Appendix F – Summary of Immune Effects

The following table summarizes the findings of previous epidemiological studies of PFAS and immune-related effects – specifically hospitalization due to infectious diseases, risk of respiratory tract infections, and decreased vaccine response. These studies include occupational exposure studies, studies of communities living near a PFOA manufacturing facility with high levels of PFOA in the drinking water, and studies of populations exposed to background levels of perfluoroalkyls (referred to as general population studies). This summary reflects the relevant available literature as of 2/16/2021, including some studies that suggest PFAS exposure may impact the immune system and susceptibility to viral infections.

Compound	Acronym	CAS Registry Number
Perfluorooctanoic acid	PFOA	335-67-1
Perfluorononanoic acid	PFNA	375-95-1
Perfluorodecanoic acid	PFDA	335-76-2
Perfluoroundecanoic acid	PFUnA	2058-94-8
Perfluorododecanoic acid	PFDoDA	375-73-5
Perfluorohexane sulfonic acid	PFHxS	1763-23-1
Perfluorooctane sulfonic acid	PFOS	754-91-6
N-Methylperfluorooctane sulfonamidoacetic acid	MeFOSAA	2355-31-9

Study Overview	Serum PFAS Levels	Outcome	Results
Abraham et al. 2020	Mean child serum PFAS levels	Correlation between PFAS	Correlation between adjusted
	PFOA, formula-fed: 3.8 μg/L	levels and specific vaccine	antibody levels and PFOA
Cross-sectional study of 101	PFNA, formula-fed: 0.2 μg/L	antibodies for tetanus,	Hib: <i>r</i> = -0.32, <i>p</i> = 0.001
healthy 1-year old children (21	PFHxS, formula-fed: 1.7 μg/L	diptheria, and Haemophilus.	Tetanus: <i>r</i> = -0.25, <i>p</i> = 0.01
formula-fed and 80	PFOS, formula-fed: 6.8 μg/L		Diphtheria: <i>r</i> = -0.23, <i>p</i> = 0.02
breastfed)at the end of the			
1990s; specific vaccine	PFOA, breastfed: 16.8 μg/L		There were no significant correlations
antibodies for tetanus,	PFNA, breastfed: 0.6 μg/L		of levels of PFOS, PFHxS, and PFNA
diptheria, and Haemophilus.	PFHxS, breastfed: 2.1 μg/L		with levels of the vaccine antibodies
influence type b (Hib) were	PFOS, breastfed: 15.2 μg/L		

Study Overview	Serum PFAS Levels	Outcome	Results
measured			
Ait Bamai et al. 2020	Median maternal serum PFAS levels	Risk of infectious diseases in early life, based on mothers'	PFOA: OR (95% CI) Pneumonia: 1.17 (1.01, 1.37)
Prospective cohort study of	PFOA: 1.94 ng/mL	self-administered	
2,689 children enrolled in the	PFNA: 1.14 ng/mL	questionnaire:	<u>PFDoDA:</u> OR (95% CI)
Hokkaido Study from 2003-	PFDA: 0.51 ng/mL	Chickenpox	Chicken Pox: 0.85 (0.72, 1.00)
2012; children were monitored	PFUnA: 1.43 ng/mL	Otitis media	
for physician-diagnosed	PFDoDA: 0.17 ng/mL	Pneumonia	No other significant associations were
infectious disease up to 7 years	PFTrDA: 0.33 ng/mL	RSV	observed between maternal PFAS
of age.	PFHxS: 0.30 ng/mL		levels and any of the infectious
	PFOS: 5.12 ng/mL		diseases.
Bulka et al. 2021	Mean serum PFAS levels by	Presence of antibodies to	Prevalence Ratio (95% CI) per
	pathogen burden	cytomegalovirus, Epstein Barr,	doubling increase in serum PFAS, in
Cross-sectional study of 8778		hepatitis C and E, herpes	adults (20-49 years)
participants (3189 adolescents,	<u>Adolescents</u>	simplex 1 and 2, HIV, T. gondii,	PFOA
5589 adults) in NHANES 1999-	PFOA, uninfected: 2.16 ng/mL	and Toxacara spp.;	Herpes 1: 1.03 (1.01, 1.06)
2016; looked at presence of	PFOA, 3+ pathogens: 3.30ng/mL	Seropositivity was summed to	Herpes 2: 1.11 (1.05, 1.17)
antibodies to cytomegalovirus,	PFNA, uninfected: 0.72 ng/mL	calculate a pathogen burden	<u>PFNA</u>
Epstein Barr, hepatitis C and E,	PFNA, 3+ pathogens: 0.89ng/mL	score reflecting the total	Herpes 1: 1.05 (1.02, 1.08)
herpes simplex 1 and 2, HIV, T.	PFHxS, uninfected: 1.51 ng/mL	number of infections.	<i>Toxocara</i> spp.: 1.40 (1.13, 1.73)
gondii, and Toxacara spp.;	PFHxS, 3+ pathogens: 1.93ng/mL		<u>PFHxS</u>
Seropositivity was summed to	PFOS, uninfected: 5.58 ng/mL		<i>Toxocara</i> spp.: 1.21 (1.06, 1.37)
calculate a pathogen burden	PFOS, 3+ pathogens: 16.14ng/mL		PFOS
score reflecting the total			Herpes 1: 1.04 (1.01, 1.06)
number of infections.	Adults		<i>Toxocara</i> spp.: 1.57 (1.26, 1.96)
	PFOA, uninfected: 2.63 ng/mL		
	PFOA, 3+ pathogens: 2.75ng/mL		No significant associations between
	PFNA, uninfected: 0.82 ng/mL		persistent infections were found in
	PFNA, 3+ pathogens: 0.93ng/mL		adolescents.
	PFHxS, uninfected: 1.53 ng/mL		
	PFHxS, 3+ pathogens: 1.35ng/mL		Total Pathogen Burden, adults (20-49
	PFOS, uninfected: 8.14 ng/mL		<u>years); OR (95% CI)</u>
	PFOS, 3+ pathogens: 12.69ng/mL		PFOA: 1.09 (1.06, 1.12)

Study Overview	Serum PFAS Levels	Outcome	Results
			PFNA: 1.03 (1.00, 1.05)
			PFHxS: 1.02 (1.00, 1.05)
			PFOS: 1.10 (1.07, 1.12)
			Total Pathogen Burden, adolescents
			<u>(12-19 years); OR (95% CI)</u>
			PFOA: 1.36 (1.27, 1.45)
			PFNA: 1.11 (1.03, 1.19)
			PFHxS: 1.11 (1.06, 1.15)
			PFOS: 1.30 (1.25, 1.36)
Dalsager et al. 2016	Median maternal serum PFAS	The following outcomes were	Odds of number of days above
	level (measured at gestational	evaluated both as odds of	median, Fever:
Prospective cohort study of 359	age 10-16 weeks)	number of days above median	PFOA: OR 1.97 (1.07–3.62), 3 rd tertile
children (aged 1–4 years)	PFOA: 1.68 ng/mL	and number of days (incidence	PFOS: OR 2.35 (1.34–4.11), 3 rd tertile
participating in the Odense	PFOS: 8.07 ng/mL	rate):	
Child Cohort in Denmark	PFHxS: 0.32 ng/mL	Fever (>101.3°F)	Rate of number of days, Fever:
(women enrolled 2010-2012);	PFNA: 0.70 ng/mL	Cough	PFOS: IRR 1.65 (1.24–2.18), 3 rd tertile
parents responded to texts	PFDA: 0.27 ng/mL	Nasal Discharge	
every other week regarding the		Diarrhea	Odds of number of days above
child's symptoms of infection.		Vomiting	median, Nasal Discharge:
			PFNA: OR 0.53 (0.31–0.92), 2 nd tertile;
			OR 0.55 (0.31–0.97), 3 rd tertile
			No significant associations (p>0.05)
			between maternal serum levels of
			PFAS and any other outcomes.
Dalsager et al. 2021	Median maternal serum PFAS	Risk of hospitalizations for	Hazard ratios (95% CI) for
	level (measured at gestational	infectious disease categorized	hospitalization per doubling of
Prospective cohort study of	age 10-16 weeks)	as:	maternal PFAS concentrations
1,503 mother-child pairs (aged	PFOA: 1.68 ng/mL	Upper respiratory tract	
1–5 years) participating in the	PFOS: 7.52 ng/mL	infections (URTI)	Any infection
Odense Child Cohort in	PFHxS: 0.36 ng/mL	Lower respiratory tract	PFOS: 1.23 (1.05, 1.44)
Denmark (women enrolled	PFNA: 0.64 ng/mL	infections (LRTI)	

Serum PFAS Levels	Outcome	Results
PFDA: 0.29 ng/mL	Gastrointestinal infections (GI)	LRTI
	Other infections	PFOA: 1.27 (1.01, 1.59)
		PFOS: 1.54 (1.11, 2.15)
		No significant associations (p>0.05)
		between maternal serum levels of
		PFAS and any other outcomes.
Mean maternal serum PFAS level	Rate of hospitalization for	PFOA:
(measured at gestation week 12)	infectious disease in young	IRR 0.96 (0.87-1.06) for trend
PFOA: 5.6 ng/mL	children	IRR 1.21 (1.04-1.42) for trend, girls
PFOS: 35.3 ng/mL		IRR 0.83 (0.73-0.95) for trend, boys
		PFOS:
		IRR 1.00 (0.91-1.09) for trend
		IRR 1.18 (1.03-1.36) for trend, girls
		IRR 0.90 (0.80-1.12) for trend, boys
		<u>PFOA:</u> p=0.39 for trend, OR (95% CI)
· ·		4 th quartile: 1.11 (0.806–1.54)
-		
0		PFOS: p=0.008 for trend, OR (95% CI):
-		2 nd quartile: 1.44 (1.06–1.96)
-		3 rd quartile: 1.28 (0.949–1.73)
0		4 th quartile: 1.61 (1.18–2.21)
e		
-		PFHxS: p=0.93 for trend, OR (95% CI)
PFDoDA: 0.19 ng/mL		4 th quartile: 0.96 (0.73–1.41)
	RSV	
		PFNA: p=0.92 for trend, OR (95% CI)
		4 th quartile: 0.92 (0.67–1.25)
		<u>PFDA:</u> p=0.11 for trend, OR (95% CI)
		4^{th} quartile: 0.80 (0.59–1.08)
	PFDA: 0.29 ng/mL Mean maternal serum PFAS level (measured at gestation week 12) PFOA: 5.6 ng/mL	PFDA: 0.29 ng/mLGastrointestinal infections (GI) Other infectionsMean maternal serum PFAS level (measured at gestation week 12) PFOA: 5.6 ng/mLRate of hospitalization for infectious disease in young childrenPFOS: 35.3 ng/mLRisk of "total infectious diseases" in early life, based on mothers' self-administered questionnaire. "Total infectious diseases" was defined as at

Study Overview	Serum PFAS Levels	Outcome	Results
			<u>PFUnA:</u> p=0.79 for trend, OR (95% CI) 4 th quartile: 1.03 (0.76–1.40)
			<u>PFDoDA:</u> p=0.50 for trend, OR (95% CI) 4 th quartile: 1.07 (0.79–1.46)
Grandjean et al. 2012	Geometric mean maternal	Tetanus and Diphtheria	Changes in antibodies were assessed
	serum PFAS levels	antibody levels at ages 5 and 7	per two-fold increases in PFAS level.
Prospective cohort study of	PFOA: 3.20 ng/mL	, ,	
children living in the Faroe	PFOS: 27.3 ng/mL		<u>Tetanus antibody at age 5, β</u>
Islands; children were	PFHxS: 4.41 ng/mL		PFOS (age 5): -28.5% (-45.4, -6.1)
examined prior to receiving	PFNA: 0.60 ng/mL		PFHxS (age 5): -19.0% (-29.8, -6.6)
vaccine boosters (5 years of	PFDA: 0.28 ng/mL		PFDA (age 5): -19.9% (-33.1, -3.9)
age, n=532) for tetanus and			
diphtheria, 4 weeks after	Median serum PFAS levels at age		<u>Tetanus antibody at age 7, β</u>
receiving the 5-year vaccine	5		PFOA (age 5): -35.8% (-51.9, -14.2)
booster (n=456), and at age 7	PFOA: 4.1 ng/mL		PFHxS (age 5): -19.7% (-31.6, -5.7)
(n=464).	PFOS: 17.3 ng/mL		PFHxS (age 7): -22.3% (-36.3, -5.2)
	PFHxS: 0.6 ng/mL		PFDA (age 5): -22.3% (-35.8, -5.8)
	PFNA: 1.00 ng/mL		
	PFDA: 0.28 ng/mL		<u>Diphtheria antibody at age 5, β</u>
			PFNA (age 5): -16.1% (-28.8, -1.0)
	Median serum PFAS levels at age		
	7		<u>Diphtheria antibody at age 7, β</u>
	PFOA: 4.4 ng/mL		PFOA (age 5): -25.2% (-42.9, -2.0)
	PFOS: 15.5 ng/mL		PFOA (age 7): -25.4% (-40.9, -5.8)
	PFHxS: 0.5 ng/mL		PFOS (age 5): -27.6% (-45.8, -3.3)
	PFNA: 1.1 ng/mL		PFOS (age 7): -30.3% (-47.3, -7.8)
	PFDA: 0.4 ng/mL		
			No other significant associations were
			noted between maternal or child

Study Overview	Serum PFAS Levels	Outcome	Results
			(ages 5 or 7 years) PFAS levels and
			antibody levels
Grandjean et al. 2017	Median serum PFAS levels at age	Tetanus and Diphtheria	Significant association between
	7	antibody levels at ages 7 and 13	serum PFDA and antibodies for
Prospective study of 516	PFOA: 4.4 ng/mL		tetanus at age 7 (p=0.022), but not at
children living in the Faroe	PFOS: 15.3 ng/mL		age 13 (p=0.258).
Islands; serum antibodies to	PFHxS: 0.5 ng/mL		
diphtheria and tetanus were	PFNA: 1.1 ng/mL		No other significant associations
measured at age 13 and	PFDA: 0.4 ng/mL		reported between any serum PFAS
compared to serum			levels at ages 7 or 13 and diphtheria
perfluoroalkyl levels at age 7	Median serum PFAS levels at age		or tetanus antibody levels.
and 13.			
	PFOA: 2.0 ng/mL		
	PFOS: 6.7 ng/mL		
	PFHxS: 0.4 ng/mL PFNA: 0.7 ng/mL		
	PFDA: 0.3 ng/mL		
Granum et al. 2013	Maternal median serum PFAS	Rubella antibody levels	Rubella antibody levels, β
Grandin et al. 2013	levels (measured at delivery)	Hemophilus influenza type B	PFOA: -0.40 (-0.64, -0.17)
Prospective birth cohort study,	PFOA: 1.1 ng/mL	antibody levels	PFOS: -0.08 (-0.14, -0.02)
subcohort of the Norwegian	PFOS: 5.5 ng/mL	Tetanus antibody levels	PFHxS: -0.38 (-0.66, -0.11)
Mother and Child Cohort study,	PFHxS: 0.3 ng/mL	Number of episodes of	PFNA: -1.38 (-2.35, -0.40)
56 children examined annually	PFNA: 0.3 ng/mL	common cold (3-year period)	
to age 3 years; exclusion		Wheezing	Number of episodes of common cold,
criteria included maternal use			β
of steroids or anti-			PFOA: 0.42 (0.21, 0.62)
inflammatory drugs during			PFNA: 0.74 (0.05, 1.43)
pregnancy, as well as maternal			
autoimmune disease.			No other significant associations were
			reported between any PFAS and
			outcome
Huang et al. 2020	Median cord blood PFAS levels	Respiratory tract infections	There were no significant associations

Study Overview	Serum PFAS Levels	Outcome	Results
	PFOA: 6.68 ng/mL	(RTIs) (both lower and upper) in	between any PFAS and total number
Prospective birth cohort,	PFOS: 2.44 ng/mL	the first 5 years of life.	of RTIs, nor were there any significant
Shanghai Prenatal Cohort;	PFHxS: 0.16 ng/mL	Assessed through self-report	associations between PFAS levels and
mothers were enrolled at 29-41	PFNA: 0.63 ng/mL	from parents, review of medical	incidence of recurrent respiratory
weeks of gestation and children	PFDA: 0.35 ng/mL	records, and IgG and IgE levels	tract infections. There were no
were followed-up at 5 years of	PFUnA: 0.39 ng/mL	as biomarkers of humoral	associations between PFAS and IgG or
age (n=344).	PFDoDA: 0.09 ng/mL	immunity	IgE concentrations.
Impinen et al. 2018	Mean cord PFAS levels:	Number of common colds (0-2	<u>Common Cold, β</u>
	PFOA: 1.8 ng/mL	years of age)	PFUnA: 0.11 (0.08, 0.14)
Prospective study of 641	PFOS: 5.6 ng/mL	Number of lower respiratory	
infants participating in the	PFHxS: 0.3 ng/mL	infections (0-10 years of age)	<u>LRTI, β</u>
Environment and Childhood	PFNA: 0.2 ng/mL		PFOA: 0.28 (0.22, 0.35)
Asthma study in Norway;	PFUnA: 0.1 ng/mL		PFOS: 0.50 (0.42, 0.57)
health outcomes were			PFNA: 0.09 (0.03, 0.14)
evaluated at 2 and 10 years of			PFUnA: 0.18 (0.13, 0.23)
age.			
Impinen et al. 2019	Median maternal serum PFAS	Parent-reported number of	<u>Common cold, RR (95% CI)</u>
	levels (collected mid-pregnancy)	episodes of infections from 0-3	PFOA: 0.96 (0.94, 0.99)
Prospective birth cohort,	PFOA: 2.54 ng/mL	years of age and from 6-7 years	PFOS: 0.94 (0.92, 0.97)
subcohort of the Norwegian	PFOS: 12.87 ng/mL	of age. Infections included	
Mother and Child Cohort Study	PFHxS: 0.65 ng/mL	common cold (3 years only) and	Bronchitis/pneumonia at 3 years of
enrolled between 1999-2008.	PFNA: 0.45 ng/mL	bronchitis/RS-virus/pneumonia	<u>age, RR (95% CI)</u>
Children were followed-up at 3	PFUnA: 0.20 ng/mL	(3 and 7 years)	PFOA: 1.27 (1.12, 1.43)
years (n=1,270) and 7 years			PFOS: 1.20 (1.07, 1.34)
(n=972).			PFHxS: 1.15 (1.06, 1.24)
			No other associations were found for
			PFAS and episodes of infection.
Kvalem et al. 2020	Mean serum PFAS levels at 10	Parent reported number of	Number of common colds in last 12
E 11 11 11 11 11	years of age	episodes of common cold and	months at age 16, OR (95% CI) per
Prospective birth cohort, the	PFOA: 4.62 ng/mL	number of episodes of	IQR increase in PFAS from
Environment and Childhood	PFOS: 20.9 ng/mL	bronchitis and pneumonia	multinomial logistic regression
Asthma (ECA) study; PFAS	PFHxS: 3.33 ng/mL	between 10-16 years of age	models

Study Overview	Serum PFAS Levels	Outcome	Results
exposure measured at age 10 and health outcomes collected at age 16 for 378 children.	PFNA: 0.63 ng/mL PFDA: 0.19 ng/mL PFUnA: 0.18 ng/mL	and in last 12 months	Results \geq 3 vs. 0 colds, PFOS: 0.67 (0.47, 0.96) \geq 3 vs. 0 colds, PFNA: 0.63 (0.44, 0.91) \geq 3 vs. 0 colds, PFDA: 0.56 (0.37, 0.84) \geq 3 vs. 0 colds, PFUA: 0.64 (0.44, 0.92) LRTI between 10-16 years of age, RR (95% CI) per IQR increase in PFAS PFOA: 1.10 (1.02, 1.19) PFOS: 1.34 (1.17, 1.55) No other significant associations found between PFAS and common
Kielsen et al. 2016 Prospective study of 12 adults in Denmark administered a booster vaccine for tetanus and diphtheria; antibody concentrations were measured 4- and 10-days post- vaccination.	Median serum PFAS levels PFOA: 1.69 ng/mL PFOS: 9.52 ng/mL PFHxS: 0.37 ng/mL PFNA: 0.66 ng/mL PFDA: 0.30 ng/mL PFUnA: 0.21 ng/mL PFDoDA: 0.039 ng/mL	Tetanus and diphtheria antibody levels	cold or LRTIDiphtheriaPFOS: inverse association (p=0.044),unadjustedPFNA: Inverse association (p=0.004),unadjustedPFDA: Inverse association (p=0.009),unadjustedPFUnA: Inverse association (p=0.036),unadjustedPFDoDA: Inverse association (p=0.036),unadjustedPFDoDA: Inverse association (p=0.038), unadjusted
Looker et al. 2014	Geometric mean serum PFOA: 33.74 ng/mL	Seroprotection from influenza A H3N2 virus	TetanusPFUnA: Inverse association (p=0.039),unadjustedPFDoDA: Inverse association(p=0.038), unadjustedInfluenza A H3N2 virusseroprotection, OR (95% CI)

Study Overview	Serum PFAS Levels	Outcome	Results
Cross-sectional study of 411	• 1 st quartile: 0.25-13.7 ng/mL	Seroprotection from influenza	PFOA Q2: 0.34 (0.14, 0.83)
adults participating in a follow-	• 2 nd quartile: 13.8-31.5	A H1N1 virus	PFOA Q3: 0.28 (0.11, 0.70)
up study to the C8 Health	ng/mL	Seroprotection from influenza	PFOA Q4: 0.39 (0.15, 0.99)
Project; all participants	• 3 rd quartile: 31.6-90 ng/mL	type B virus	
received an influenza vaccine	• 4 th quartile: 90.4–2,140	Cold or flu infection	No other significant associations were
and were examined prior to	ng/mL	Frequency of colds	reported for PFAS levels (continuous
vaccination and 21 days post			or categorical) and the outcomes of
vaccination; 755 adults in the	Geometric mean serum PFOS:		interest.
follow-up study participated in	8.32 ng/mL		
a survey evaluating self-			
reported colds and influenza			
episodes.			
Manzano-Salgado et al. 2019	Mean maternal serum PFAS	Mother-reported occurrence of	There were no associations between
	levels (collected during first	LRTI at 1.5, 4, and 7 years of	LRTIs and log-transformed PFAS
Prospective cohort – Spanish	trimester)	age (defined as occurrence of	levels.
INMA birth cohort study (2003-	PFOA: 2.67 ng/mL	bronchitis, bronchiolitis, or	
2008) – of children with follow-	PFOS: 6.41 ng/mL	pneumonia)	
ups at 1.5 years (n=1,188), 4	PFHxS: 0.67 ng/mL		
years (n=1,188) and 7 years (n-	PFNA: 0.74 ng/mL		
1,071).			
Okada et al. 2012	Median maternal serum PFAS	Infectious disease during the	There were no statistically significant
	(measured after second	first 18 months of life	associations between maternal serum
Prospective cohort study of 343	trimester)		PFAS levels and infectious diseases
pregnant women in Japan; cord	PFOA: 1.3 ng/mL		during the first 18 months of life.
blood samples collected at	PFOS: 5.2 ng/mL		
delivery to measure total IgE			
levels; infant allergies and			
infectious disease information			
collected during first 18 months			
of age.			
Pilkerton et al. 2018	Mean serum PFAS	Rubella IgG titers (log titers in	<u>Adults,</u> β
	PFOA	analyses)	Men
Cross-sectional study of	Men: 6.0 ng/mL		PFOA Q3: -0.55 (-0.81, -0.28)

Study Overview	Serum PFAS Levels	Outcome	Results
participants (≥ 12 years of age)	Women: 4.3 ng/mL		PFOA Q4: -0.45 (-0.84, -0.05)
from NHANES 1999-2000 and	Youth: 4.8 ng/mL		
2003-2004 data (n=581 women	PFOS		No significant associations with
and 621 men adults [19-49	Men: 28.1 ng/mL		Rubella titers for PFOA or PFOS in
years], and 1012 youth [12-18	Women: 22.1 ng/mL		youth or in adult women. No
years]).	Youth: 25.1 ng/mL		associations between PFOS and
			Rubella titers in adult men.
Shih et al. 2021	Median serum PFAS	Serum antibody concentrations	No significant associations between
	concentrations at birth	for Hepatitis A, Hepatitis B,	PFAS collected at any age were found
Prospective cohort study of	PFOA:1.11 ng/mL	Diphtheria, and Tetanus;	with HAV or HB at age 28 years.
adults (28 years old) in the	PFNA: 0.11 ng/mL	assessed 6 months after the	
Faroese Island cohort recruited	PFDA: 0.07 ng/mL	booster for diphtheria and	Positive associations of diphtheria
in 1986-1987; blood samples	PFHxS: 0.21 ng/mL	tetanus and 6 months after the	were found with cord-blood PFAS and
collected in cord blood, at 7,	PFOS: 5.96 ng/mL	completion of the 3-dose	PFAS at ages 22 and 28 years.
14, 22, and 28-years; 401		hepatitis vaccine.	
cohort members for Hepatitis A	Median serum PFAS		No evidence of associations was
and B and 281 cohort members	concentrations at age 28 years		observed for tetanus.
for tetanus and diphtheria	PFOA: 1.28 ng/mL		
were included.	PFNA: 0.98 ng/mL		
	PFDA: 0.34 ng/mL		
	PFHxS: 0.41 ng/mL		
	PFOS: 6.85 ng/mL		
Stein et al. 2016a	Geometric mean serum PFAS	Serum cytokines (IFN-α2, IFN-γ,	Significant associations between
	PFOA: 2.28 ng/mL	TNF- α , IP 10) and chemokines	serum PFHxS and changes in the
Cross-sectional study of 78	PFOS: 5.22 ng/mL	(MCP-1, MIP1a) were measured	serum cytokines IFN- γ (p=0.05) and
healthy adults in New York city	PFHxS: 1.1 ng/mL	pre-vaccination and 3- and 30-	TNF-α (p=0.04).
vaccinated during the 2010-	PFNA: 0.77	days post vaccination; nasal	
2011 season with the intranasal		cytokine (IP-10), chemokine	No other associations between serum
FluMist influenza vaccine.		(MCP-1), and nasal mucosal IgA	PFAS and seroconversion as
		were measured 3- and 30-days	measured by hemagglutinin inhibition
		post vaccination	or immunohistochemistry. No other
			associations between serum PFAS
			and changes in serum cytokine or

Study Overview	Serum PFAS Levels	Outcome	Results
			chemokine levels or nasal cytokine, chemokine, or IgA levels.
Stein et al. 2016b Cross-sectional study of adolescents (12–19 years of age) utilizing NHANES 1999– 2000 and 2003–2004 data (n=1,191).	Geometric mean serum PFAS PFOA: 4.13 ng/mL PFOS: 20.8 ng/mL PFHxS: 2.47 ng/mL PFNA: 0.765 ng/mL	Antibody titers for measles, mumps, and rubella in whole cohort as well as in seropositive subcohort	Full Cohort, β per 2-fold increase inPFASMumpsPFOS: -7.4% (-12.8, -1.7)Seropositive subcohort, β per 2-foldincrease in PFASMumpsPFOA: -6.6% (-11.7, -1.5)PFOS: -5.9% (-9.9, -1.6)RubellaPFOA: -8.9% (-14.6, -2.9)PFOS: -13.3% (-19.9, -6.2)PFHxS: -6.0% (-9.6, -2.2)No other significant associations were observed in the full cohort or
Timmermann et al. 2020	Madian corum DEAS lovals	Moosles antibady titors at	subcohort.
Timmermann et al. 2020 Subset analysis of RCT of early	Median serum PFAS levels collected from children at baseline (4-7 months of age)	Measles antibody titers at baseline (no measles vaccination), 9-month visit (1	<u>9-month visit, control, β</u> PFOS: -27% (-44, -4)
measles vaccination conducted in Guinea-Bissau from 2012- 2015 (n=237). Intervention (n=135) included two doses of measles vaccine (at 4-7 months and at 9 months); control (n=102) included the usual single vaccination (at 9 months).	PFOA: 0.68 ng/mL PFOS: 0.77 ng/mL PFHxS: 0.10 ng/mL PFNA: 0.21 ng/mL PFDA: 0.19 ng/mL PFUnA: 0.12 ng/mL	measles vaccination in intervention vs. none in controls), and at 2 years of age (2 measles vaccinations in intervention vs. 1 measles vaccination in control) with a doubling of serum PFAS at inclusion.	9-month visit, intervention, βPFOS: -20% (-35, -1)PFDA: -25% (-42, -4)No other significant associationsbetween percentage difference inmeasles antibody concentrations forany PFAS at any time points.

Study Overview	Serum PFAS Levels	Outcome	Results
		Presence of fever, coughing,	Coughing, OR (95% CI)
		diarrhea, and any morbidity at	PFOA: 1.87 (1.02, 3.45)
		inclusion and at 9-month visit	PFHxS: 2.15 (1.17, 3.97)
		with doubling of serum PFAS	
		concentrations at baseline	Any Morbidity, OR (95% CI)
			PFOA: 2.02 (1.20, 3.41)
			PFHxS: 1.82 (1.06, 3.11)
Zeng et al. 2019	Cord blood serum PFAS levels	Antibody titers against 2 Hand,	CA16 antibody in cord blood, OR
	PFOA: 1.22 ng/mL	Foot, and Mouth Disease	<u>(95%CI)</u>
201 mother-infant pairs in the	PFNA: 0.16 ng/mL	(HFMD) viruses, enterovirus 71	PFOA: 1.56 (1.13, 2.14)
Guangzhou Birth Cohort Study,	PFDA: 0.12 ng/mL	(EV71) and coxsackievirus A 16	PFOS: 1.75 (1.16, 2.63)
enrolled in 2013.	PFUnA: 0.13 ng/mL	(CA16), were assessed in cord	Total PFAS: 2.24 (1.30, 3.85)
	PFDoDA: 0.05 ng/mL	blood and blood serum at 3	
	PFHxS: 3.96 ng/mL	months of age	CA16 antibody in 3-month infants, OR
	PFOS: 3.17 ng/mL		<u>(95%CI)</u>
	Total PFAS: 10.56 ng/mL		PFOA: 1.73 (1.08, 2.75)
			PFNA: 1.50 (1.04, 2.17)
			PFDA: 2.22 (1.42, 3.47)
			PFUnA: 1.45 (1.02, 2.08)
			PFOS: 1.71 (1.12, 2.60)
			Total PFAS: 2.74 (1.33, 5.61)
			EV71 antibody in cord blood, OR
			<u>(95%CI)</u>
			PFOA: 1.49 (1.09, 2.05)
			PFNA: 1.44 (1.02, 2.02)
			PFDA: 1.49 (1.03, 2.16)
			PFOS: 1.66 (1.12, 2.45)
			Total PFAS: 1.90 (1.14, 3.16)
			EV71 antibody in cord blood, OR
			(95%CI)
			PFOA: 2.11 (1.27, 3.48)

Study Overview	Serum PFAS Levels	Outcome	Results
			PFNA: 1.01 (1.04, 2.16)
			PFDA: 2.05 (1.33, 3.18)
			PFUnA: 1.48 (1.04, 2.10)
			PFHxS: 1.49 (1.06, 2.10)
			PFOS: 2.25 (1.44, 3.51)
			Total PFAS: 4.55 (2.06, 10.06)
Zeng et al. 2020	Median serum concentrations of	Change in hepatitis B surface	Mean difference (95% CI) in serum
	PFAS among seropositive	antibody (HBsAb) per log serum	HBsAb
Cross-sectional study of 605	PFOA: 5.10 ng/mL	PFAS.	PFOS: -0.51 (-0.84, -0.18)
participants from the Isomer C8	PFOS: 10.1 ng/mL		
Health Project in China (2015-			No associations were found for PFOA
2016). The population included	Median serum concentrations of		
509 participants who were	PFAS among seronegative		
HBsAb seropositive and 96	PFOA: 5.54 ng/mL		
participants who were HBsAb	PFOS: 14.1 ng/mL		
seronegative			

OR=odds ratio, IRR=incidence rate ratio, IQR=interquartile range