFOA was detected in four wells, including Well 26 at 0.349 μg/L, close to the 0.4 µg/L PHAL for PFOA. Wells 13 and 26 were shut down in June 2014. Well 10 was shut down in September 2014. On May 19, 2016, wells 2, 14 and 15 were removed from service due to the EPA new lifetime health advisory level for PFOA/FPOS (ATSDR 2016).

PFOS levels above the PHAL were also detected in private drinking water samples. As of September 2015, 100 private wells (94 residential and 6 non-residential) were identified and sampled within an approximate 1-3 mile radius of the site. At least one PFAS was detected in the majority (93 out of 100) of these private water wells. Of the 94 residential private water wells, five were non-detect for PFOA and PFOS, 18 had detections of PFOA only, and 71 had both PFOA and PFOS. Eleven exceeded the PFOS PHAL, ranging from $0.152 \,\mu g/L$ to $0.729 \,\mu g/L$. The PFOS PHAL exceedances are

in two general locations: one location is south of the Jacksonville Road and East Bristol Road intersection and the other location is in the area of York Road and W Street. Six residential wells with PFOS levels that range from 0.102 to $0.109 \,\mu g/L$ (50% of the PHAL) are located at the Jacksonville/East Bristol Roads intersection (ATSDR 2016).

The Navy and EPA provided a limited number of residents whose private well water was at or above EPA's PHAL (with rounding up to one significant digit) with bottled water to use for drinking and cooking water, and is currently working to connect these locations to public water (ATSDR 2016).

Using currently available water-distribution system information, ATSDR determined that for "present day" conditions, the southwestern part of the Warminster system typically received water that did not contain PFOA and PFOS concentrations. If any customers in this part of the system received water containing PFOA and PFOS concentrations, it was at levels below the EPA LTHA. The northwestern part of the Warminster system typically received water containing PFOA and PFOS concentrations at or below the EPA LTHA. Some areas in the eastern parts of the Warminster system received water containing PFOA and PFOS concentrations at levels up to three times the EPA LTHA, and areas in the central part received water containing concentrations at level up to 15 times the EPA LTHA. More detailed analyses of the system need to be conducted to estimate historical PFAS concentrations at specific housing areas. These analyses would involve looking at the water-distribution system operating conditions, historical monthly well pumping records, and customer consumption information in more detail.

Although some WMA customers received the majority of their water from one of the contaminated wells, the majority of water customers likely received water that either did not contain PFAS or had levels less than the PHALs (but levels may be higher than the EPA LTHA for PFOS/PFOA). If one assumes that all the wells supply a similar amount of water to the system (each well typically supplied 5-10% of the water to the system), then the number of customers potentially exposed to elevated PFAS in their drinking water could be approximately 7,000.

Warminster Municipal Authority (WMA) UCMR 2013-2014 data*

Well	PFOS (μg/L)	PFHxS (μg/L)	PFOA (µg/L)	PFNA (µg/L)
Well 2	0.06	0.03	0.03	
Well 10	0.19	0.10	0.09	-
Well 13	0.16	0.09	0.12	-
Well 14	0.06	0.03	0.02	-
Well 15	0.06	0.04	0.02	-
Well 26	1.09	0.39	0.35	-

^{*}Wells 2, 10, 13, 14, and 15 were sampled 11/19/2013; Well 26 was sampled 6/9/2014

Other drinking water contaminants

Samples taken in 1979 showed maximum levels of contamination in on-site supply wells of 36 ppb for PCE and 293 ppb for TCE. These wells were closed in 1979. Contamination levels in samples taken from off-site municipal supply wells found 17 ppb for PCE and 67.8 ppb for TCE; past off-base residents may have been exposed to these VOCs between 1974, when the well first began supplying water, until it was closed in 1979. Sampling of VOCs in off-site private wells detected PCE at 31 ppb; as

a result, affected homes were connected to municipal water supplies or groundwater treatment systems were installed (ATSDR 2002b).

Because the TCE- and PCE-contaminated wells were shut down in 1979, military service personnel and DOD civilian workers who began service/employment at NAWC after 1979 might be eligible for a PFAS study. More information is needed to determine when the water supply may have been contaminated with PFAS.

More detailed analyses will help determine which specific housing areas received water containing PFOA and PFOS from the NASJRB and ARS at Willow Grove and the NAWC in Warminster. To conduct more detailed analyses, including modeling, additional information and specific data pertinent to each water system's operations needs to be obtained from site visits to the water utilities.

Appendix tables

Table A1. Maximum levels (parts per billion) of combined PFHxS and PFOS from the US EPA's Third Unregulated Contaminant Monitoring Rule (UCMR-3)

Water Utility Name	<u>State</u>	<u>Size</u>	PFHxS & PFOS sum
Commonwealth Utilities Corp. (Saipan)	MP	L	8.60
Artesian Water Company	DE	L	2.48
Security WSD	CO	L	1.89
Horsham Water & Sewer Authority	PA	L	1.59
Warminster Municipal Authority	PA	L	1.479
Oatman Water Company	AZ	S	1.03
Warrington Township Water & Sewer Department	PA	L	0.91047
Issaquah Water System	WA	L	0.841
Hyannis Water System	MA	L	0.7
Suffolk County Water Authority	NY	L	0.67
United Water PA	PA	L	0.572
Emerald Coast Utilities Authority	FL	L	0.56
GU Waterworks Authority - Northern System	GU	L	0.55
Widefield WSD	CO	L	0.54
Oakdale	MN	L	0.4913
City of Tucson	AZ	L	0.476
City of Cleveland Heights	ОН	L	0.4
Sanford Water District	ME	L	0.4
Wright-Patterson AFB Area A/C	ОН	L	0.36
Liberty Water LPSCO	AZ	L	0.33
Westfield Water Department	MA	L	0.33
City of Zephyrhills	FL	L	0.32
Bemidji	MN	L	0.32
City of Fountain	CO	L	0.29
City of Stuart Water Plant	FL	L	0.259

Water Utility Name	<u>State</u>	<u>Size</u>	PFHxS & PFOS sum
City of Tempe	AZ	L	0.245
CA American Water Co Suburban	CA	L	0.241
City of Newburgh	NY	L	0.24
CA Water Service - Visalia	CA	L	0.212
Eastern Municipal Water District	CA	L	0.202
New Windsor Consolidated Water District	NY	L	0.1936
VAW Water System, Inc.	AL	L	0.18
Freeport	IL	L	0.18
La Crosse Waterworks	WI	L	0.172
Salt River Public Works	09 [*]	L	0.166
City of Martinsburg	WV	L	0.157
Dyer Water Department	IN	L	0.1437
Atlantic City MUA	NJ	L	0.142
West Morgan - East Lawrence Water Authority	AL	L	0.13
City of Greensboro	NC	L	0.124
Rome	GA	L	0.12
Dover Water Department	NH	L	0.12
CA Water Service - Chico	CA	L	0.118
Moore County Public Utilities - Pinehurst	NC	L	0.118
Rhinelander Water & Wastewater	WI	S	0.1173
Bayleaf Master	NC	L	0.11
City of Ocala	FL	L	0.104
NJ American Water Co Raritan	NJ	L	0.103
Mahwah Water Department	NJ	L	0.098
City of Abilene	TX	L	0.09781
West Lawrence Water Co-op	AL	L	0.09
Hampton Bays Water District	NY	L	0.082
Fort Drum	NY	L	0.08

Water Utility Name	<u>State</u>	<u>Size</u>	PFHxS & PFOS sum
City of Lathrop	CA	L	0.076
Northeast Alabama Water System	AL	L	0.07
City of Anaheim	CA	L	0.07
Fair Lawn Water Department	NJ	L	0.06603
City of Orange	CA	L	0.0659
Montebello Land & Water Company	CA	L	0.065
Vienna	WV	L	0.0641
Chatsworth	GA	L	0.06303
Bethany	OK	L	0.063
City of Pico Rivera Water Department	CA	L	0.062
Camp Pendleton (South)	CA	L	0.062
Montgomery County Water Services #2	ОН	L	0.061
Rainbow City Utilities Board	AL	L	0.06
Florence Water-Wastewater Department	AL	L	0.06
Plainfield Township	MI	L	0.06
Pendleton County Water District #1/South	KY	S	0.05853
City of Miami Beach	FL	L	0.058
Ridgewood Water	NJ	L	0.058
Woodbury	MN	L	0.0577
Montgomery County Water Services #1	ОН	L	0.0542
CA Water Service - East Los Angeles	CA	L	0.054
Town of Nashville	NC	S	0.05312
Metropolitan DWID	AZ	L	0.053
City of Downey Water Department	CA	L	0.053
Pierre	SD	L	0.053
Park Water Company - Bellflower/Norwalk	CA	L	0.051
Washington Township MUA	NJ	L	0.0503

Water Utility Name	<u>State</u>	<u>Size</u>	PFHxS & PFOS sum
Colbert County Rural Water System	AL	L	0.05
Gadsden Waterworks & Sewer Board	AL	L	0.05
Southside Waterworks	AL	L	0.05
City of North Miami	FL	L	0.05
Kennebunk, Kennebunkport & Wells WD	ME	L	0.05
Bell Arthur Water Corp.	NC	S	0.05
City of Garden Grove	CA	L	0.0496
City of Lauderhill	FL	L	0.049
FKAA	FL	L	0.049
Yorba Linda Water District	CA	L	0.0474
City of Miramar	FL	L	0.047
Miami International Airport	FL	L	0.047
City of Corona	CA	L	0.046
Orchard Dale Water District	CA	L	0.045
Lima City Water	ОН	L	0.045
Pico Water District	CA	L	0.044
Golden State Water Co Norwalk	CA	L	0.043
MDWASA - Main System	FL	L	0.043
Ann Arbor	MI	L	0.043
City of Fullerton	CA	L	0.0412
Cliffdale West	NC	L	0.041
Central ASG	AS	L	0.04
City of DeFuniak Springs Water System	FL	L	0.04
Cottage Grove	MN	L	0.0381
City of Great Bend	KS	L	0.037
City of Pleasanton	CA	L	0.036
Sacramento Suburban Water District	CA	L	0.036

Water Utility Name	<u>State</u>	<u>Size</u>	PFHxS & PFOS sum
Mashpee Water District	MA	L	0.033
Belvidere	IL	L	0.03167

L=large system (serves >10,000); S=small system (serves <10,000)

^{*} Tribal nation located in Arizona

Table A2. PFAS studies on cancers and other chronic diseases in adults.

Reference	Exposure*	Outcome	RR (SIR, SMR, OR, HR) &	Duration/Intensity/cumulative exp.
			95% CI	
Alexander 2003	POSF/PFOS	Bladder Cancer (and other	3 cases	
Decatur, AL	High exposure job	urinary organs) mortality	SMR=12.77 (2.63, 37.35)	
POSF plant	≥1 yr		SMR=16.12 (3.32, 47.14)	
Geometric mean		Urinary Cancers mortality		
serum level:	High exposure job		SMR=4.02 (0.83, 11.75) (3 cases)	
PFOS=0.9 ppm	≥1 yr		SMR=5.11 (1.05, 14.93)	
PFOA=1.13 ppm				
Olsen 2004	POSF/PFOS	Colon Cancer	RR=5.4 (0.5, >100) 4 exposed	Long term: RR=12 (0.8, >100)
Decatur, AL		Rectal Cancer	RR=1.8 (0.3, 12.4) 4 exposed	Long term: RR=11 (0.8, >100)
POSF plant		Prostate Cancer	RR=7.7 (0.9, >100) 5 exposed	Long term: RR=8.2 (0.8, >100)
Grice 2007	POSF/PFOS		Odds Ratios:	Low/high L/H (>1 yr) High (>1 yr)
Decatur, AL		Colon Cancer		1.2 (0.5, 2.9) 1.4 (0.6, 3.3) 1.7 (0.7, 4.2)
POSF plant		Prostate Cancer		1.3 (0.6, 2.9) 1.4 (0.6, 3.0) 1.1 (0.4, 2.7)
Alexander 2007	POSF/PFOS	Bladder Cancer (11 cases)		Cumulative exposure:
Decatur, AL	Ever low		SIR=2.26 (0.91, 4.67)	Low RR=0.83 (0.15, 4.65)
POSF plant	Ever low or high		SIR=1.70 (0.77, 3.22)	Medium RR=1.92 (0.30, 12.06)
_	Low or high (≥1 yr)		SIR=1.31 (0.48, 2.85)	High RR=1.52 (0.21, 10.99)
	Ever high		SIR=1.74 (0.64, 3.79)	
	High (≥1 yr)		SIR=1.12 (0.23, 3.27)	
Eriksen 2009	PFOS	Prostate Cancer	Trend: RR=1.05 (0.97, 1.14)/10 ppb	RRs PFOS PFOA
Danish Pop.	PFOA		Trend: RR=1.03 (0.99, 1.07)/1 ppb	Low: 1.35 (0.97, 1.87) 1.09 (0.78, 1.53)
Median serum		(PFOS & PFOA: no association	(plasma concentration)	Med: 1.31 (0.94, 1.82) 0.94 (0.67, 1.32)
level:		for cancers of bladder or liver)		High: 1.38 (0.99, 1.93) 1.18 (0.84, 1.65)
PFOS=35 ng/ml				
PFOA=6.9 ng/ml		Pancreatic Cancer		Low: 0.88 (0.49, 1.57)
	PFOS		No association with PFOS	Med: 1.33 (0.74, 2.38)
	PFOA		Trend: RR=1.03 (0.98, 1.10)/1 ppb	High: 1.55 (0.85, 2.80)
Hardell 2014	PFAS (the median	Prostate Cancer		ORs: PSA ≥11 Heredity, PFC>median
Swedish Pop.	level of each PFAS	(case-control)	PFH _x S: OR=1.3 (0.8, 1.9)	1.5 (0.9, 2.6) 4.4 (1.7, 12)
Median serum	among the controls		PFOS: OR=1.0 (0.6, 1.5)	0.8 (0.4, 1.3) 2.7 (1.04, 6.8)
level: (ng/ml)	was used as the cut		PFOA: OR=1.1 (0.7, 1.7)	1.3 (0.8, 2.1) 2.6 (1.2, 6.0)
PFOS=9.0	point for the		PFNA: OR=1.2 (0.8, 1.8)	1.2 (0.7, 2.1) 2.1 (0.9, 4.8)
PFOA=2.0	calculation of the		PFDA: OR=1.4 (0.9, 2.1)	1.2 (0.7, 2.0) 2.6 (1.1, 6.1)
PFHxS=0.91	OR.)		PFU _n DA: OR=1.2 (0.8, 1.9)	1.0 (0.6, 1.6) 2.6 (1.1, 5.9)

Reference	Exposure*	Outcome	RR (SIR, SMR, OR, HR) & 95% CI	Duration/Intensity/cumulative exp.
Raleigh 2014	PFOA	Cancer:	Cumulative exposure, Mortality	Cumulative exposure, Incidence
APFO production		Prostate	Q1: HR=0.34 (0.25, 1.60)	Q1: HR=0.80 (0.57, 1.11)
workers, Cottage			Q2: HR=1.12 (0.53, 2.37)	Q2: HR=0.85 (0.61, 1.19)
Grove, 3M			Q3: HR=0.36 (0.11, 1.17)	Q3: HR=0.89 (0.66, 1.21)
- · · · , ·			Q4: HR=1.32 (0.61, 2.84)	Q4: HR=1.11 (0.82, 1.49)
Geometric mean				
serum level:		Kidney	Q1-Q2: HR=0.38 (0.11, 1.23)	Q1-Q2: HR=1.07 (0.46, 2.46)
PFOA=815 ng/ml			Q3-Q4: HR=0.39 (0.11, 1.32)	Q3-Q4: HR=0.85 (0.36, 2.06)
l		_		
PFOA-related		Pancreas	Q1: HR=0.32 (0.08, 1.35)	Q1-Q2: HR=0.13 (0.02, 1.03) (1 case)
manufacturing,			Q2: HR=0.89 (0.34, 2.31)	Q3-Q4: HR=1.36 (0.59, 3.11)
PFOA=2,538			Q3: HR=0.82 (0.32, 2.12)	
ng/ml			Q4: HR=1.23 (0.50, 3.00)	
		Bladder	Q1-Q2: HR=1.03 (0.27, 3.96)	Q1: HR=0.81 (0.36, 1.81)
		Bradder	Q3-Q4: HR=1.96 (0.63, 6.15)	Q2: HR=0.78 (0.33, 1.85)
			Q3-Q4. 11K=1.90 (0.03, 0.13)	Q3: HR=1.50 (0.80, 2.81)
				Q4: HR=1.66 (0.86, 3.18)
				Q4. 11K=1.00 (0.00, 3.18)
		Breast	Q1-Q2: HR=0.61 (0.25, 1.48)	Q1-Q2: HR=0.46 (0.25, 0.87)
			Q3-Q4: HR=0.54 (0.15, 1.94)	Q3-Q4: HR=1.27 (0.70, 2.31)
		Chronic Diseases:		
		Ischemic Heart Disease	Q1: HR=0.93 (0.73, 1.18)	
			Q2: HR=0.87 (0.66, 1.13)	
			Q3: HR=0.88 (0.68, 1.13)	
			Q4: HR=0.89 (0.66, 1.21)	
l				
		Cerebrovascular Disease	Q1: HR=0.57 (0.32, 1.02)	
			Q2: HR=0.70 (0.39, 1.24)	
			Q3: HR=0.93 (0.57, 1.53)	
			Q4: HR=0.98 (0.53, 1.81)	
		Diabetes	Q1: HR=0.27 (0.10, 0.76)	
		Diadetes	Q2: HR=0.42 (0.17, 1.04)	
			Q3: HR=0.80 (0.42, 1.51)	
			Q4: HR=0.72 (0.34, 1.52)	
			113	

Reference	Exposure*	Outcome	RR (SMR, HR) & 95% CI	Duration/Intensity/cumulative exp.		
Lundin 2009	PFOA	Mortality	Job classification	Cumulative Exposure-Years		
APFO production						
workers, Cottage		Prostate Cancer	Low (ref.)	<1 (ref.)		
Grove, 3M [€]			Med HR=3.0 (0.9, 9.7)	≥1 HR=2.0 (0.7, 5.3)		
			High HR=6.6 (1.1, 37.7)			
		Pancreatic Cancer	Low (ref.)	<1 (ref.)		
		Fancieatic Cancer	Med/hi HR=1.6 (0.5,4.8)	≥1 HR=1.8 (0.6, 5.6)		
			Wed/III TIK=1.0 (0.3,4.8)	≥ 1 11K-1.8 (0.0, 5.0)		
		Bladder Cancer	Low (ref.)	<1 (ref.)		
			Med/hi HR=0.7 (0.2, 3.4)	≥1 HR=1.7 (0.4, 7.8)		
		Cerebrovascular Disease	Low (ref.)	<1 (ref.)		
			Med HR=1.8 (0.9, 3.7)	1-4.9 HR=0.6 (0.2, 2.2)		
			High HR=4.6 (1.3, 17.0)	≥5 HR=2.1 (1.0, 4.6)		
		Diabetes	Low (ref.)	<1 (ref.)		
		Diaoctes	Med/hi HR=3.4 (1.3, 9.3)	≥1 HR=1.3 (0.6, 3.1)		
Steenland and	PFOA	Mortality	111cd/iii 111c-3.1 (1.3, 7.3)	cumulative exposure-years (SMR¥≠)		
Woskie 2012				Q1 Q2 Q3 Q4		
Dupont plant,		Liver Cancer	SMR=1.07 (0.51, 1.96)	2.39 0 2.01 0.32		
Parkersburg WV		Pancreatic Cancer	SMR=1.04 (0.62, 1.64)	1.18 1.02 1.09 0.92		
(Washington		Lung Cancer	SMR=0.78 (0.62, 1.64)	0.58 0.63 1.09 0.75		
Works Plant)		Breast Cancer	SMR=0.65 (0.13, 1.90)	1.49 0 0.87 0		
		Prostate Cancer	SMR=0.76 (0.47, 1.16)	1.07 0.82 0.65 0.57		
Median serum		Kidney Cancer	SMR=1.28 (0.66, 2.24)	1.07 1.37 0 2.66		
level: (1979-2004)		Bladder Cancer	SMR=1.08 (0.52, 1.99)	1.24 2.49 0.39 0.36		
PFOA=580 ng/ml		Mesothelioma	SMR=2.85 (1.05, 6.20)	0 0 1.73 6.27		
		NHL	SMR=1.05 (0.57, 1.76)	1.54 0.99 0.85 0.96		
Nonexposed		Leukemias	SMR=1.05 (0.57, 1.76)	0.28 2.34 0.57 1.03		
workers:		Diabetes	SMR=1.90 (1.35, 2.61)	1.85 1.47 2.30 1.90		
PFOA=160 ng/ml		Ischemic Heart Disease	SMR=0.97 (0.86, 1.09)	1.07 1.02 0.87 0.93		
Directly exposed		Stroke	SMR=0.86 (0.64, 1.14)	0.63 0.78 1.34 0.69		
workers:		COPD	SMR=1.05 (0.75, 1.42)	0.93 1.00 1.30 0.93		
PFOA=2,880		Chronic Liver Disease	SMR=1.09 (0.54, 1.95)	1.32 2.10 0.37 0.72		
ng/ml		Chronic Renal Disease	SMR=3.11 (1.66, 5.32)	0 3.79 1.83 8.60		

Reference	Exposure*	Outcome	cumulative exposure-years (RR & 95% CI), 10 year exposure lag			
			Q1 (ref.) Q2		Q3	Q4
Steenland 2015	PFOA	Ulcerative colitis	3.00 (0.82,	, 11.0)	3.26 (0.70, 15.1)	6.57 (1.47, 29.4)
Dupont plant,		Rheumatoid arthritis	1.74 (0.45,	, 6.77)	2.12 (0.40, 11.1)	2.62 (0.47, 14.7)
Parkersburg WV		Bladder cancer	0.55 (0.12,	, 2.61)	0.47 (0.10, 2.21)	0.31 (0.06, 1.54)
		Colorectal cancer	0.31 (0.90)	, 1.11)	0.99 (0.33, 2.94)	1.06 (0.34, 3.31)
Median serum		Prostate cancer	1.92 (0.56,	, 6.58)	1.89 (0.57, 6.34)	2.15 (0.64, 7.26)
level: (2005/2006)		Melanoma	0.85 (0.27,	, 2.71)	1.10 (0.34, 3.58)	0.75 (0.21, 2.67)
PFOA=113 ng/ml		Liver disease (non-hepatitis)	1.46 (0.42,	, 5.04)	2.13 (0.59, 7.71)	2.02 (0.50, 8.10)
		Thyroid disease (males)	1.23 (0.57,	, 2.66)	1.70 (0.74, 3.91)	1.71 (0.68, 4.25)
		Thyroid disease (females)	0.79 (0.42,	, 1.50)	0.87 (0.37, 2.02)	0.23 (0.05, 1.01)
		Coronary heart disease	1.20 (0.82)	, 1.75)	1.06 (0.71, 1.58)	0.93 (0.61, 1.41)
		Hypertension	0.95 (0.81,	, 0.98)	0.91 (0.75, 1.16)	0.95 (0.77, 1.16)
		High cholesterol	0.93 (0.79)	, 1.10)	1.01 (0.84, 1.22)	0.96 (0.78, 1.18)
		Osteoarthritis	0.74 (0.49)	, 1.10)	0.56 (0.34, 0.93)	0.67 (0.39, 1.14)
		Stroke	1.48 (0.56,	, 3.69)	1.53 (0.60, 3.89)	1.33 (0.51, 3.43)
		COPD	0.75 (0.38,	, 1.48)	1.16 (0.60, 2.26)	0.77 (0.38, 1.57)
		Asthma	0.48 (0.10,	, 2.29)	0.57 (0.11, 2.93)	0.52 (0.09, 2.83)
		Kidney disease	1.32 (0.32,	, 5.43)	0.50 (0.11, 2.34)	0.67 (0.15, 3.05)
		Diabetes	1.06 (0.75)	, 1.49)	1.10 (0.76, 1.61)	1.12 (0.76, 1.66)

Reference	Exposure	Outcome	Odds Ratio & 95% CI	Odds Ratios & 95	% CI
Bonefeld-	PFCs	Breast Cancer (case-control)	(per ng/ml, serum)		
Jorgensen 2011	PFOS		OR=1.03 (1.00, 1.07)		
Greenlandic Inuit	PFOA		OR=1.20 (0.77, 1.88)		
Median serum	$sumPFSA^{\mathfrak{t}}$		OR=1.03 (1.00, 1.05)		
level:	$sumPFCA^{\pm}$		OR=1.07 (0.96, 1.18)		
PFOS=45.6 ng/ml					
PFOA=2.5 ng/ml					
Bonefeld-		Breast Cancer	Quartile 1 (ref.)	Age, $Dx \leq 40$	Age, $Dx > 40$
Jorgensen 2014	PFOS, Q2		OR=1.51 (0.81, 2.71)	OR=1.2 (0.5, 2.9)	OR=2.3 (0.9, 5.6)
Case-control	Q3		OR=1.51 (0.82, 2.84)	OR=1.4 (0.6, 3.3)	OR=1.9 (0.7, 5.0)
study,	Q4		OR=1.13 (0.59, 2.04)	OR=0.8 (0.3, 1.9)	OR=2.2 (0.9, 5.7)
premenopausal	Q5		OR=0.90 (0.47, 1.70)	OR=1.0 (0.4, 2.5)	OR=0.9 (0.3, 2.4)
mothers nested in					
the Danish	PFOA, Q2		OR=0.97 (0.53, 1.75)	OR=0.7 (0.3, 1.6)	OR=1.8 (0.7, 4.3)
National Birth	Q3		OR=1.02 (0.56, 1.89)	OR=1.3 (0.5, 3.2)	OR=0.9 (0.4, 2.3)
Cohort	Q4		OR=1.14 (0.62, 2.12)	OR=0.8 (0.4, 2.0)	OR=1.9 (0.8, 4.8)
	Q5		OR=0.94 (0.51, 1.76)	OR=0.8 (0.3, 1.9)	OR=1.2 (0.5, 2.9)
Bonefeld-					
Jorgensen 2014		Breast Cancer	Quartile 1 (ref.)	Age, $Dx \leq 40$	Age, $Dx > 40$
(cont.)	PFNA, Q2		OR=1.10 (0.60, 2.02)	OR=1.1 (0.4, 2.5)	OR=1.2 (0.5, 3.0)
	Q3		OR=0.75 (0.41, 1.40)	OR=0.5 (0.2, 1.3)	OR=1.3 (0.5, 3.3)
Mean serum	Q4		OR=1.08 (0.58, 1.99)	OR=0.8 (0.4, 1.9)	OR=1.9 (0.7, 4.9)
level:	Q5		OR=0.80 (0.43, 1.47)	OR=0.6 (0.2, 1.4)	OR=1.1 (0.5, 2.8)
PFOS=30.6 ng/ml					
PFOA=5.2 ng/ml	PFHxS, Q2		OR=0.64 (0.34, 1.18)	OR=0.4 (0.2, 0.9)	OR=1.2 (0.4, 3.4)
PFHxS=1.2 ng/ml	Q3		OR=0.70 (0.38, 1.29)	OR=0.6 (0.2, 1.4)	OR=1.0 (0.4, 2.7)
PFNA=0.5 ng/ml	Q4		OR=0.38 (0.20, 0.70)	OR=0.3 (0.1, 0.7)	OR=0.5 (0.2, 1.4)
PFOSA=3.5 ng/ml	Q5		OR=0.61 (0.33, 1.12)	OR=0.4 (0.2, 1.0)	OR=1.0 (0.4, 2.5)
	PFOSA, Q2		OR=1.38 (0.75, 2.52)	OR=1.5 (0.7, 3.3)	OR=1.3 (0.5, 3.6)
	Q3		OR=0.91 (0.49, 1.66)	OR=1.0 (0.5, 2.4)	OR=1.0 (0.4, 2.5)
	Q4		OR=1.11 (0.60, 2.05)	OR=1.1 (0.5, 2.6)	OR=1.4 (0.5, 3.6)
	Q5		OR=1.89 (1.01, 3.54)	OR=2.5 (1.0, 6.0)	OR=1.6 (0.6, 4.3)

Reference	Exposure	Outcome	Hazard Ratio (95% CI) 10 yr. lag	Community	Occupational
Barry et al 2013	$PFOA^\Delta$	Bladder Cancer	HR=0.98 (0.88, 1.10)	0.90 (0.75, 1.09)	0.73 (0.55, 0.98)
C8 Health Project		Brain Cancer	HR=1.06 (0.79, 1.41)	1.02 (0.68, 1.52)	0.73 (0.32, 1.67)
Entire		Breast Cancer	HR=0.93 (0.88, 0.99)	0.95 (0.89, 1.01)	1.03 (0.59, 1.79)
cohort=32,254		Cervical Cancer	HR=0.98 (0.69, 1.38)	1.02 (0.72, 1.43)	one case
		Colorectal Cancer	HR=0.99 (0.92, 1.07)	0.98 (0.89, 1.09)	1.08 (0.84, 1.39)
Community		Esophageal Cancer	HR=0.97 (0.72, 1.31)	1.01 (0.67, 1.52)	1.17 (0.19, 7.36)
Cohort = $28,541$		Leukemia	HR=1.02 (0.88, 1.18)	0.92 (0.75, 1.13)	1.30 (0.78, 2.18)
		Liver Cancer	HR=0.74 (0.43, 1.26)	0.53 (0.21, 1.34)	one case
Worker Cohort =		Lung Cancer	HR=0.92 (0.81, 1.04)	0.89 (0.76, 1.05)	1.04 (0.68, 1.58)
3,713		Lymphoma	HR=0.98 (0.88, 1.10)	1.02 (0.89, 1.17)	1.10 (0.73, 1.65)
		Melanoma	HR=1.04 (0.96, 1.13)	1.02 (0.92, 1.14)	0.93 (0.73, 1.18)
Median serum		Oral Cancer	HR=0.66 (0.43, 1.02)	0.77 (0.47, 1.27)	one case
level:		Ovarian Cancer	HR=0.90 (0.69, 1.16)	0.94 (0.73, 1.22)	no cases
Community,		Pancreatic Cancer	HR=0.96 (0.75, 1.22)	0.98 (0.72, 1.34)	1.14 (0.33, 3.89)
PFOA=24.2 ng/ml		Prostate Cancer	HR=0.99 (0.94, 1.05)	0.98 (0.90, 1.06)	0.98 (0.83, 1.16)
Worker,		Soft Tissue Cancer	HR=0.72 (0.48, 1.09)	0.64 (0.36, 1.13)	0.91 (0.25, 3.33)
PFOA=112.7		Stomach Cancer	HR=0.77, 0.49, 1.22)	0.74 (0.41, 1.31)	one case
ng/ml		Uterine Cancer	HR=0.99 (0.86, 1.15)	0.99 (0.84, 1.16)	0.96 (0.42, 2.18)
		Kidney Cancer	HR=1.09 (0.97, 1.21)	1.11 (0.96, 1.29)	0.99 (0.67, 1.46)
		Q2	HR=0.99 (0.53, 1.85)	0.94 (0.45, 1.99)	1.22 (0.28, 5.30)
		Q3	HR=1.69 (0.93, 3.07)	1.08 (0.52, 2.25)	3.27 (0.76, 14.1)
		Q4	HR=1.43 (0.76, 2.69)	1.50 (0.72, 3.13)	0.99 (0.21, 4.68)
		Testicular Cancer	HR=1.28 (0.95, 1.73)	1.53 (1.09, 2.15)	1.61 (0.21, 12.2)
		Q2	HR=0.87 (0.15, 4.88)	0.98 (0.13, 7.14)	two cases total
		Q3	HR=1.08 (0.20, 5.90)	1.54 (0.19, 12.2)	in the cohort
		Q4	HR=2.36 (0.41, 13.7)	4.66 (0.52, 41.6)	
		Thyroid Cancer	HR=1.04 (0.89, 1.20)	1.00 (0.84 (1.20)	1.12 (0.61, 2.05)
		Q2	HR=2.06 (0.93, 4.56)	2.09 (0.91, 4.82)	1.65 (0.09, 31.5)
		Q3	HR=2.02 (0.90, 4.52)	1.92 (0.82, 4.50)	4.52 (0.10, 198)
		Q4	HR=1.51 (0.67, 3.39)	1.42 (0.60, 3.37)	5.85 (0.13, 257)

Reference	Exposure	Outcome	Odds Ratio & 95% CI	Odds Ratios & 95% CI		
Vieira 2013	PFOA	Cancers [©] :	OR for the Little Hocking system	OR for Very High serum level category		
C8 Health Project		Breast Cancer	OR=1.2 (0.8, 2.0)	OR=1.4 (0.9, 2.3) (non-monotonic trend)		
		Kidney Cancer	OR=1.7 (0.9, 3.3)	OR=2.0 (1.0, 3.9) (non-monotonic trend)		
Median serum		NHL	OR=1.6 (0.9, 2.8)	OR=1.8 (1.0, 3.4) (non-monotonic trend)		
level:		Ovarian Cancer	OR=1.8 (0.7, 4.4)	OR=2.1 (0.8, 5.5) (non-monotonic trend)		
PFOA=28.2 ng/ml		Prostate Cancer	OR=1.4 (0.9, 2.3)	OR=1.5 (0.9, 2.5) (non-monotonic trend)		
Estimate PFOA		Testicular Cancer	OR=5.1 (1.6, 15.6)	OR=2.8 (0.8, 9.2) (non-monotonic trend)		
median serum						
level in 1995 at						
Little Hocking						
WD = 125 ng/ml						
Reference	Exposure*	Outcome	SMR & 95% CI	Cumulative exposure (SMR)		
Consonni 2013	APFO (PFOA)	Mortality:		No Exp. Low Med High		
TFE synthesis &		Esophageal cancer	1.44 (0.72, 2.57)	0 1.62 1.54 1.16		
polymerization		Liver cancer	1.43 (0.57, 2.94)	0.72 0.70 1.25 2.14		
workers (including		Pancreatic cancer	1.05 (0.51, 1.94)	1.66 0 1.30 1.84		
the WV Dupont		Lung cancer	0.73 (0.54, 0.97)	0.75 0.91 0.75 0.54		
plant workers and		Kidney/other urinary cancers	1.69 (0.81, 3.11)	0 1.57 1.50 2.00		
6 other production		Leukemias	1.61 (0.80, 2.88)	0.79 1.64 1.35 1.85		
sites in NJ and		Stomach cancer	0.52 (0.17, 2.21)			
Europe)		Colon cancer	0.48 (0.19, 0.99)			
		Rectal cancer	1.03 (0.38, 2.25)			
		Laryngeal cancer	0.76 (0.09, 2.75) (2 cases)			
		Prostate cancer	0.24 (0.05, 0.70) (3 cases)			
		Testicular cancer	1.35 (0.03, 7.49) (1 case)			
		Bladder cancer	0.55 (0.11, 1.60) (3 cases)			
		Brain cancer	0.64 (0.17, 1.63)			
		NHL	0.79 (0.26, 1.84)			
				No Exp. Low Med High		
	APFO (PFOA)	Multiple myeloma	0.66 (0.08, 2.39) (2 cases)	THO DAY. DOW MICH INCH		
	1110 (11011)	Diabetes mellitus	0.57 (0.23, 1.17)			
		Circulatory diseases	0.88 (0.77, 1.00)			
		Respiratory diseases	0.63 (0.42, 0.89)			
		Liver cirrhosis	1.00 (0.60, 1.56)	1.19 1.49 0.92 0.52		
		Nephritis, nephrosis	0.92 (0.25, 2.37)	0 0.67 1.49 0.67		

Mortality: Alexander 2003, Grice 2007, Alexander 2007, Lundin 2009, Steenland & Woskie 2012, Consonni 2013, Raleigh 2014

Incidence: Grice 2007, Alexander 2007, Eriksen 2009, Barry 2013, Vieira 2013, Hardell 2014, Raleigh 2014, Steenland 2015

^{*} Exposures are occupational unless otherwise noted.

⁶ The cohort evaluated in the Lundin 2009 study is the same cohort included in the Raleigh 2014 study. The Lundin 2009 study is included in the table because it provides additional information that can be used to interpret the more recent Raleigh 2014 study.

^{*} Reference rates are from other Dupont workers.

[≠] Exposure was not lagged.

[£] sumPFSA: sum of PFOS, PFHxS and PFOSA

[±] sumPFCA: sum of PFHpA, PFOA, PFNA, PFDA, PFUnA, PFDoA and PFTrA

^Δ The hazard ratio is per unit of log estimated cumulative PFOA serum concentration (ng/ml). For cancers of the kidney, testes and thyroid, hazard ratios are also provided for quartiles of cumulative PFOA serum concentration with the first quartile as referent.

[©] Except for kidney cancer, these are cancers with ORs at Little Hocking ≥ the ORs for the other water systems and also with elevated ORs in the very high PFOA serum exposure category. For kidney cancer, the OR for the Tuppers Plains system was 2.0 compared to the OR for Little Hocking of 1.7. Other cancers not listed were not elevated in the Little Hocking system and/or were not elevated in the very high serum category.

Table A3. Other Adult Diseases

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Dhingra 2016a C8	28,240 ≥20 years of age			28.2	Chronic kidney disease 397 cases (retrospective) 212 cases (prospective)	Hazard ratios for quintiles of cumulative exposure Retrospective 2 nd : 1.26 (0.90, 1.75) 3 rd : 1.12 (0.80, 1.55) 4 th : 1.12 (0.81, 1.56) 5 th : 1.24 (0.88, 1.75) Hospital exposure 1.36 (0.89, 2.09) 0.94 (0.62, 1.45) 1.08 (0.70, 1.66) 1.12 (0.72, 1.75)
Dhingra 2016b C8	29,641 ≥20 years of age 6,342 women, aged 30-65, who had not had a hysterectomy			28.2	Estimated glomerular filtration rate (eGFR) (mL/min/1.73 m²); Earlier menopause (reported as of 2005/2006); and Reverse causation	Measured eGFR (β±S.E.) Menopause (OR, 95% CI) 2^{nd} : $-0.64 ± 0.268$ 1.68 (1.21, 2.35) 3^{rd} : $-1.03 ± 0.269$ 1.45 (1.04, 2.02) 4^{th} : $-0.84 ± 0.271$ 1.39 (1.00, 1.93) 5^{th} : $-0.98 ± 0.274$ 1.58 (1.14, 2.19) Modeled serum PFOA quintiles eGFR (β±S.E.) Menopause (OR, 95% CI) 2^{nd} : $-0.08 ± 0.268$ 0.98 (0.70, 1.37) 3^{rd} : $0.37 ± 0.268$ 1.05 (0.75, 1.45) 4^{th} : $0.21 ± 0.269$ 0.78 (0.56, 1.08) 5^{th} : $0.23 ± 0.271$ 0.92 (0.65, 1.30)
Steenland 2010 C8	53,458 aged ≥20 years	20.2		27.9	Uric Acid mg/dL Hyperuricemia	Highest decile serum PFOA: 0.28 ± 0.02 Highest decile serum PFOS: 0.22 ± 0.02 Monotonic exposure-response for PFOA and PFOS Highest quintile serum PFOA: OR=1.47 (1.37, 1.58) Highest quintile serum PFOS: OR=1.26 (1.17, 1.35) Monotonic exposure-response for PFOA and PFOS
Shankar 2011 NHANES, 1999- 2000; 2003-2006	3,883 aged ≥20 years	17.9		3.5	Uric Acid mg/dL hyperuricemia	4 th quartile serum PFOA: 0.44 (0.32, 0.56) 4 th quartile serum PFOS: 0.27 (0.13, 0.41) Monotonic exposure-response for PFOA and PFOS 4 th quartile serum PFOA: OR=1.97 (1.44, 2.70) 4 th quartile serum PFOS: OR=1.48 (0.99, 2.22) Monotonic exposure-response for PFOA

Reference, Location	Study population	PFOS serum	PFHxS serum	PFOA serum	Outcome	Difference detected (95% CI)
2000000	population	level	level	level		
Gleason 2015 NHANES, 2007- 2010 4,333 aged ≥12 year	4,333 aged ≥12 years	11.3	1.8	3.7	Hyperuricemia	Serum PFAS quartiles, odds ratios PFOA PFOS PFHxS 2nd 1.46 (1.16, 1.85) 1.18 (0.88, 1.56) 0.83 (0.64, 1.09) 3rd 1.74 (1.35, 2.25) 1.09 (0.81, 1.47) 1.14 (0.88, 1.48) 4th 1.88 (1.37, 2.58) 1.20 (0.88, 1.63) 1.13 (0.89, 1.43)
					Elevated ALT	2 nd 1.42 (1.11, 1.83) 1.29 (0.93, 1.78) 1.35 (1.01, 1.80) 3 rd 1.55 (1.14, 2.10) 1.26 (0.88, 1.81) 1.37 (1.06, 1.77) 4 th 1.51 (1.18, 1.94) 1.23 (0.87, 1.74) 1.18 (0.94, 1.49)
				Elevated GGT	2 nd 1.09 (0.79, 1.52) 1.16 (0.87, 1.55) 0.97 (0.80, 1.81) 3 rd 1.11 (0.80, 1.52) 1.14 (0.87, 1.51) 1.02 (0.81, 1.30) 4 th 1.34 (1.00, 1.80) 1.10 (0.82, 1.47) 0.91 (0.75, 1.11)	
					Elevated AST	2 nd 1.31 (1.03, 1.65) 1.03 (0.78, 1.36) 1.29 (0.97, 1.71) 3 rd 1.26 (0.97, 1.64) 1.14 (0.87, 1.50) 1.29 (0.96, 1.73) 4 th 1.39 (1.06, 1.80) 0.91 (0.69, 1.21) 1.30 (0.94, 1.78)
					Total Bilirubin	2 nd 1.31 (0.98, 1.75) 1.44 (1.12, 1.84) 1.10 (0.86, 1.40) 3 rd 1.66 (1.19, 2.32) 1.65 (1.25, 2.18) 1.40 (1.04, 1.88) 4 th 1.68 (1.31, 2.14) 1.51 (1.06, 2.15) 1.32 (0.97, 1.81)
Gallo 2012 C8	46,452 ≥18 years	20.3		28.0	Ln-ALT (IU/L) Ln-GGT (IU/L) Ln-Direct bilirubin (mg/dL) Elevated ALT Elevated GGT Elevated Direct bilirubin	$\begin{array}{c} \text{Ln-PFOA} & \text{Ln-PFOS} \\ \beta = 0.022 \ (0.018, 0.025) & \beta = 0.020 \ (0.014, 0.026) \\ \beta = 0.015 \ (0.010, 0.019) & \beta = 0.008 \ (-0.000, 0.016) \\ \beta = 0.001 \ (-0.002, 0.004) & \beta = 0.029 \ (0.024, 0.034) \\ \\ \text{tenth decile odds ratios;} \\ 1.54 \ (1.33, 1.78) & 1.25 \ (1.08, 1.44) \\ 1.06 \ (0.92, 1.20) & 0.94 \ (0.83, 1.07) \\ 1.01 \ (0.66, 1.53) & 1.23 \ (0.82, 1.83) \\ \text{No monotonic exposure-response relationships} \end{array}$

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Darrow 2016 C8	30,723 aged ≥20 years	lever	level	28.2	Ln-ALT (IU/L) Ln-GGT (IU/L) Ln-Direct bilirubin (mg/dL)	Fifth quintile, estimated cumulative serum PFOA $\beta = 0.058 \ (0.040, 0.076)$ $\beta = 0.020 \ (-0.004, 0.044)$ $\beta = -0.017 \ (-0.032, -0.001)$ monotonic exposure-response for ALT Fifth quintile, estimated cumulative serum PFOA
					Elevated ALT Elevated GGT Elevated Direct bilirubin	OR=1.16 (1.02, 1.33) OR=0.96 (0.85, 1.09) OR=0.95 (0.66, 1.37) No monotonic exposure-response relationships
					Any liver disease	5 th quintile cumulative exposure: OR=0.95 (0.70, 1.27)
Melzer 2010 NHANES, 1999- 2000, 2003-2006	3,974 aged ≥20 years	17.9		3.5	Ever reported: Arthritis Asthma COPD Diabetes Heart Disease Liver disease (current) Women: Thyroid disease ever Thyroid disease with current medication Men: Thyroid disease ever Thyroid disease with current medication	4 th quartile serum concentrations, ORs: PFOA PFOS 1.28 (0.97, 1.68) 0.74 (0.53, 1.04) 0.93 (0.64, 1.36) 0.79 (0.50, 1.26) 0.85 (0.54, 1.34) 0.58 (0.43, 0.76) 0.69 (0.41, 1.16) 0.87 (0.57, 1.31) 1.08 (0.70, 1.69) 0.91 (0.50, 1.64) 0.61 (0.21, 1.78) 0.95 (0.39, 2.29) 1.64 (1.09, 2.46) 1.15 (0.70, 1.91) 1.86 (1.12, 3.09) 1.31 (0.72, 2.36) Exposure-response trends were non-monotonic 1.58 (0.74, 3.39) 1.58 (0.72, 3.47) 1.89 (0.60, 5.90) 1.89 (0.72, 4.93) Exposure-response trends were non-monotonic
Lin 2010 NHANES, 1999- 2000, 2003-2004	2,216 aged ≥18 years				ALT (IU/L) Ln-GGT (IU/L) Total bilirubin (µM)	Regression coefficient (s.e.) per unit increase in log serum PFAS PFOA PFOS PFHxS 1.86 (0.62) 1.01 (0.53) 0.19 (0.48) 0.08 (0.03) 0.01 (0.03) -0.00 (0.02) -0.09 (0.20) -0.30 (0.24) 0.38 (0.20)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Winquist 2014b C8	32,254 aged ≥20 years			26.1	Hypertension Hypercholesterolemia Men 40-59 Coronary artery disease	Fifth quintile, cumulative serum PFOA, HR=0.98 (0.91, 1.06) Fifth quintile, cumulative serum PFOA, HR=1.19 (1.11, 1.28) Fifth quintile, cumulative serum PFOA, HR=1.44 (1.28, 1.62) Fifth quintile, cumulative serum PFOA, HR=1.07 (0.93, 1.23) None of the analyses had a monotonic trend.
Mattsson 2015 Sweden: male cohort of farmers and rural residents	231 cases of coronary heart disease and 231 controls	22.4	1.6	4.1	Coronary heart disease	Quartile of serum PFAS, Odds ratios (95% CI): PFOS PFOA PFHxS 2nd 0.82 (0.46, 1.45) 0.79 (0.44, 1.43) 0.91 (0.51, 1.63) 3rd 1.30 (0.74, 2.26) 1.18 (0.67, 2.06) 1.00 (0.56, 1.77) 4th 1.07 (0.60, 1.92) 0.88 (0.50, 1.55) 0.95 (0.54, 1.67)
Shankar 2012 NHANES, 1999- 2000, 2003-2004	1,216 aged ≥40 years			4.2	Cardiovascular disease Peripheral arterial disease Either CVD or PAD Coronary heart disease	Serum PFOA level, Odds ratios (95% CI) CVD PAD Either CVD or PAD 2nd 1.58 (0.80, 3.12) 0.75 (0.37, 1.52) 1.41 (0.81, 2.45) 3rd 1.77 (1.04, 3.02) 1.18 (0.47, 2.96) 1.72 (1.13, 2.64) 4th 2.01 (1.12, 3.60) 1.78 (1.03, 3.08) 2.28 (1.40, 3.71) Coronary heart disease 2nd 0.90 (0.37, 2.23) 4.39 (1.44, 13.4) 3rd 1.90 (0.89, 4.08) 3.94 (1.48, 10.5)
Steenland 2009 C8	46,294 aged ≥18 years	19.6		26.6	Stroke Log total cholesterol	1.70 (0.89, 4.06) 3.94 (1.46, 10.3) 4 th 2.24 (1.02, 4.94) 4.26 (1.84, 9.89) 10 th decile, serum PFAS: change in the log total cholesterol, (SE) PFOA: 0.05 (0.004) PFOS: 0.06 (0.004) (equivalent to an increase of 11-12 mg/dL in total cholesterol)
					Log total cholesterol Log HDL Log LDL Log triglycerides Log total cholesterol/HDL	Linear regression coefficient (SD) Log serum PFOA Log serum PFOS 0.01112 (0.00076) 0.02660 (0.00140) 0.00276 (0.00094) 0.00355 (0.00173) 0.01499 (0.00121) 0.04176 (0.00221) 0.00169 (0.00219) 0.01998 (0.00402) 0.00831 (0.00110) 0.02290 (0.00202) PFOA (OR, 95% CI) PFOS 2nd 1.21 (1.12, 1.31) 1.14 (1.05, 1.23)
					High total cholesterol (≥240 mg/dL)	2 nd 1.21 (1.12, 1.31) 1.14 (1.05, 1.23) 3 rd 1.33 (1.23, 1.43) 1.28 (1.19, 1.39) 4 th 1.38 (1.28, 1.50) 1.51 (1.40, 1.64)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Fitz-Simon 2013 C8 (longitudinal study)	560 aged >20 years	8.2		30.8	Total cholesterol LDL HDL Triglycerides	Percentage <u>decrease</u> in lipid per halving of serum PFAS PFOA: 1.65 (0.32, 2.97) PFOS: 3.20 (1.63, 4.76) PFOA: 3.58 (1.47, 5.66) PFOS: 4.99 (2.46, 7.44) PFOA: 1.33 (-0.21, 2.85) PFOS: 1.28 (-0.59, 3.12) PFOA: -0.78 (-5.34, 3.58) PFOS: 2.49 (-2.88, 7.57)
Nelson 2010 NHANES, 2003- 2004	860 aged ≥20 years	21.0	1.8	3.9	Total Cholesterol	Mean difference in lipid PFOS PFOA PFHxS 2 nd 6.12 (-4.45, 16.7) 5.40 (-2.11, 12.9) -3.22 (-11.8, 5.30) 3 rd 5.07 (-4.24, 14.4) 7.50 (-3.71, 18.7) -2.27 (-8.95, 4.41) 4 th 13.42 (3.83, 23.0) 9.76 (-0.23, 19.7) -7.01 (-13.2, -0.79)
					LDL cholesterol	2 nd 2.78 (-12.7, 18.3) 6.07 (-8.65, 20.8) -4.49 (-12.4, 3.40) 3 rd 0.38 (-9.64, 10.4) 0.71 (-12.6, 14.0) -4.07 (-13.4, 5.28) 4 th 8.50 (-7.10, 24.1) 2.94 (-10.8, 16.7) -9.67 (-20.1, 0.71)
					Non-HDL cholesterol	2 nd 4.64 (-6.87, 16.2) 7.41 (-0.97, 15.8) -3.94 (-12.2, 4.37) 3 rd 4.97 (-5.30, 15.2) 9.11 (-2.44, 20.7) -2.02 (-9.32, 5.28) 4 th 12.55 (1.62, 23.5) 11.03 (1.20, 20.9) -9.32 (-15.9, -2.77)
Fisher 2013 Canadian Health Measures Survey	2,700 aged ≥18 years 2,345	8.4	2.18	2.46	High cholesterol	Odds ratios (95% CI) PFOA PFOS 2 nd 1.61 (1.02, 2.53) 3 rd 1.26 (0.76, 2.07) 4 th 1.50 (0.86, 2.62) 1.36 (0.87, 2.12) 1.57 (0.93, 2.64)
Fu 2014 China	133 aged 0 to 88 years	1.47		1.43	High total cholesterol	Odds ratios PFOA PFOS 2nd 0.82 (0.14, 4.81) 0.57 (0.12, 2.81) 3rd 2.60 (0.56, 12.1) 0.82 (0.17, 3.91) 4th 0.55 (0.09, 3.31) 2.27 (0.47, 10.9)
					High LDL cholesterol	2 nd 0.55 (0.11, 2.82) 1.06 (0.25, 4.53) 3 rd 1.70 (0.40, 3.49) 1.11 (0.25, 4.82) 4 th 0.71 (0.14, 3.49) 2.27 (0.50, 10.4)
					High Triglycerides	2 nd 1.73 (0.57, 5.21) 0.52 (0.17, 1.56) 3 rd 1.03 (0.33, 3.20) 0.93 (0.31, 2.80) 4 th 1.97 (0.59, 6.55) 1.26 (0.41, 3.90)

Reference, Location	Study population	PFOS serum	PFHxS serum	PFOA serum	Outcome	Difference detected (95% CI)
		level	level	level		
Winquist 2014a C8	32,254 aged ≥20 years (community & worker cohort)			26.1	Functional thyroid disease	Quintiles of serum PFOA, cumulative exposure, hazard ratios All Females Males 2nd 1.21 (1.01, 1.45) 1.24 (1.02, 1.51) 1.12 (0.69, 1.79) 3rd 1.17 (0.97, 1.41) 1.27 (1.04, 1.55) 0.83 (0.51, 1.37) 4th 1.27 (1.06, 1.52) 1.36 (1.12, 1.66) 1.01 (0.63, 1.62) 5th 1.28 (1.06, 1.53) 1.37 (1.11, 1.68) 1.05 (0.66, 1.66)
					Hyperthyroidism	2 nd 0.94 (0.62, 1.42) 1.04 (0.65, 1.67) 0.71 (0.29, 1.74) 3 rd 1.12 (0.75, 1.68) 1.33 (0.84, 2.11) 0.57 (0.23, 1.43) 4 th 1.22 (0.82, 1.82) 1.45 (0.92, 2.28) 0.70 (0.30, 1.66) 5 th 1.20 (0.80, 1.81) 1.39 (0.86, 2.26) 0.74 (0.33, 1.65)
28,541 (community cohort only)			24.2	Hypothyroidism	2 nd 1.31 (1.06, 1.63) 1.32 (1.04, 1.67) 1.43 (0.77, 2.66) 3 rd 1.27 (1.01, 1.58) 1.33 (1.05, 1.69) 1.12 (0.59, 2.14) 4 th 1.30 (1.04, 1.62) 1.34 (1.06, 1.70) 1.32 (0.71, 2.45) 5 th 1.40 (1.12, 1.75) 1.47 (1.15, 1.88) 1.36 (0.74, 2.48)	
			24.2	Functional thyroid disease	2 nd 1.24 (1.03, 1.49) 1.24 (1.02, 1.52) 1.06 (0.62, 1.80) 3 rd 1.21 (1.00, 1.46) 1.27 (1.04, 1.56) 0.80 (0.46, 1.40) 4 th 1.32 (1.09, 1.59) 1.36 (1.11, 1.66) 1.02 (0.59, 1.75) 5 th 1.36 (1.11, 1.66) 1.36 (1.10, 1.69) 1.21 (0.69, 2.12)	
					Hyperthyroidism	2 nd 0.96 (0.62, 1.47) 1.07 (0.66, 1.73) 0.60 (0.22, 1.64) 3 rd 1.13 (0.74, 1.73) 1.36 (0.85, 2.18) 0.47 (0.17, 1.32) 4 th 1.23 (0.81, 1.88) 1.45 (0.91, 2.33) 0.57 (0.21, 1.58) 5 th 1.28 (0.81, 2.01) 1.45 (0.88, 2.40) 0.70 (0.24, 2.06)
					Hypothyroidism	2 nd 1.34 (1.07, 1.68) 1.31 (1.03, 1.66) 1.53 (0.76, 3.11) 3 rd 1.31 (1.04, 1.64) 1.32 (1.04, 1.68) 1.26 (0.60, 2.61) 4 th 1.35 (1.07, 1.70) 1.33 (1.04, 1.69) 1.55 (0.75, 3.18) 5 th 1.49 (1.17, 1.89) 1.43 (1.10, 1.84) 1.89 (0.90, 3.97)
					community cohort only: Functional thyroid disease Hyperthyroidism Hypothyroidism	Prospective Analyses: cumulative exposure, quintile 5 results: 1.13 (0.83, 1.54)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Starling 2014 Norway Knox 2011 C8	891 pregnant women, aged 19-44 50,113 aged ≥20 years	13.03 Median o	o.60 r geometric r PFOA and	2.25	per ln-ng/ml PFAS LDL cholesterol per ln-ng/ml PFAS Ln-triglycerides per ln-ng/ml PFAS Thyroxine (total T4) T3 uptake Thyroid stimulating hormone (TSH) Serum albumin	Linear regression coefficients for lipids (mg/dL) PFOA PFOS PFHxS 2nd 1.49 (-6.49, 9.48) -3.35 (-10.3, 3.64) 0.65 (-6.87, 8.17) 3rd 3.54 (-4.51, 11.6) 3.06 (-4.93, 11.1) 1.62 (-6.08, 9.32) 4th 3.90 (-5.00, 12.8) 7.59 (-0.42, 15.6) 4.25 (-3.88, 12.4) 2.58 (-4.32, 9.47) 8.96 (1.70, 16.2) 3.00 (-1.75, 7.76) 2nd 0.94 (-6.08, 7.96) -3.23 (-9.28, 2.83) 0.44 (-6.19, 7.08) 3rd 4.16 (-3.19, 11.5) 2.60 (-4.49, 9.70) 0.50 (-6.15, 7.16) 4th 3.35 (-4.35, 11.1) 5.51 (-1.62, 12.6) 1.48 (-5.89, 8.85) 2.25 (-3.97, 8.48) 6.48 (-0.07, 13.0) 1.92 (-2.50, 6.33) 2nd 0.03 (-0.04, 0.11) 0.00 (-0.06, 0.07) -0.04 (-0.11, 0.02) 3rd 0.01 (-0.08, 0.09) -0.03 (-0.10, 0.05) -0.02 (-0.10, 0.05) 4th -0.04 (-0.12, 0.04) 0.00 (-0.07, 0.07) -0.02 (-0.09, 0.05) 0.00 (-0.07, 0.06) -0.02 (-0.09, 0.04) -0.01 (-0.05, 0.03) No confidence intervals were provided. (Many p-values were presented as less than a value, so CIs could not be estimated.) PFOA results: PFOA was associated with increased thyroxine and decreased T3 uptake in women of all ages and in men who were >50 years of age. PFOA was associated with a slight increase in albumin among all participants. T3 uptake was lower in women than in men; TSH was lower in men than in women; T4 was higher in women than in men; and albumin was higher in men.
						PFOS results: PFOS was associated with increased thyroxine, decreased T3 uptake, and increased albumin in all participants. Thyroxine was higher in women than in men; T3 update was lower in women than in men; and albumin was lower in women than in men.

Reference, Location	Study population	PFOS serum	PFHxS serum	PFOA serum	Outcome	Difference detected (95% CI)
Locution	population	level	level	level		
Wen 2013 NHANES, 2007- 2010	1,181 aged ≥20 years 672 males 509 females	17.14 11.09	2.60 1.41	4.70 3.53	Total T4 (µg/mL) Ln-free T4 (ng/dL) Total T3 (ng/dL) Ln-free T3 (pg/mL) Ln TSH (mIU/L)Ln Thyroglobulin (ng/mL) Total T4 (µg/mL) Ln-free T4 (ng/dL) Total T3 (ng/dL) Ln-free T3 (pg/mL) Ln TSH (mIU/L) Ln Thyroglobulin (ng/mL) Subclinical hypothyroidism: Men: Women: Subclinical hyperthyroidism: Men:	Regression coefficient per unit increase in log PFAS, Men: PFOA PFOS PFHxS 0.000 (-0.280, 0.280) -0.020 (-0.223, 0.183) -0.032 (-0.175, 0.111) -0.010 (-0.041, 0.022) -0.009 (-0.034, 0.017) -0.016 (-0.029, -0.003) 0.775 (-3.048, 4.598) -1.111 (-3.856, 1.634) -0.081 (-1.698, 1.536) 0.013 (-0.004, 0.031) 0.002 (-0.008, 0.012) 0.005 (-0.003, 0.012) 0.004 (-0.081, 0.090) 0.003 (-0.070, 0.076) 0.019 (-0.057, 0.524) -0.096 (-0.258, 0.066) -0.047 (-0.149, 0.055) -0.049 (-0.185, 0.087) Women: 0.082 (-0.369, 0.532) 0.087 (-0.143, 0.318) 0.260 (0.108, 0.413) -0.004 (-0.047, 0.039) 0.009 (-0.019, 0.036) 0.003 (-0.024, 0.030) 6.628 (0.545, 12.712) 1.453 (-1.987, 4.891) 4.074 (2.232, 5.916) 0.016 (-0.018, 0.051) -0.007 (-0.024, 0.010) 0.003 (-0.021, 0.026) -0.030 (-0.215, 0.154) -0.048 (-0.156, 0.060) -0.019 (-0.128, 0.090) 0.095 (-0.111, 0.302) 0.135 (-0.007, 0.277) -0.018 (-0.122, 0.087) Odds ratio 1.29 (0.40, 4.10) 1.98 (1.19, 3.28) 1.57 (0.76, 3.25) 7.42 (1.14, 48.1) 3.03 (1.14, 8.07) 3.10 (1.22, 7.86)
Webster 2016 NHANES, 2007- 2008	1,525 aged ≥18 years	14.2	2.0	4.2	Free T3 Free T4 Free T3/Free T4 Thyroid stimulating hormone (TSH) Total T3 Total T4	0.99 (0.13, 7.59) 1.90 (0.53, 6.80) 2.27 (1.07, 4.80) Low iodine & high thyroid peroxidase antibody (TPOAb): percent difference per interquartile ratio increase in log serum PFAS PFOS PFOA PFHxS 4.7 (3.9, 5.5) 4.8 (3.7, 5.8) 3.9 (2.3, 5.5) -4.4 (-7.6, -1.1) -2.7 (-6.1, 0.8) -8.3 (-15.8, -0.2) 9.5 (5.8, 13.2) 7.7 (3.6, 12.0) 13.3 (4.4, 22.9) 17.1 (6.6, 17.7) 16.2 (5.1, 28.5) 27.3 (0.7, 60.9) 12.0 (6.7, 17.7) 12.4 (7.0, 18.1) 13.8 (6.0, 22.1) 2.5 (-1.3, 6.5) 3.9 (-0.3, 8.3) 1.8 (-3.9, 7.8)
Shrestha 2015 Hudson River area, NY	87 aged 55-74	29.8		9.3	Thyroid stimulating hormone (TSH) Free T4 Total T4 Total T3	Regression coefficient, log serum PFAS PFOS 0.129 (-0.023, 0.281) 0.054 (0.002, 0.106) 0.766 (0.327, 1.205) 2.631 (-2.248, 7.510) PFOA 0.102 (-0.047, 0.250) 0.016 (-0.036, 0.069) 0.380 (-0.070, 0.830) 3.032 (-1.725, 7.789)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Ji 2012 Korea	633 aged >12 years	7.96	1.51	2.74	Thyroid stimulating hormone (TSH) Total T4	Regression coefficients, serum PFAS PFOS PFOA PFHxS 0.062 (-0.069, 0.192) -0.066 (-0.220, 0.089) 0.013 (-0.094, 0.120) -0.021 (-0.048, 0.005) -0.020 (-0.051, 0.012) -0.007 (-0.029, 0.015)
Wang 2014 Taiwan	285 pregnant women (average age=29)	12.73	0.81	2.39	Thyroid stimulating hormone (TSH) Free T4 Total T4 Total T3	Regression coefficients, serum PFAS PFOS PFOA PFHxS -0.005 (-0.024, 0.013) 0.011 (-0.057, 0.078) 0.105 (0.002, 0.207) 0.001 (-0.002, 0.003) -0.003 (-0.012, 0.005) -0.010 (-0.023, 0.003) 0.019 (-0.016, 0.053) 0.011 (-0.108, 0.130) -0.130 (-0.316, 0.057) 0.000 (-0.002, 0.001) -0.000 (-0.002, 0.009) -0.002 (-0.005, 0.001)
Webster 2014 Canada	152 pregnant women without thyroid disease, aged ≥25 years	4.8	1.0	1.7	Thyroid stimulating hormone (TSH) Free thyroxine (fT4)	Percent change (compared to the median thyroid function in the study population) per interquartile increase in PFAS among those with high TPOAb: PFOS PFOA PFHxS 69% (15%, 123%) 54% (8%, 100&) 2% (-45%, 48%) -7% (-18%, 3%) -4% (-14%, 5%) -5% (-15%, 4%)
Berg 2015 Norway	375 pregnant women aged 18-43	8.03	0.44	1.53	Thyroid stimulating hormone (TSH) mIU/L Subclinical hypothyroidism (%)	Mean difference in TSH, and proportion with subclinical hypothyroidism per quartile of PFOS 1st 12.8% (n=12) 2nd 0.18 (0.06, 0.31) 17.8% (n=16) 3rd 0.26 (0.13, 0.40) 25.3% (n=24) 4th 0.35 (0.21, 0.50) 31.3% (n=30)
Berg 2016 Norway	370 pregnant women aged 18-43	8.03	0.44	1.53	Thyroid stimulating hormone (TSH) mIU/L	Percent change in TSH per quartile of PFOS 2 nd 4 (-3.1, 11.4) 3 rd 8 (0.6, 15.4) 4 th 10 (1.6, 16.9)
Kato 2016 Japan	392 pregnant women	5.2		1.2	Ln-Thyroid stimulating hormone (TSH) Ln-Free Thyroxine (T4)	Regression coefficient per Ln-PFAS (confidence intervals are estimated) PFOS PFOA -0.214 (p-value presented as <0.001) 0.039 (-0.067, 0.144) 0.061 (-0.039, 0.161) 0.004 (-0.102, 0.110)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Steenland 2013 C8 28,541 community cohort 3,713 worker cohort Total=32, 254 aged ≥20 years	cohort 3,713 worker cohort Total=32, 254 aged			24 113 26	Ulcerative colitis	10 year lagged cumulative PFOA quartiles, RR: Retrospective: Prospective: 2nd 1.71 (0.89, 3.27) 1.21 (0.43, 3.34) 3rd 2.05 (1.07, 3.91) 2.16 (0.80, 5.81) 4th 3.05 (1.56, 5.96) 1.51 (0.43, 4.30)
				Rheumatoid arthritis	2 nd 1.53 (0.61, 2.58) 0.31 (0.14, 0.71) 3 rd 1.73 (1.10, 2.71) 0.90 (0.41, 0.72) 4 th 1.35 (0.87, 2.11) 0.32 (0.14, 0.72) 10 year lagged cumulative PFOA quartiles, retrospective analyses, RR: 2 nd Q 3 rd Q 4 th Q	
					Crohn's disease Type 1 diabetes Lupus Multiple sclerosis	0.80 (0.32, 1.99) 0.97 (0.36, 2.60) 0.69 (0.26, 1.82) 0.50 (0.05, 4.90) 1.32 (0.14, 12.4) 0.71 (0.07, 7.14) 0.79 (0.27, 2.34) 1.26 (0.40, 4.03) 0.61 (0.19, 1.91) 1.16 (0.54, 2.47) 1.62 (0.75, 3.52) 1.32 (0.61, 2.84)
Innes 2011 C8	49,432 aged ≥21 years	20.3		28.2	Osteoarthritis	PFOA odds ratio PFOS 2 nd 1.16 (1.03, 1.31) 0.91 (0.81, 1.01) 3 rd 1.21 (1.07, 1.36) 0.94 (0.84, 1.06) 4 th 1.42 (1.26, 1.59) 0.76 (0.68, 0.85) 1.07 (1.04, 1.1) per 1-unit increment in ln-PFOA
Uhl 2013 NHANES, 2003- 2008	4,102 aged 20-84	24.6 (mean)		5.4 (mean)	Osteoarthritis	PFOA All Female Male 2nd 1.32 (0,78, 2.23) 1.44 (0.80, 2.62) 0.97 (0.42, 2.27) 3rd 1.20 (0.72, 2.00) 1.18 (0.67, 2.08) 0.98 (0.46, 2.08) 4th 1.55 (0.99, 2.43) 1.98 (1.24, 3.19) 0.82 (0.40, 1.70) Per ln-PFOA: 1.20 (0.96, 1.49) 1.35 (1.02, 1.79) 0.89 (0.67, 1.19) PFOS 2nd 1.04 (0.58, 1.85) 0.88 (0.46, 1.70) 1.32 (0.41, 4.25) 3rd 1.99 (1.14, 3.49) 1.92 (0.98, 3.75) 1.86 (0.55, 6.25) 4th 1.77 (1.05, 2.96) 1.73 (0.97, 3.10) 1.56 (0.54, 4.53) Per ln-PFOS: 1.15 (0.94, 1.40) 1.22 (0.94, 1.58) 0.95 (0.73, 1.23)

Reference, Location	Study population	PFOS serum	PFHxS serum	PFOA serum	Outcome	Difference detected (95% CI)
Location	population	level	level	level		
Lin 2014 NHANES, 2005- 2008 2,339 aged ≥20 years 1,192 men 1,147 women 842 not in menopause 305 in menopause	15.32		3.96	Total Lumbar spine bone mineral density (BMD) (g/cm²)	Regression coefficient for change in BMD per unit-increase in ln-PFOA Men Women not in menopause Women in menopause 0.006 (-0.014, 0.026) 0.001 (-0.020, 0.022) 0.018 (-0.014, 0.049) Regression coefficient for change in BMD per unit-increase in ln-PFOS Men Women not in menopause Women in menopause 0.000 (-0.013, 0.013) -0.022 (-0.038, -0.007) -0.004 (-0.026, 0.034)	
				Total hip BMD (g/cm ²)	Regression coefficient for change in BMD per unit-increase in ln-PFOA Men Women not in menopause Women in menopause -0.002 (-0.023, 0.019) 0.008 (-0.010, 0.027) 0.022 (-0.011, 0.055) Regression coefficient for change in BMD per unit-increase in ln-PFOS Men Women not in menopause Women in menopause -0.003 (-0.016, 0.010) 0.000 (-0.017, 0.017) 0.017 (-0.012, 0.047)	
		All frac		All fractures	OR per unit-increase in ln-PFOA Men Women not in menopause Women in menopause 0.84 (0.67, 1.07) 0.98 (0.75, 1.28) 1.53 (0.63, 3.74) OR per unit-increase in ln-PFOS Men Women not in menopause Women in menopause	
				Hip fracture	0.92 (0.73, 1.16) 0.97 (0.75, 1.24) 1.59 (0.88, 2.86) OR per unit-increase in ln-PFOA Men Women not in menopause Women in menopause 0.64 (0.39, 1.06) 1.59 (0.57, 4.46) 0.48 (0.06, 4.16) OR per unit-increase in ln-PFOS Men Women not in menopause Women in menopause	
					Wrist fracture	1.07 (0.76, 1.52) 1.12 (0.62, 2.03) 0.83 (0.23, 3.00) OR per unit-increase in ln-PFOA Men Women not in menopause Women in menopause 1.12 (0.75, 1.70) 1.07 (0.65, 1.77) 1.21 (0.46, 3.13) OR per unit-increase in ln-PFOS Men Women not in menopause Women in menopause
			Spine fracture	1.09 (0.72, 1.66) 1.04 (0.63, 1.72) 1.22 (0.61, 2.45) OR per unit-increase in ln-PFOA Men Women not in menopause Women in menopause 1.54 (0.85, 2.79) 1.83 (0.59, 5.61) 0.84 (0.46, 1.53) OR per unit-increase in ln-PFOS Men Women not in menopause Women in menopause 1.27 (0.67, 2.42) 0.52 (0.15, 1.86) 1.12 (0.26, 4.78)		

Reference,	Study	PFOS	PFHxS	PFOA	Outcome Difference detected (95% CI)
Location	population	serum	serum	serum	· '
		level	level	level	
Khalil 2016 NHANES, 2009- 2010	1,914 aged ≥12 years 956 men 959 women 590 premenopausal 368 postmenopausal	9.70 Means: 12.70 15.10 10.30	1.70 Means: 2.50 3.10 1.90	5.20 Means: 3.70 4.10 3.30	Total femur

			T	1	1	
					3 rd -0.026 4 th -0.023 Per Ln-PFOS -0.011 PFHxS Men 2 nd 0.015 3 rd 0.021	3 (-0.064, 0.018)
					Odds ratio for osteoporosis	s in women
					PFOA	PFOS PFHxS
					2 nd 1.25 (0.38, 4.06)	0.42 (0.13, 1.32) 9.29 (1.81, 47.6)
					3 rd 1.23 (0.37, 4.05)	0.83 (0.45, 1.51) 8.06 (1.84, 35.3)
					4 th 2.59 (1.01, 6.67) Per Ln: 1.84 (1.17, 2.90)	1.07 (0.36, 3.19) 13.20 (2.72, 64.2) 1.14 (0.68, 1.94) 1.64 (1.14, 2.38)
Reference,	Study	PFOS	PFHxS	PFOA	, , , , , , , , , , , , , , , , , , , ,	nce detected (95% CI)
					Outcome Differen	ice delected (95% C1)
Location	population	serum	serum	serum		
Y 1 2011	102 1 10	level	level	level		
Looker 2014 C8	403 aged >18 years	8.32		33.74	Influence Tema D	PFOA quartiles regression coefficient, log10-PFOA
C8					Influenza Type B Geometric mean	GMT log10-titer rise log10-titer ratio 1st 49.5 (38.1, 64.1)
					antibody titer (GMT)	2 nd 46.0 (35.3, 60.0)03 (19, .13) .05 (09, .19)
					Log10-titer rise	3 rd 43.6 (33.1, 57.3)02 (19, .15) .07 (07, .22)
					Log10-titer ratio:	4 th 20.9 (16.6, 28.2)07 (24, .10)03 (17, .12)
					Postvaccine / prevaccine	Regression coefficient per unit increase in log10-PFOA:
						02 (13, .09)02 (11, .08)
						PFOA Odds ratio: Seroconversion Seroprotection
						2 nd 1.43 (0.76, 2.70) 0.76 (0.40, 1.45)
						3 rd 1.39 (0.73, 2.66) 1.13 (0.57, 2.23)
						4 th 0.71 (0.38, 1.36) 0.77 (0.39, 1.50)
						Odds ratio per unit-increase in log10-PFOA:
						0.80 (0.53, 1.21) 1.04 (0.68, 1.60)
						PFOS quartiles regression coefficient, log10-PFOS
i						GMT log10-titer rise log10 titer ratio
						GMT log10-titer rise log10-titer ratio
						1st 42.3 (33.4, 53.4)

	Regression coefficient per unit increase in log10-PFOS:
	.05 (11, .21) .05 (09, .18)
	PFOS Odds ratio: Seroconversion Seroprotection
	2 nd 0.72 (0.39, 1.33) 0.67 (0.35, 1.25)
	3 rd 0.81 (0.42, 1.53) 0.82 (0.42, 1.59)
	4 th 0.87 (0.43, 1.74) 0.73 (0.36, 1.47)
	Odds ratio per unit-increase in log10-PFOS:
	1.17 (0.63, 2.17) 0.85 (0.44, 1.64)
	PFOA quartiles regression coefficient, log10-PFOA
	GMT log10-titer rise log10-titer ratio
Influenza Type	
Geometric mea	
antibody titer (
Log10-titer rise	
Log10-titer rati	
Postvaccine / p	
T ostvaceme / p	PFOA Odds ratio: Seroconversion Seroprotection
	2 nd 0.74 (0.34, 1.59) 0.74 (0.17, 3.28)
	3 rd 1.11 (0.49, 2.50) 1.59 (0.33, 7.70)
	4 th 2.23 (0.90, 5.53) 6.47 (0.91, 45.9)
	2.22 (0.50, 2.62)
	Odds ratio per unit-increase in log10-PFOA: 1.51 (0.89, 2.56) 2.34 (0.91, 6.07)
	PFOS quartiles regression coefficient, log10-PFOS
	GMT log10-titer rise log10-titer ratio 1st 342.3 (256.0, 457.7)
	2 nd 280.4 (197.6, 397.9)04 (21, .14)07 (28, .13)
	3 rd 417.7 (319.0, 547.1) .13 (04, .31) .03 (18, .24)
	4 th 341.8 (258.0, 452.8) .10 (09, .29) .03 (19, .26)
	Regression coefficient per unit increase in log10-PFOS:
	.15 (02, .32) .10 (11, .30)
	PFOS Odds ratio: Seroconversion Seroprotection
	2 nd 0.97 (0.44, 2.14) 0.55 (0.13, 2.37)
	3rd 0.78 (0.35, 1.75) 1.81 (0.32, 10.2)
	4 th 0.94 (0.38, 2.31) 1.26 (0.24, 6.61)
	Odds ratio per unit-increase in log10-PFOS:
	•
	1.10 (0.51, 2.37) 0.93 (0.23, 3.71)
	PFOA quartiles regression coefficient, log10-PFOA
	GMT log10-titer rise log10-titer ratio
	1st 228.9 (161.5, 324.3)
Influenza Type	
	3 rd 104.1 (72.5, 149.6)37 (60,13)07 (28, .14)
	4 th 183.7 (127.3, 265.2)12 (36, .13)22 (43,01)

	755 aged >18 years				Geometric mean antibody titer (GMT) Log10-titer rise Log10-titer ratio: Postvaccine / prevaccine Self-reported cold or flu in last 12 months	Regression coefficient per unit increase in log10-PFOA:
		2200				0.85 (0.62, 1.16) 0.90 (0.55, 1.48)
Reference,	Study	PFOS	PFHxS	PFOA	Outcome	Difference detected (95% CI)
Location	population	serum	serum	serum		
	P - P	_				
Kielsen 2016		level	level	level		
Denmark	12 adults	9.52	0.37	1.69	Diphtheria	% change in antibody per doubling of PFAS PFOA PFOS PFHxS -8.22 (6.44, -20.85) -11.90 (-0.33, -21.92) -13.31 (0.29, -25.07)

Reference,	Study	PFOS	PFHxS	PFOA	Outcome	Reference,
Location	population	serum	serum	serum	Difference detected	Location
		level	level	level	(95% CI)	
Stein 2016b	75 aged 21-49 years	5.22	1.1	2.28	FluMist Seroconversion	RR
New York					measured by immuno-	PFOS PFOA PFHxS
					histochemistry	2 nd 2.6 (0.9, 7.4) 0.6 (0.2, 2.0) 1.1 (0.4, 2.9)
						3 rd 2.4 (0.9, 6.6) 1.8 (0.7, 4.3) 1.7 (0.6, 4.8)
					Serum Immune Markers:	
					Interferon-α2	Mean change between baseline and FluMist response:
						2 nd 8.69 (-23, 40.5) 2.54 (-28, 32.9) -21 (-51, 8.61)
						3 rd -5.6 (-37, 25.7) 10.9 (-19, 40.4) -29 (-64, 5.20)
					Interferon-γ	
						2 nd 8.00 (-34, 49.6) -23 (-61, 14.1) -40 (-76, -3.7)
						3 rd 9.95 (-28, 47.9) -23 (-61, 16.1) -40 (-84, 2.69)
					Tumor necrosis factor-α	
						2 nd -0.06 (-4.6, 4.45) 1.28 (-2.8, 5.39) -5.3 (-9.2, -1.3)
						3 rd 0.59 (-3.6, 4.81) -0.96 (-5.2, 3.26) -4.8 (-9.4, -0.1)
					Interferon-γ-inducible	
					protein 10 (IP-10)	2 nd 12.0 (-44, 67.6) -3.4 (-55, 48.7) -32 (-84, 20.1)
						3 rd -42 (-94, 10.4) -28 (-82, 25.0) -15 (-76, 46.9)
					Monocyte chemo-	
					attractant protein-1	2 nd 4.75 (-60, 69.7) -9.9 (-70, 49.8) -39 (-97, 19.5)
						3 rd 19.0 (-42, 79.8) -2.7 (-64, 58.6) 20.6 (-48, 89.5)
					Macrophage inflam-	
					matory protein-1a	2 nd -1.6 (-22, 18.8) 4.25 (-15, 23.7) -6.6 (-25, 12.1)
						3 rd -0.51 (-20, 19.2) 5.09 (-15, 24.8) -5.0 (-28, 18.4)
					Granulocyte colony-	
					stimulating factor	2 nd -8.7 (-60, 42.8) -1.4 (-49, 45.7) -10 (-56, 35.9)
						3 rd -0.36 (-49, 47.9) -13 (-62, 35.2) 36.6 (-18, 91.3)
					Nasal Secretion Immune	
					Markers:	
					IP-10	2 nd 429 (-1309, 2166) -30 (-1623, 1563) -691 (-2279, 896)
					3.6	3 rd -215 (-1841, 1412) -564 (-2200, 1072) -713 (-2596, 1170)
					Monocyte chemo-	and a 24 (0.2, 12.0) = 0.72 (0.0, 14.2) = 0.77 (14.40.5)
					attractant protein-1	2 nd 2.34 (-9.2, 13.8) 0.72 (-9.9, 11.3) -0.7 (-11, 10.6)
					1.	3 rd -6.7 (-17, 4.04) -6.6 (-18, 4.24) -3.6 (-16, 9.08)
					Mucosal immune-	and 70.0 (01.6 0.6) 17 (00.5 0.5) 20.1 (0.5 0.5)
					globulin A	2 nd 73.9 (-216, 364) -17 (-285, 251) 294 (35.2, 553)
				1		3 rd -88 (-359, 183) 28.7 (-246, 304) 238 (-69, 545)

Table A4. PFAS studies on infertility/subfertility.

Reference	Exposure	Outcome	OR/FR/β & 95% CI
Fei 2009	PFOA and PFOS in blood samples at	Subfecundity	Infertility OR = $1.77 (1.06, 2.95)$ and $2.54 (1.47, 4.39)$ for
Danish study	gestational week 17	(n= 1240 women)	highest vs lowest quartile of PFOS and PFOA, respectively
			Fecundity OR (FOR) = 0.74 (0.58, 0.93) and 0.60 (0.47, 0.76) for highest vs lowest quartile of PFOS and PFOA, respectively FORs <1 indicate decreased fecundity and a longer TTP
Whitworth 2012a	PFOA and PFOS in blood samples at	Subfecundity	ORs = $0.7 \ 0.4-1.3$) for highest PFOS quartile and $0.5 \ (0.2-1.2)$
Norway study	gestational week 17	(n=910)	for highest PFOA quartile among primiparous women and there
1 tor way staay	gesturional work 17	(11) 10)	was a monotonic exposure-response relationship for PFOS
Buck Louis 2013	PFOS, PFOA, PFNA and 4 other PFCs in	Fecundability ORs (TTP)	FORs about 1 for PFOS, PFOA, and PFNA
LIFE study	serum	(n=501 couples followed for 12	
ZII Z study	544441	months)	
Velez 2015	PFOS	Female fecundity odds ratio (FOR)	FOR = 0.89 (0.83–0.94), 0.91 (95% CI 0.86–0.97), and 0.96
MIREC study	PFOA	as measured by TTP	(0.91–1.02) per one SD increase in log-transformed serum
	PFHxS	Infertility	concentrations of PFOA, PFHxS, and PFOS, respectively
	Measured in the 1 st trimester		
		(n= 1743 women recruited before	ORs for infertility = $1.31 (1.11-1.53)$ for PFOA
		14 weeks of gestation)	1.14 (0.98–1.34) for PFOS, and 1.27 (1.09–1.48) for PFHxS
Jorgensen 2014	PFOA, PFOS, PFHxS and PFNA	Fecundability ratios (FRs) infertility (n= 938 women; 448 were from Greenland, 203 from Poland, and 287 from Ukraine)	log-scale FR = 0.80 (0.69-0.94) and 0.90 (0.76, 1.07) for PFNA and PFOS, respectively, in pooled sample FRs in Greenland ranged from 0.71-0.90 for categorical analyses and there were monotonic exposure-response relationships for PFOA, PFOS, and PFNA FRs in Poland were 0.90 and 0.94 for categorical analyses of PFOS and PFHxS, respectively FRs in Ukraine were 0.93 and 0.88 for categorical analyses of PFOS and PFNA, respectively log-scale OR = 1.53 (1.08-2.15), 1.11 (0.74, 1.66), and 1.39 (0.93, 2.07) for PFNA, PFOA, and PFOS, respectively, and infertility in pooled sample
			ORs in Greenland ranged from 1.22-2.15 for categorical analyses and there was a monotonic exposure-response relationships for PFOA

			ORs in Poland were 1.41 and 1.92 for categorical analyses of PFOA and PFOS, respectively, and there were monotonic exposure-response relationships for PFOA
			ORs in Ukraine ranged from 1.17-1.22 for categorical analyses of PFOA, PFOS, and PFNA
			associations were weaker in a sensitivity analysis of primiparous women
Reference	Exposure	Outcome	OR/FR/β & 95% CI
Bach 2015a	plasma PFOS and PFOA	Time to pregnancy (TTP)	PFOA:
Danish study	r	r	FRs ranging from 0.74-0.86 comparing the highest and lowest
		(n=440 for sample 1 and n= 1161 for sample 2)	quartiles
			FRs was 0.74 and 0.82 comparing the highest and lowest
			quartiles in nulliparous women in sample 2 and the pooled
			analysis, respectively, and the log FR were 0.67 and 0.84 and
			there was an exposure-response relationship for sample 2
			FRs ranged from 0.74-0.76 comparing the highest and lowest
			quartiles in parous women and the log FRs ranged from 0.66-0.72 and there was an exposure-response relationship in the pooled sample
			PFOS: FRs ranged from 0.69-0.97 comparing the highest and lowest
			quartiles in nulliparous women and the log FR was 0.62 in sample 2 and 0.78 in the pooled analysis
			FRs was 0.93 and 0.97 comparing the highest and lowest quartiles in parous women in sample 2 and the pooled analysis, respectively, and the log FR ranged from 0.85-0.91
Vested 2013	In utero exposure to PFOA and PFOS	Adult male semen quality,	PFOA was associated with a lower percentage of adjusted sperm
Danish study	measured in maternal blood samples from	testicular volume, and reproductive	concentration (-34%, -58, 5), total sperm count (-34%, -62,
	pregnancy week 30	hormone levels (n=169)	12), and morphologically normal spermatozoa (-19%, -42, 13) and with higher adjusted levels of luteinizing hormone (6 IU/L,
	Maternal and adult son questionnaires on		-11, 27) and follicle-stimulating hormone (15 IU/L, -8, 44).
	dietary/health and lifestyle habits		Monotonic exposure-response relationships for sperm concentration and LH and FSH levels
	Mothers' median plasma concentrations of		
	PFOA and PFOS were 3.8 ng/mL (2.8–4.7		

	ng/mL) and 21.2 ng/mL (17.4–26.5		PFOS was associated with a lower percentage of adjusted total
	ng/mL), respectively		sperm count (-23%, -56, 38) and morphologically normal
			spermatozoa (-14%, -39, 20) and with higher adjusted levels of
			follicle-stimulating hormone (20 IU/L, -5, 51). Monotonic
			exposure-response relationships for morphologically normal
			spermatozoa and FSH levels
Bach 2016a systematic review	PFOS and PFOA measured in blood	Fertility measured by:	Men: Inconsistent results for semen volume, sperm
			concentration, total sperm count, motility, and morphology;
	Men: Average exposure	Reproductive hormones and TTP	levels of testosterone, free androgen index/free testosterone,
	levels in the non-occupational studies	in men and women	estradiol, SHBG, LH, FSH, and inhibin B; and TTP
	ranged from 4.6-44.7 ng/mL for PFOS and	Semen characteristics	
	1.3-9.2 ng/mL for		Women: Inconsistent results except for mostly positive
	PFOA	Men: n's varied from 56-857	associations for infertility and fecundability in parous women
	Women: Average exposure levels ranged	Women: TTP n's varied from 222-	
	from 3.8-36.3 ng/mL for PFOS and 1.5-5.6	1743; hormone n's varied from	
	ng/mL for PFOA	178-825	

Table A5. PFAS studies on pregnancy-induced hypertension/pre-eclampsia.

Reference	Exposure	Outcome	OR/HR & 95% CI
Stein 2009	PFOA	Pre-eclampsia	For exposures >90 th percentile:
C8 project	PFOS		Pre-eclampsia: OR = 1.6 (1.2, 2.3) for PFOS; OR <1 for PFOA
		(n=1,845 pregnancies for PFOA	
		and 5,262 pregnancies for PFOS)	
Savitz 2012a	Serum PFOA levels at the time of	Preeclampsia	OR = 1.2 (1.0-1.6) for highest vs lowest quartile
C8 project	pregnancy from drinking water		
	contaminated by chemical plant releases	(n= 11,737 pregnancies)	
	and d'a ser malalada an an PECA		
	analysis uses modeled serum PFOA estimates		
Starling 2014	PFOA, PFOS, and PFHxS in	Preeclampsia	PFOS: HR = 1.09 (0.75, 1.58) for highest vs lowest quartile and
Norwegian Mother and Child	maternal plasma extracted midpregnancy	Trecelampsia	HR = 1.13 (0.84, 1.52) for per ln-unit
Cohort Study	maternar plasma extracted inapregnancy	(n= 466 cases, 510 noncases)	1114 – 1.13 (0.04, 1.32) for per in unit
Conort Study		(ii= 100 cases, 510 noneases)	PFHxS: HRs <1
Avanasi 2015	Estimated and simulated PFOA	Preeclampsia	OR in 12 simulations ranged between 1.10 and 1.12
C8 project	concentrations		
		(n= 10,149 participants for each	OR = 1.11 (0.99, 1.24) from original exposure assignments
		of the 12 Monte Carlo simulations	
		[500 iterations per simulation])	
Savitz 2012b	Historical estimates of serum PFOA from	Pregnancy-induced hypertension	OR = 1.2 (0.8, 1.7) using survey data and comparing the highest
C8 project	a fate and transport model using address at	(PIH) (n=224 cases and n=3616	vs lowest quintile of PFOA (Bayesian calibration)
	delivery (birth records) and a survey with	controls)	
	residential history data		
Darrow 2013	PFOA and PFOS measurements	Pregnancy-induced hypertension	ORs = 3.16 (1.35, 7.38) and 1.56 (0.72, 3.38) for the highest vs
C8 project	11 Off and 11 Ob measurements	(n=106 and n=1630 total births)	lowest quintile of PFOA and PFOS, respectively
1 J			1

Table A6. PFAS studies on adverse birth outcomes.

Reference	Exposure	Outcome	OR/β & 95% CI
Apelberg 2007	PFOA and PFOS in cord serum	Gestational age, birth weight,	birth weight per ln-unit: $\beta = -69$ g (-149 to 10) for PFOS
Baltimore THREE study	samples	and birth size	and -104 g (-213 to 5) for PFOA
		(n = 293)	
			ponderal index per ln-unit: $\beta = -0.074 \text{ g/cm}^3 \times 100 \text{ (}-$
			$0.123 \text{ to } -0.025) \text{ for PFOS and} = -0.070 \text{ g/cm}^3 \times 100 \text{ (} -$
			0.138 to -0.001) for PFOA
			head circumference per ln-unit: $\beta = -0.32$ cm (-0.56 to $-$
			0.07) for PFOS and -0.41 cm (-0.76 to -0.07) for PFOA
			,
			length per ln-unit: β = -0.10 cm (-0.64 to 0.44) for PFOA
Fei 2007	PFOS and PFOA	Preterm birth, low birth	$\beta = -10.63 \text{ g } (-20.79, -0.47) \text{ for PFOA and birth weight}$
Danish study		weight, SGA	
		(1100)	OR = 4.82 (0.56–41.16) and 2.44 (0.27–22.25) for LBW
		(n=1400)	and highest vs lowest quartiles of PFOS and PFOA,
			respectively
			OR = 1.43 (0.50–4.11) and 1.71 (0.55–5.28) for preterm
			birth and highest vs lowest quartiles of PFOS and PFOA,
			respectively
Stein 2009	PFOA	Preterm birth	For exposures >90 th percentile:
C8 project	PFOS	Low birth weight	Pre-term birth: $OR = 1.4 (1.1, 1.7)$ for PFOS and there
			was a monotonic exposure-response relationship and OR
		(n=1,845 pregnancies for	<1 for PFOA
		PFOA and 5,262 pregnancies	Low birth weight $OR = 1.8$ (1.2, 2.8) for PFOS and there
		for PFOS)	was a monotonic exposure-response relationship and OR
Fei 2008	PFOS and PFOA in maternal blood	Placental weight	<1 for PFOA Placental weight $β = -21.3 g (-46.1, 3.4)$ and $-10.8 g$
Danish study	samples taken early in pregnancy	Birth length	(-33.4, 11.8) for highest vs lowest quartiles of PFOA and
Damon study	samples taken earry in pregnancy	Head and abdominal	PFOS, respectively; monotonic exposure-response
		circumferences	relationship for PFOA
		Ponderal index	r
			Birth length β = -0.49 cm (-0.81, -0.16) for highest vs
		(n=1400)	lowest quartiles of PFOA

Reference Nolan 2009 Little Hocking communities	Exposure PFOA	Outcome Mean birthweight, mean gestational age, low birthweight, and preterm birth	Head circumference β = -0.14 cm (-0.39, 0.12) for highest vs lowest quartiles of PFOA Abdominal circumference β = -0.29 cm (-0.63, 0.06) for highest vs lowest quartiles of PFOA; monotonic exposure-response relationship for PFOA OR/ β & 95% CI β = -8.81 g (-86.1–68.5) for mean birth weight comparing LHWA only to no LHWA water service
Washino 2009 Japan	PFOS and PFOA in maternal serum	(n=1555) Birth weight Birth length	β = -269.4 g (-465.7 to -73.0 g) and -76.7 (-234.7 to 81.3) for PFOS and PFOA per log10 unit and birth weight
		Chest circumference Head circumference (n=428)	only in female infants
Andersen 2010 Danish study	maternal plasma levels of PFOS and PFOA and cord blood samples	Weight, length, and body mass index development during 1 st year of life (n=1400 born in 1996-2002)	Weight: $a\beta$ = -0.8 g (-4.2, 2.6) at 5 months and -5.8 g (-10.4,-1.2) at 12 months for PFOS; -9.4 g (-28.6, 9.9) at 5 months and -19.0 g (-44.9, 6.8) at 12 months for PFOA
Hamm 2010 Canada	PFOS PFOA PFHxS	Birth weight Fetal growth SGA	adjusted changes in birth weight per natural log (ng/ml) of PFOA were -37.4 g (-86.0 to 11.2 g)
		Preterm birth (n=252)	Difference of -0.086 (- 0.62 to 0.45) gestation week for highest vs lowest tertile of PFOA RR= 2.35 (0.63–8.72) for highest vs lowest tertile of PFHxS and SGA RR = 1.31 (0.38-4.45) and 1.11 (0.36–3.38) for preterm birth comparing highest vs lowest tertile of PFOA and PFOS, respectively; monotonic exposure-response relationship for PFOS

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Reference	Exposure	Outcome	OR/β & 95% CI
Chen 2012	Cord blood for PFOA, PFOS, PFNA,	Gestational age	per ln unit:
Taiwan	and PFUA	Birth weight	β = -0.37 (-0.60, -0.13) weeks for gestational age and -
		Birth length	0.17 cm
		Head circumference Ponderal	(-0.42, 0.09) for birth length and PFOS
		index (indicator of	
		disproportionate or	βs ranged from -19.2 to -110.2 g for birth weight and
		asymmetric growth restriction)	-0.05 to -0.25 cm for head circumference and PFOA,
		Preterm birth	PFOS, and PFUA; there was a monotonic exposure-
		Low birth weight SGA	response relationship for head circumference and PFOS when exposures were categorized into quartiles
		SGA	when exposures were categorized into quartiles
		(n=429)	βs ranged from -0.01 to -0.02 for Ponderal index and
			PFOA, PFOS, PFNA, and PFUA
			ORs of preterm birth and low birth weight = $2.45 (1.47,$
			4.08) and 2.61 (0.85, 8.03), respectively, for PFOS
			OD - 2.27 (1.25 4.15) - 1.1.24 (0.75 2.05) for CCA - 1.1
			ORs = 2.27 (1.25, 4.15) and 1.24 (0.75, 2.05) for SGA and PFOS and PFOA, respectively
			FFOS and FFOA, respectively
			An exposure response relationship was observed for PFOS
			and head circumference and preterm birth
Savitz 2012a	Serum PFOA levels at the time of	Preterm birth, term low	$OR \le 1$ for preterm birth and term low birthweight
C8 project	pregnancy from drinking water	birthweight	_
	contaminated by chemical plant		
	releases	(n= 11,737 pregnancies)	
	analysis uses modeled serum PFOA		
Maisonet 2012	estimates PFOS	Fetal and postnatal growth in	Birth weight: β ranged from -107.93 to -14.01 g for the
Avon Longitudinal Study	PFOA	girls	highest vs lowest tertiles of PFOS, PFOA, and PFHxS,
(Britain)	PFHxS	Sur	respectively; exposure-response relationships were
		(n=447)	observed for all 3 chemicals
		()	
			Birth length: β ranged from -0.44 to -0.82 cm for the
			highest vs lowest tertiles of PFOS, PFOA, and PFHxS;

			exposure-response relationships were observed for PFOA and PFHxS Gestational age: β ranged from -0.15 to -0.24 weeks for the highest vs lowest tertiles of PFOS, PFOA, and PFHxS, respectively; exposure-response relationships were observed for all 3 chemicals At 20 months, girls usually weighed more for the highest vs lowest tertiles of the chemicals
Reference	Exposure	Outcome	OR/β & 95% CI
Whitworth 2012b Norway study	Maternal plasma samples of PFOS and PFOA obtained around 17 weeks of gestation	Birth weight z scores, preterm birth, SGA and large for gestational age (LGA)	β = -0.18 (-0.41, 0.05) and -0.21 (-0.45, 0.04) for adjusted birth weight z scores and highest vs lowest quartiles of PFOS and PFOA, respectively
		(n=901)	OR = 1.3 (0.5, 3.4 0.51) for SGA and highest vs lowest quartiles of PFOS
Bach 2015b meta-analysis	PFOA or PFOS in maternal blood during pregnancy or umbilical cord	Birth weight	PFOA exposure was associated with decreased measures of continuous birth weight in all 14 studies PFOS exposure and birth weight were associated in some studies
Wu 2012	PFOA	Gestational age	Adjusted results for 1 lg-unit change in PFOA:
China		Birth weight	Gestational age: -15.99 days (-27.72 to -4.25)
		Birth length	Birth weight: -267.30 g (-573.27 to -37.18)
		Apgar scores	Birth length: -1.91 cm (-3.31 to -0.52)
		(n=167)	5-minute Apgar score: -1.37 (-2.42 to -0.32)
Darrow 2013	PFOA and PFOS measurements in	Preterm birth (n=158), low	β = -54 g (-124, 17) for birth weight in full-term infants
C8 project	2005-2006	birth weight (n=88), and birth	and the highest vs lowest quintile of PFOS
		weight among full-term infants	$(\beta = -105 \text{ g} [-196, -13] \text{ when restricted to births}$
		(, , , , , , , , , , , , , , , , , , ,	conceived after the blood sample collection
		(n=1630)	OD 122 (0.52, 2.22) for most one blade and blade
			OR = 1.32 (0.53, 3.32) for preterm birth and highest vs lowest quintile of PFOA restricted to births conceived
			after the blood sample collection
			OR = 1.33 (0.60, 2.96) for LBW and the highest vs lowest quintile for PFOS

Reference	Exposure	Outcome	OR/β & 95% CI
Kishi 2015	prenatal PFOS and PFOA levels were	Birthweight	β = - 186.6 g (-363.4, -9.8) for females comparing the 4 th
Hokkaido Study	measured in maternal serum samples	(205 1 111 1)	and 1st quartiles of PFOS
Bach 2015b	samue lavale of DELLEC DELLEC DEOC	(n= 306 mother-child pairs)	βs for birthweight ranged from -8 to -23 g for the highest
Danish study	serum levels of PFHxS, PFHpS, PFOS, PFOA,	Birth weight Birth length	vs lowest quartiles of PFHxS, PFHpS, PFOS, and PFUnA
Damsh study	PFNA, PFDA, and PFUnA	Head circumference	vs towest quartiles of 1111x5, 1111p5, 11 O5, and 11 OhA
	measured between 9-20 completed	Gestational age at birth	βs for birthweight for girls ranged from -39 to -76 g for
	gestational weeks	Preterm birth	the highest vs lowest quartiles of PFHxS, PFHpS, PFOS,
		(n=1507 mother-child pairs)	PFNA, PFDA, and PFUnA
			Association between PFAA exposures and birth length,
			head circumference, and gestational age were all close to
			zero
			OR = 1.18 (0.65, 2.16) for preterm birth for the highest vs
			lowest quartile of PFNA
Lenters 2015	PFHpA, PFHxS, PFOS, PFOA, PFNA,	Term birth weight (n=1250)	ln–PFOA β = -63.77 g (-122.83, -4.71); represents change
Greenland, Poland and	PFDA, PFUnDA, and PFDoDA		per a 2-SD increase in ln–transformed exposure biomarker
Ukraine	DECA 1 DECG	D' d d d	or untransformed continuous covariate levels
Verner 2015 Meta-analysis	PFOA and PFOS	Birth weight	summary β coefficients for g birth weight per ng/ml increase in PFOA and PFOS levels were -14.7 g (-21.7, -
Wicta-analysis			7.8) and -5.0 g (-8.9, -1.1), respectively
Lee 2016	PFBS	Birthweight in (n=85 births)	InPFOS $\beta = -0.14$ (95% CI: -0.33, 0.03)
South Korea	PFHxS		lnPFOA $\beta = -0.03$ (95% CI: -0.25 , 0.18)
	PFHpS		InPFNA $\beta = -0.14 (95\% \text{ CI:} -0.39, 0.10)$
	PFOS PFOA		InPFDA $\beta = -0.12$ (95% CI: -0.39 , 0.14)
	PFNA		lnPFDoA $\beta = -0.03 (95\% \text{ CI: } -0.36, 0.30)$
	PFDA		
	PFUnA		
	PFDoA		

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Reference	Exposure	Outcome	OR/β & 95% CI
Callan 2016	PFOS	Birth weight	OR = 3.5 (1.1-11.5) for being 95% of their calculated
Western Australia	PFOA	Birth length	optimal birth weight comparing the highest to lowest
	PFHxS	Head circumference	tertile of PFHxS
	And 11 other PFAS measured in whole		
	blood	(n=98 pregnant women)	β = - 69 g (-231, 94), -48 g (-203, 108) and -103 g (-221, 15) for birthweight and ln-unit increase in PFOS, PFOA,
	Median (in μg/L):	Proportion of optimal birth	and PFHxS
	PFOS 1.99	weight (POBW), proportion of	
	PFHxS 0.33	optimal birth length (POBL)	
	PFOA 0.86	and proportion of optimal head	
		circumference	
		(POHC)	
		(n=82-89 infants)	
Lauritzen 2017	PFOS and PFOA	Birth weight, birth length,	Sweden:
		head circumference,	PFOA:
	Median serum levels (ng/ml):	gestational age, SGA	Birthweight β = -359 g (-596, -122)
	PFOA: 2.33 in Sweden and 1.62 In		Birth length β = -1.3 (-2.3, -0.3)
	Norway	424 mother-child pairs,	Head circumference β = -0.4 (-1.0, 0.1)
	PFOS: 16.4 in Sweden and 9.74 in	excluding 1st time mothers:	Gestational age β = -0.3 (-0.9, 0.3)
	Norway	143 SGA births and 281 non-	SGA OR=5.25 (1.68-16.4)
		SGA controls	
			Differences more pronounced in boys
			PFOS:
			Birthweight β = -292 (-500, -84)
			Birth length β = -1.2 (-2.1, -0.3)
			Head circumference β = -0.4 (-0.9, 0.04)
			Gestational age β = -0.4 (-0.9, 0.2) 0.201
			SGA OR=2.51 (0.93-6.77)
			5011 011-2.51 (0.75-0.77)
			In Norway, SGA ORs < 1 and β s > 1 or very close to 0

 $\label{thm:condition} \textbf{Table A7. PFAS studies on congenital malformations.}$

Reference	Exposure	Outcome	OR/β/RR & 95% CI
Nolan 2010	PFOA	Congenital anomaly	Serviced entirely by contaminated LHWA water:
Little Hocking communities		(n=168 served by LHWA only and 1171 no LHWA)	OR = 7.0 (0.4-113) for both heart and circulatory defect $OR = 21 (0.9-517)$ for club foot
Savitz 2012a C8 project	Serum PFOA levels at the time of pregnancy from drinking water contaminated by chemical plant releases	Birth defects (n= 11,737 pregnancies)	$OR \le 1$ for birth defects
	analysis uses modeled serum PFOA estimates		
Stein 2014a C8 project	Modeled PFOA	Maternally reported birth defects (n = 325) among 10,262 births	Brain defect $OR = 2.6$ (1.3-5.1) for IQR increase from 25th to 75^{th} percentile and $OR = 16.1$ (0.8, 325) for highest vs lowest tertile Limb defect $ORs = 1.2$ (0.7, 2.0) for IQR increase and 1.5 (0.2, 9.7) for highest vs lowest tertile Eye defect $OR = 1.3$ (0.2-8.4) for highest vs lowest tertile Heart defect $ORs = 1.2$ (0.8, 1.7) for IQR increase and 1.4 (0.4, 5.1) for highest vs lowest tertile
Vesterholm Jensen 2014	Cord blood PFAS levels	Cryptorchidism (n=29 Danish cases and 30 matched controls and 78 Finnish cases and 78 matched controls)	OR = 1.14 (0.19–6.95) and 1.30 (0.27–6.39) for ln PFOA and PFOS in Danish cases, respectively OR = 2.34 (0.16–34.67) for the highest vs lowest tertile of PFOS in Denmark
Toft 2016 Danish study	Amniotic fluid PFOS level	Cryptorchidism Hypospadias (n= 270 cryptorchidism cases, 75 hypospadias cases, and 300 controls)	ORs for cryptorchidism and hypospadias were < 1
Kim 2016 South Korea	16 PFAS in infant sera Mean concentrations were: PFOS 4.05ng/mL), PFOA	Congenital hypothyroidism (CH) measured by serum thyroid stimulating hormone (TSH),free	large difference in PFOA concentrations between cases and controls (2.12 ng/mL in controls and 5.40 ng/mL in cases)

	(2.12ng/mL), PFHxS 1.17ng/mL)	thyroxine (FT4), total T3, thyroglobulin antibody (TGAb), relevant microsomal antibodies (microAb), and thyroid stimulating immuno-globulin (TSI) (n=27 infants with CH and 13 healthy infants)	mean concentrations of PFOA, PFNA, PFDA, PFUnDA, and total PFAS in cases (0.525–16.8ng/mL) were "significantly" higher than in controls (0.298–10.0ng/mL) (data presented in figure only) in CH infants, correlations were -0.482 and -0.642 for TSI and PFOA and PFHxS, respectively (results not shown for PFOS)
Reference	Exposure	Outcome	OR/β/RR & 95% CI
Liew 2014 Danish study	PFASs in maternal plasma collected in early or midpregnancy: PFOS, PFOA, PFHxS, PFNA, PFHpS, PFDA Median (ng/ml) PFOS: 27.40 PFOA: 4.00 PFHxS: 0.92	Cerebral palsy (n=156 cases and 550 controls)	per 1-unit (natural-log ng/mL) increase in boys: RR = 1.7 (1.0, 2.8) for PFOS RR = 2.1 (1.2, 3.6) for PFOA RR = 1.2 (0.9, 1.7) for PFHxS RR = 1.2 (0.6, 2.5) for PFNA RR = 1.5 (1.0, 2.2) for PFHpS RR = 1.1 (0.7, 1.7) for PFDA and there was an exposure response relationship for PFHxS, PFNA, and PFHpS when exposure was categorized RRs generally increased for boys born at term

Table A8. PFAS studies on adverse health outcomes in children ages ≥ 2 years.

Reference,	Studies on adverse nearth Study population	PFOS	PFHxS	PFOA	Outcome	Difference detected (95% CI)
Location	Study population	serum	serum	serum	Outcome	Enterence detected (50 % O1)
		level	level	level		
Pease Tradeport	children <12 years, N=366	8.3	4.2	3.6		
r ouse rruncpore	children <18 years, N=396	8.1	4.0	3.4		
Frisbee 2010	1,971 boys <12 years	19.9	n/a	35.1		< 12 years 12 - < 18 yrs
C8 study	2,773 boys 12 - <18	20.3		30.1	Total cholesterol	PFOA: $+6.3^{\$}$ $\beta=1.6 (0.4) +4.8^{\$}$ $\beta=1.1 (0.4)$
· ·					(mg/dL)	PFOS: $+6.2^{\$} \beta = 1.2 (0.5) +9.3^{\$} \beta = 2.1 (0.4)$
	1,886 girls <12 years	21.7	n/a	30.7	Total cholesterol	PFOA: $+5.8^{\$}$ $\beta=1.1 (0.4) +3.9^{\$}$ $\beta=1.0 (0.4)$
	2,520 girls 12 - <18	18.2		22.9	(mg/dL)	PFOS: $+4.6^{\P}$ $\beta=1.3 (0.5) +9.4^{\P}$ $\beta=1.9 (0.4)$
					≥170 mg/dL	OR=1.6 (1.4, 1.9), PFOS (5th quintile)
						OR=1.2 (1.1, 1.4) PFOA (5th quintile)
Zeng 2015	102 boys, age 12-15	29.9	1.4	0.5	Total cholesterol (all	4th quartile: 23.1 mg/dL increase for PFOS,
Taiwan	123 girls, age 12-15	28.8	1.2	0.5	children)	12 mg/dL increase for PFOA.
	225 total age 12-15	28.9	1.3	0.5		Ln PFHxS β = 1.1 (-0.7, 2.9)
						Ln PFNA $\beta = 12.9 (0.7, 25.1)$
						Ln PFOS $\beta = 0.3 (0.2, 0.5)$
						Ln PFOA $\beta = 6.6 (2.7, 10.4)$
Maisonet 2015a	Maternal serum	20.0		3.6	Total cholesterol	3 rd tertile β for PFOS & PFOA were <0.
Avon, UK	(N=199 girls aged 7 and 15)					Mean differences at age 15, 3 rd tertile vs 1 st tertile, for
G : 2011	0.15 1.11 1.10 1.0	45.5		1.0	m . 1 . 1 1	PFOA and PFOS = 8.1 and 19.1, respectively.
Geiger 2014a	815 children, aged 12-18	17.7		4.2	Total cholesterol	PFOS: 5.9 mg/dL increase (0.1, 11.6), 3 rd tertile
NHANES	1999-2008	(mean)		(mean)	III -1 -1 -1 -1 -1 -1	PFOA: 7.0 mg/dL increase (1.4, 12.6), 3 rd tertile
					High cholesterol	PFOS: OR=1.53 (1.07, 2.19), 3 rd tertile
Nelson 2010	222 hours 12 10 (2002 2004)	19.9	2.4	4.0		PFOA: OR=1.49 (1.05, 2.12) 3 rd tertile 4 th quartile mean difference (vs 1 st quartile), (mg/dL)
NHANES	322 boys, 12-19 (2003-2004) 263 girls, 12-19 (2003-2004)	19.9	2.4	4.0		Boys Girls
MIMILS	203 girls, 12-19 (2003-2004)				Total cholesterol	PFOS: 3.6 (-8.5, 15.7) -0.4 (-9.3, 8.6)
					Total choicsteroi	PFOA: 5.0 (-2.3, 12.2) 3.3 (-4.2, 10.8)
						PFHxS: -3.2 (-15.4, 9.0) -12.7 (-23.4, -2.0)
						1111AD. 3.2 (13.4, 7.0) 12.7 (23.4, 2.0)
					Non-HDL cholesterol	PFOS: 2.0 (-8.1, 12.1) -4.1 (-12.9, 4.7)
					2	PFOA: 6.8 (-1.3, 14.9) -1.0 (-9.1, 7.2)
						PFHxS: -3.3 (-14.6, 8.1) -16.5 (-29.5, -3.4)
Lopez-Espinosa 2012	1,078 children, ages 1–5 years	16.3		33.8	thyroid stimulating	PFOS, 4th vs 1st quartile: 3.1% (0.0, 6.2) increase in
C8 study	3,132 children ages 6–10 years	21.8		32.2	hormone, TT ₄	TSH, 2.3% (1.2, 3.3) increase in TT ₄
•	6,447 ages >10–17 years	19.6		26.9	<u> </u>	
	1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1				Thyroid disease	PFOA: OR=1.44 (1.02, 2.03)
					_	PFOS: OR=0.80 (0.62, 1.08)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Lin 2013	212 aged 12-19	7.0		2.8	Free T ₄	5% increase in free T ₄ for PFNA(same level as
Taiwan						Pease) [¥]
Qin 2016					Uric acid ≥6 mg/dL;	PFOA PFOS PFHxS
Taiwan	102 boys, aged 12-15	29.9	1.4	0.5	14.7% prevalence	2.8 (1.4, 5.6) 1.4 (0.9, 2.2) 1.65 (1.01, 2.69)
	123 girls, aged 12-15	28.8	1.2	0.5	(Odds ratios)	1.6 (0.7, 3.9) 1.5 (0.8, 2.9) 1.3 (0.7, 2.3)
	225 total, aged 12-15	28.9	1.3	0.5		2.2 (1.3, 3.6) 1.35 (0.95, 1.93) 1.4 (0.9, 2.1)
Geiger 2013	1,772 aged 12-18	16.6		4.1	Serum uric acid (4th	PFOA: .30 mg/dL increase
NHANES	1999-2008 data				quartile)	PFOS: .12 mg/dL increase
					Hyperuricemia (16%)	PFOA: OR=1.62 (4q vs 1q)
						PFOS: OR=1.65 (4q vs 1q)
Kataria 2015	1,960 aged 12-18	12.8	2.0	3.5	Serum uric acid (4th	PFOA: .21 mg/dL increase (0.06, 0.37)
NHANES	2003-2010				quartile	PFOS: .19 mg/dL increase (0.03, 0.34)
						PFHxS: .05 mg/dL decrease (-9.22, 0.11)
					eGFR (4 th quintile)	PFOA: -6.61 (-11.39, -1.83)
					$(mL/min/1.73 m^2)$	PFOS: -9.47 (-14.68, -4.25)
					(1112/11111/11117)	PFHxS: -0.32 (-4.44, 3.81)
Lopez-Espinosa 2016					Percent difference	PFOS PFOA PFHxS
C8	1,169 boys aged 6-9 years	22.4	8.1	34.8	Ln testosterone	-5.8 (-9.4, -2.0) -4.9 (-8.7, -0.8) -2.7 (-6.4, 1.2)
					Ln estradiol	-4.0 (-7.7, -0.1) 4.3 (-0.4, 9.1) -1.3 (-5.5, 3.1)
					Ln IGF-1	-5.9 (-8.3, -3.3) -0.4 (-3.4, 2.7) -2.5 (-5.2, 0.3)
	1,123 girls aged 6-9 years	20.9	7.0	30.1	Ln testosterone	-6.6 (-10.1, -2.8) -2.5 (-6.7, 1.8) 0.2 (-3.5, 4.0)
					Ln estradiol	-0.3 (-4.6, 4.2) 4.2 (-0.7, 9.4) 2.1 (-2.2, 6.5)
					Ln IGF-1	-5.6 (-8.2, -2.9) -3.6 (-6.6, -0.5) -2.1 (-4.8, 0.7)
Tsai 2015	95 children aged 12-17	7.12		3.03	Ln SHBG	PFOA: decline among girls
Taiwan					Ln FSH	PFOS: declines both sexes
					Ln testosterone	PFOS: decline among girls
Maisonet 2015b	72 girls, 15 years of age $^{\Delta}$	19.2	1.6	3.6	Total testosterone	Increase in total testosterone by about .20 nmol/L for
Avon, UK						PFOS, PFOA, and PFHxS (95% CI: .01, .38)
					SHBG	Declines for PFOA and PFHxS in 3rd tertile but
					Silbo	increases in 2nd tertile
Zhou 2016	102 boys, aged 12-15	29.9	1.4	0.5	Ln testosterone	Declines in both sexes for PFOA; decline in boys,
Taiwan						PFOS
	123 girls, aged 12-15	28.8	1.2	0.5	Ln estradiol	Increase in both sexes, PFOA
Christensen 2011	218 girls (puberty <11.5 yrs)	19.8	1.6	3.7	Early age at puberty	PFOA: OR=1.29 (0.86, 1.93)
Avon, UK	230 controls (aged 13 yrs) $^{\Delta}$					PFOS: OR=0.83 (0.56, 1.23)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Lopez-Espinosa 2011 C8	3,072 boys, ages 8-18	20		26	Reaching puberty: Odds ratio # days delay	4 th quartile PFOS PFOA 0.46 (0.29, 0.71) 0.75 (0.49, 1.15) 190 days delay 69 day delay
	2,903 girls, ages 8-18	18		20	Odds ratio # days delay	0.55 (0.35, 0.87) 0.57 (0.37, 0.89) 138 days delay 130 days delay
Kristensen 2013 Denmark	343 women, 20 years old $^{\Delta}$	21.1		3.6	Reaching puberty, Months delay	3rd tertile: PFOS, 1.5 (-2.5, 5.4) months delay; PFOA, 5.3 (1.3, 9.3) months delay
Wang 2015 Taiwan	120 at age 5^{Δ} §	13.25	0.69	2.5	VIQ, PIQ, FSIQ®	Age 5: VIQ PIQ FSIQ PFOS:-1.7 (-4.0, 0.7) -2.2 (-4.7, 0.3) -1.9 (-4.3, 0.5) PFOA: 0.9 (-1.4, 3.3) 1.0 (-1.4, 3.4) 1.2 (-1.0, 3.5) PFNA: 0.7 (-1.3, 2.7) -1.4 (-3.4, 0.6) -0.2 (-2.1, 1.7)
	120 at age $8^{\Delta \epsilon}$	12.28	0.69	2.5		Age 8: VIQ PIQ FSIQ PFOS: -1.3 (-3.6, 1.1) -1.6 (-4.0, 0.7) -1.9 (-4.3, 0.4) PFOA: 0.5 (-1.5, 2.5) -1.1 (-3.2, 1.0) -0.4 (-2.5, 1.7) PFNA:-2.1 (-3.9, -0.2) -1.5 (-3.5, 0.4) -1.5 (-3.4, 0.4)
Stein 2013 C8	320 children, 6-12 years			35	IQ, reading, language, memory, (etc.)	PFOA evaluated. 4 th quartile PFOA had higher IQ scores than 1 st quartile and decreased scores for ADHD characteristics.
Lien 2016 Taiwan	282 children, 7 years old Cord blood levels	4.79		1.55	Hyperactivity symptoms	Slight, inconsistent results for PFOS and PFOA.
Stein 2011 C8	10,546 children ages 5-18	20.2	5.2	28.2	ADHD Learning problem	4th quartile, ORs: PFOS, 1.3 (1.0, 1.6); PFHxS, 1.6 (1.2, 2.1); PFOA, 0.7 (0.6, 0.9); PFNA, 1.2 (0.9, 1.5) 4th quartile, ORs: PFHxS, 1.2 (1.0, 1.4); PFOS, 0.9 (0.7, 1.0); PFOA, 0.9 (0.8, 1.1); PFNA, 0.7 (0.6, 0.9)
Stein 2014b C8	320 children, 6-12 years			35	ADHD behaviors	Inconsistent results (parents vs teachers; boys vs girls)
Fei 2011 Denmark	787 children, 7 years old [∆]	34.4		5.4	Hyperactivity $^{\Psi}$ Coordination $^{\Psi}$	Conduct problem: PFOS OR=1.45 (0.77, 2.72); PFOA OR=1.29 (0.67, 2.52) Coordination problem: PFOS OR=1.39 (0.65, 3.00); PFOA OR=1.14 (0.46, 2.81)
Hoffman 2010 NHANES, 1999- 2000, 2003-2004	571 children aged 12-15	22.6	2.2	4.4	ADHD, ORs for IQR	PFOS: OR=1.60 (1.10, 2.31) PFOA: OR=1.35 (1.04, 1.77) PFHxS: OR=1.19 (1.05, 1.34) PFNA: OR=1.15 (0.93, 1.42)
Ode 2014 Sweden	203 ADHD cases and 205 controls (cord blood PFASs)	6.8		1.8	ADHD	PFOA, ≥75th percentile: OR=1.07 (0.67, 1.70) (PFOS OR < 1.0)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Liew 2015 Denmark 545 controls [△]	215 ADHD [∆]	26.8	0.8	4.1	ADHD	PFOA, 4th quartile: OR=1.14 (0.92, 1.40). OR=2.0 (1.5, 2.8) when all six PFAS included in model; PFOS & PFHxS ORs <1.0. PFNA OR < 1.0, but when all 6 PFAS in model, the OR for PFNA=1.6 (1.2, 2.1)
Serum levels: PFOS=27.4 PFHxS=0.9 PFOA=4.0	213 ASD ^Δ	25.4	0.9	3.9	ASD	PFHxS, 4th quartile: OR=1.07 (0.73, 1.56). When all 6 PFAS in model, OR=1.3 (0.8, 2.1). Per ln(PFHxS), OR=1.10 (0.92, 1.33). OR<1.0 for PFOA, PFOS, PFNA. When all 6 PFAS in model, PFOS OR=1.2 (0.7, 2.1)
Braun 2014 Cincinnati, OH	175 children tested at age 4 and/or age 5^{Δ}	13.0	1.6	5.5	SRS	PFOS: 1.6 (-0.8, 4.1) increase in SRS score per 2-SD increase. PFHxS, 1.0 (-1.2, 3.3) increase; (decreased SRS score for PFOA)
Strøm 2014 Denmark	876 adolescents [∆]	21.4		3.4	ADHD (3.1%) Depression (11.9%)	HRs < 1.0 3rd tertile: PFOS HR=1.16 (0.69, 1.95); PFOA HR=1.03 (0.61, 1.73)
					Scholastic achievement	Slight decrements for PFOS and PFOA
Chen 2013 Taiwan	239 children aged 2 years Cord blood PFASs	7.4		2.6	Developmental delay	PFOS associated with deficits in development scores, especially for motor development: gross-motor domain, IQR= -3.7 points (-6.0, -1.5); OR for poor performance= 2.4 (1.3, 4.2). (Slight deficits to null findings for PFOA)
Forns 2015 Norway	843 toddlers Breast milk PFASs	0.11		0.04	Developmental delay	PFOA, >median: OR=1.25 (0.81, 1.95); PFOS, OR<1.0
Gump 2011 Oswego, NY	83 children, aged 9-11	8.79	3.67	3.28	Response inhibition	All PFASs measured reduced inhibition
Vuong 2016 Cincinnati, OH	256 mother-child pairs (maternal serum measured, 2 nd trimester) Children aged 5 and 8 years	12.6	1.4	5.3	Executive function: Behavioral regulation Metacognition index Global Executive composite	3rd tertile, ORs (clinical relevance): PFOA PFOS PFHxS 1.36 (0.55, 3.35) 2.45 (0.91, 6.56) 2.03 (0.80, 5.18) 1.06 (0.43, 2.60) 2.17 (0.85, 5.51) 1.53 (0.63, 3.74) 1.25 (0.51, 3.08) 2.42 (0.92, 6.35) 2.31 (0.91, 5.88)
Dong 2013 Taiwan	225 children, 12-15, w/asthma 231 children, 12-15, controls	33.9	2.5	0.5	Asthma	4 th quartile, PFOA: OR=4.05 (2.21, 7.42) 4 th quartile, PFHxS: OR=3.83 (2.11, 6.93) 4 th quartile, PFOS: OR=2.63 (1.48, 4.69) 4 th quartile, PFNA: OR=2.56 (1.41, 4.65)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected (95% CI)
Stein 2016a NHANES	1,191 children aged 12-19, 1999-2000, 2003-2004	20.8	2.47	4.13	MMR antibody	PFOS: among seropositives, a 13.3% decrease in rubella antibody (-19.9, -6.2), and a 5.9% decrease in mumps antibody (-9.9, -1.6). (declines also for PFOA and PFHxS for rubella, & PFOA for mumps)
	640 children (2005-2006)	15.0	2.09	3.59	Allergic conditions	PFOA : OR=1.28 (0.81, 2.04) for asthma (similar findings for PFOS and PFNA); OR=1.35 (1.10, 1.66) for rhinitis. For rhinitis, PFOS OR=1.16 (0.90, 1.50) and PFNA OR=1.24 (0.97, 1.60).
						PFOS & PFOA sensitivity to mold; PFOA sensitivity to rodents
Humblet 2014 NHANES	1,877 aged 12-19 years (1999-2008)				Current asthma Wheeze	PFOA, 3 rd tertile, OR=1.18 (0.90, 1.53). (ORs for PFOA, PFNA and PFHxS < 1.1.) For wheeze, all ORs for the PFAS chemicals were <1.1)
Goudarzi 2016 Japan	1,558 mother-child (aged 4 year) pairs. Maternal serum PFAS at 3 rd trimester	4.9	0.28	2.1	Eczema, wheezing, rhinoconjunctivitis	For total allergic diseases, 4 th quartile ORs for PFOA, PFOS, PFHxS and PFNA < 1.00. For wheezing, PFOA OR=1.09 (0.73, 1.65) (elevation in boys only); for PFNA, OR=1.11 (0.76, 1.63) (elevation in boys only). For PFOS and PFHxS, ORs < 1.00
Granum 2013 Norway	99 prenatal blood samples; 50 children aged 3 years	5.5	0.3	1.1	Rubella	PFOA & PFHxS: β = -0.4 optical density (-0.64, -0.11) PFOS: β = -0.08 optical density (-0.14, -0.02) PFNA: β = -1.26 optical density (-2.32, -0.20) (regression coefficients were also negative for measles but confidence intervals were wide)
					Gastroenteritis Common cold	PFOA & PFHxS: ORs > 3.0 (CIs were very wide) PFHxS: OR=1.71 (0.20, 14.8)
Buser 2016 NHANES	Children aged 12-19, 2005-2006, 2007-2010				Food allergies, sensitization (IgE)	4 th quartile ORs Self-reported allergies food sensitization PFOA: 9.09 (3.32, 24.9) 1.23 (0.57, 2.65) PFOS: 2.95 (1.21, 7.24) 0.74 (0.23, 2.40) PFHxS: 3.06 (1.35, 6.93) 1.17 (0.56, 2.44) PFNA: 1.73 (0.54, 5.52) 0.51 (0.28, 0.92)

Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected
Wang 2011 Taiwan	244 children, aged 2 years Cord blood	5.5	0.04	1.71	Atopic dermatitis	PFOS, 4th quartile: OR=2.19 (0.78, 6.17). OR for PFOA and PFNA < 1.00
Dalsager 2016 Denmark	346 children aged 1-3 years. Maternal serum PFAS <16 weeks gestation	8.07	0.32	1.68	Fever & cough	3 rd tertile ORs (above median proportion of days) PFOS: 2.35 (1.34, 4.11) PFOA: 1.97 (1.07, 3.62) PFHxS: 1.29 (0.72, 2.28) PFNA: 1.49 (0.86, 2.59) 3 rd tertile RRs (# days with fever) PFOS: 1.65 (1.24, 2.18) PFOA: 1.12 (0.82, 1.54) PFHxS: 1.20 (0.89, 1.62) PFNA: 1.12 (0.84, 1.49) 3 rd tertile RR (fever & cough, # of episodes) PFOS: 1.33 (0.99, 1.80) PFOA: 1.11 (0.80, 1.56) PFHxS: 1.13 (0.82, 1.55) PFNA: 1.02 (0.76, 1.38) 3 rd tertile RR (# episodes of diarrhea) PFHxS: 1.71 (0.92, 3.16) PFOS: 1.19 (0.67, 2.12) PFOA: 1.08 (0.55, 2.13)
Grandjean 2012 Faroes	532 children aged 5 years	16.7	0.63	4.06	Inadequate antibody (<0.1 IU/mL), age 7)	ORs Tetanus (age 5) Diphtheria (age 5) PFOA: 4.2 (1.5, 11.4) 3.3 (1.4, 7.5) PFOS: 2.6 (0.8, 8.9) 2.4 (0.9, 6.4) PFHxS: 1.8 (1.1, 2.9) 1.5 (1.0, 2.3) PFNA: 1.6 (0.7, 3.6) 1.8 (1.0, 3.4)
Grandjean 2016 Faroes	515 children aged 13 years	6.7	0.4	2.0	Diphtheria antibody	% change per doubling of age 7 serum PFAS PFOS: -31.1 (-49.8, -5.4) PFOA: -9.4 (-31.1, 19.2) PFHxS: -19.5 (-34.7, -0.7) PFNA: -17.4 (-33.7, 2.8) (Tetanus had increased % change for PFAS)
Geiger 2014b NHANES	1,655 children, 1999-2000, 2003- 2008 (aged 12-18)				Hypertension	4 th quartile OR: PFOS: 0.69 (0.41, 1.17) PFOA: 0.77 (0.37, 1.61)

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Reference, Location	Study population	PFOS serum level	PFHxS serum level	PFOA serum level	Outcome	Difference detected
Domazet 2016 Denmark	590 aged 9 years 444 aged 15 years 369 aged 21 years	42 21.5 10.5		9.3 3.5 2.9	adiposity	PFOS serum levels at age 9 was associated with indicators of adiposity in adolescence and young adulthood. PFOA serum levels at age 9 was associated with decreased β-cell function in adolescence. Later exposures were not associated with indicators of adiposity or glucose metabolism.
Mora 2016 MA	1,006 children in early childhood (3-6 years), 876 mid-childhood (6-11)	24.7	2.3	5.6	BMI, skinfold thickness, total fat mass index, waist circumference	Among girls, each interquartile increment of prenatal PFOA was associated with 0.21 kg/m ² (-0.05, 0.48) higher BMI, 0.76 mm (-0.17, 1.70) higher sum of subscapular and triceps skinfold thickness, and 0.17 kg/m ² higher total fat mass index. Similar findings for PFOS and PFHxS. No associations with boys.
Karlsen 2016 Faroes	444 children at 18 months 371 children at 5 years Maternal 2-week postpartum serum	4.68 8.04	0.34 0.19	2.22 1.37	Overweight	3 rd tertile ORs age 18 months age 5 years PFOS: 1.24 (0.98, 1.57) 0.94 (0.53, 1.66) PFOA: 1.10 (0.84, 1.46) 1.88 (1.05, 3.35) PFHxS: 1.24 (0.97, 1.58) 1.22 (0.73, 2.04)

PFASs serum levels are in micrograms per liter (µg/L)

SHBG: sex hormone-binding globulin. (Note: Testosterone circulating in the bloodstream is mostly bound to SHBG. Endocrine disruptors may bind to SHBG displacing reproductive hormones and affecting their bioavailability.)

FSH: follicle stimulating hormone

Note: For the Nelson 2010 study, a monotonic dose-response was observed only for total cholesterol and PFOS among males 12-19. The average differences in the table are based on the highest difference observed regardless of quartile observed (i.e., the largest difference could appear in quartiles 2, 3 or 4). For PFHxS, the highest quartile was negative for total cholesterol but there was considerable inconsistency between quartiles.

Note: For the Fei 2011 study, total scores for hyperactivity/behavior problem

^{¶ 5}th vs 1st quintile

[¥] PFNA=0.91 μg/L geometric in the Taiwan study, and 0.92 μg/L among Pease Tradeport children.

[△] The PFASs levels are for the mothers of this study population during their pregnancies.

^{€ 89} of the children were tested at both ages ("paired children").

 $[\]Psi$ high score on a screening test for hyperactivity/conduct problems; low scores on developmental coordination screening test.

^ø change in IQ with a doubling of PFASs level (Wang 2015)