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# ITRC PFAS Resources

## ITRC PFAS: <https://pfas-1.itrcweb.org/>

#### Guidance Document

## 13 Fact Sheets

#### External Tables

## PFAS Introductory Training

▪ Clu-In Archive: [https://www.clu](https://www.clu-in.org/conf/itrc/PFAS-Introductory/)[in.org/conf/itrc/PFAS-Introductory/](https://www.clu-in.org/conf/itrc/PFAS-Introductory/)

## Other video resources

- **E** Available through links on: [https://pfas-1.itrcweb.org](https://pfas-1.itrcweb.org/)
- **Quick Explainer Videos**
- **Example 1 Longer PFAS Training Modules**
- **EXECUTE: Archived Roundtable Sessions**





# PFAS: Beyond the Basics Training

# Fate and Transport Site Characterization Source Identification & Forensics





# Today's PFAS Trainers



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## ITRC PFAS Team: 2024 "Beyond the Basics" Training Modules



# Learning Objectives

#### Site Characterization

- Concerns for different source types
- Hydrogeological setting
- Sampling and Analysis
- Data interpretation and visualization

#### Fate and Transport

- Small-scale processes
- Movement in and between environmental media
- Differential transport of PFAS species (environmental fractionation)
- Precursor transformation

#### Forensics

- PFAS fingerprinting
- Identification of unknown sources
- Techniques for distinguishing between multiple or overlapping sources

**Conceptual** Site Model





# Fate and Transport Considerations



# Learning Objectives

### Site Characterization

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### **Fate and Transport**

- Small-scale processes
- Movement in and between environmental media
- Differential transport of PFAS species (environmental fractionation)
- Precursor transformation

#### Forensics

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**Conceptual** Site Model

# F&T: What is Fate and Transport?



- Perfluorinated vs. polyfluorinated
- Fluorinated tail: chain length
- Non-fluorinated head: functional group and charge state

# Fate & Transport

#### ▪ Nature of Release

▪ Soil properties



- Groundwater depth/velocity
- Groundwater geochemistry
- Prevailing atmospheric conditions
- Co-contaminants



# PFAA Compound Properties



PFAS figure source: M. Olson, Trihydro. Used with permission.

Energetically favorable Air Reduces the surface tension at the interface Water

PFAAs have a greater affinity than traditional surfactants

> PFAS-1, Figure 4-1. Source: D. Adamson, GSI. Used with permission.

# Microscale Partitioning

## **Electrostatic Interactions**

- pH (changing pH changes surface charges)
- Presence of polyvalent cations (Ca2+ Mg2+)
- Clay (positive surface charge)



## **Hydrophobic Interactions**

- **Organic carbon sorption** consideration
	- ↑ Chain length ↑ Sorption to organic carbon
	- **PFSA sorb more strongly** than PFCA to organic carbon



# Microscale Partitioning: Interfacial Behavior

#### The relative importance of phase partitioning changes between the source area and the plume.



PFAS-1, Section 5.2, Phase Partitioning. PFAS-1, Figure 5-1. Fate and transport processes relevant for PFAS. Source: D. Adamson, GSI. used with

# PFAA Partitioning Depends on Amount of Interfacial Area



- Degree of water saturation affects the amount of interfacial area (nonlinear)
- Larger interfacial area enhances PFAA retention
- Factors affection A-W interfacial area:
	- Soil type
	- **•** Grain size
	- **EXEC** Heterogeneity
	- Organic Content

# Microscale Factors Influencing Solid Phase Sorption



## Microscale Partitioning Summary



# Supramolecular Aggregations

- **EXECT:** At high concentrations, certain PFAS can aggregate due to surfactant properties
	- Forms micelles, hemi-micelles, and bilayers
- **If** May not be directly analogous to traditional micelle formation – more research needed.
- May be more complicated in environmental settings
	- Aggregations can occur well below CMC
	- **Interaction with charged or hydrophobic** surfaces can affect sorption
- **More research is needed to understand** how this affects fate and transport



PFAS-1, Figure 5-2. Source: D. Adamson, GSI. Used with permission.



# F&T: PFAA Precursors (environmental conditions)





See PFAS-1, Section 2.2 for more detailed information regarding nomenclature & the PFAS family tree.

# F&T: Transformation of Precursors to PFAAs





PFAA



PFAS figures source: M. Olson, Trihydro. Used with permission. PFAS-1, Section 5.4.4.2 Aerobic Biological Pathways.

# Abiotic transformations (Oxidation reactions)

- Hydrolysis of fluoropolymers can form PFOA
- Hydroxyl radicals formed in natural systems oxidize precursors
- Fluorotelomers oxidize faster than sulfonamides
- PFAAs are not always formed (not complete oxidation)
- **Indirect photolysis in the atmosphere (ex.**  $FtoH \rightarrow PFCAs$



# Aerobic Biotransformation (Oxidation)



Sulfonamides (slower rates than Fluorotelomers, **mainly** form PFSAs)



PFAS-1, Section 5.4.4.2 Aerobic Biological Pathways. Figure source: Christopher Olivares, UC Irvine. Used with permission.

# Anaerobic biotransformation

## Fluorotelomer



Fluorotelomer acids + other PFAS formed under more reducing anaerobic conditions

## Feammox (Acidimicrobium sp. Strain A6)

- **EXA** Anaerobic ammonium oxidation coupled to iron reduction
- **Reported defluorination of PFOA, PFOS with release of fluoride**

# Macroscale Fate and Transport



# Macroscale Transport: Air

- PFAS sources to air
	- Industrial facilities producing or using PFAS
	- Areas where fluorine foams are used/released
	- Waste management facilities (landfills, WWTPs, biosolids production)
- Fluorotelomer alcohols (FTOHs) and some other PFAS are volatile
- Stack emissions can include volatile PFAS and PFAS attached to particulate matter
- Atmospheric deposition (wet or dry) may result in PFAS contamination several miles from industrial emission sources
- Long-range transport processes and effects are similar to atmospheric transport of other recalcitrant contaminants



PFAS-1, Section 5.2.4, Partitioning to Air and Section 5.3.2 PFAS Transport in Air PFAS-1, Figure 5-1. Fate and transport processes relevant for PFAS. Source: D. Adamson, GSI. used with permission.

# Macroscale Transport: Vadose Zone

- **PFAS source areas: vadose-zone** retention may be significant
	- Cationic/zwitterionic precursors may strongly sorb
	- Long-chain PFAA sorption
		- Depends on soil chemistry
	- PFAAs accumulate at air/water interfaces (may retard transport)
- **Most PFAS have low volatility (little to** no partitioning to vapor phase)
- Leaching may provide long-term source to groundwater



# Macroscale Transport: Groundwater

- Readily transported once in groundwater
	- Same processes affecting sorption in vadose zone, but now with groundwater chemistry interactions
	- Dependent on chain length and functional group
	- $\bullet$  K<sub>oc</sub> important, but not sufficient to explain partitioning behavior
- Potential impacts of remedial activities targeting co-contaminants
	- Introduction of oxygen (e.g. air sparge, DO injection) can drive precursor transformation
	- Introduction of cations (e.g., ISCO) can lead to enhanced sorption/retardation of PFAA transport.



# Macroscale Transport: Surface Water

- Type of water body is important
	- Fast moving, suspended sediments
	- Slow moving, deposition
- Uptake to biota (incl. fish)
- Stratification in water column
- Dissolved PFAS in SW can move downstream, back into GW
- PFAAs aggregate at surface microlayers
	- Foam formation with wind or turbulence



# Transport in Surface Water: Foam

## PFAS-Containing Foam Considerations

- **Example 1 Fransport** as "foam islands" to a new location
- Collapse of foam and **dissolution of PFAS** back into water column
- PFAS concentration in foam > water column
- PFAS in foam potentially leads to **additional exposure pathways** – both human and ecological receptors

\*Note – foams can be produced by natural processes in the absence of PFAS as well; not all surface water foam is guaranteed to contain PFAS.





Photographs courtesy of R. Higgins, MPCA. Used with permission.

## Transport Between Surface Water and Groundwater

## Transitions zones between SW and GW tend to have changes in conditions

- $\bullet$  Low oxygen GW  $\rightarrow$  oxygen-rich SW
	- Increased microbial activity
	- Increased precursor transformation
	- Increased proportion of PFAAs
- **Differences in organic and mineral content** between soils and sediments
	- Sorption to sediments and colloids
	- Changes in salinity affect sorption
- Potential for dilution and mixing depending on type of water body



## Co-contaminant considerations

Petroleum co-contamination common at AFFF sites and can occur at other PFAS sites

#### PFAS may partition into LNAPL or accumulate at water/LNAPL interface

■ Can lead to greater retention of PFAS

Petroleum contamination leads to reducing conditions

- **EXECUTE: Slows down precursor transformation**
- **EXECUTE:** Shifts precursor transformation to anaerobic processes/pathways

If petroleum remediation has occurred, may alter redox to oxidizing conditions (e.g., air sparge) or change ionic concentrations (e.g., ISCO)

■ Can alter rates of precursor transformation

DNAPL co-contamination can also result in increased PFAS retention

# Environmental Fractionation

- Differential transport and precursor transformation lead to environmental fractionation
- As PFAS move downgradient from source area, composition of PFAS changes
- Source type and hydrogeochemical parameters influence type and rates of change







# Site Characterization



# Learning Objectives

## **Site Characterization**

- Concerns for different source types
- Hydrogeological setting
- Sampling and Analysis
- Data interpretation and visualization

## Fate and Transport

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**Conceptual** Site Model

## Forensics

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# Work Plan and Conceptual Site Model (CSM)

Initial step in Site Characterization is CSM development to inform Data Quality Objectives

CSM development should consider:

- **E** Site characteristics
- Properties of contaminants

Use CSM to identify data gaps and sampling needs



## Main Sources of PFAS Release to the Environment


## CSM for AFFF Application Sites



PFAS-1, Figure 2-19, CSM for fire training areas. Figure Adapted from figure by L. Trozzolo, TRC, used with permission.

## AFFF Contains Highly Diverse Mixtures

#### AFFF product chemistry has changed over time



Table 3-1 of the ITRC PFAS Technical Regulatory Guidance document presents types of foam and composition.

Barzen-Hanson et. Al 2017. Environ Sci Tecnol 51: 2047-2057 Figures used with permission from J. Field, Oregon State.

## Forensic Analysis of PFAS from AFFF Sources

#### **Forensic analysis of AFFF is based on fingerprinting the composition of unknown samples**

#### **Multiple lines of evidence and methods should be considered for site-specific fingerprinting:**

• AFFF use history

- AFFF manufacturing processes
- Targeted PFAS analysis, TOP analysis for precursor transformation, and TOF analysis
- The effects AFFF source commingling and cocontaminants on fate and transport behavior



## CSM for Industrial Sites



## CSM for Landfills and WWTPs



\*Leachate release from lined landfills could occur in the event of a liner leak

**O** Infiltration **O** Transformation of precursors (abiotic/biotic) KEY Atmospheric Deposition **O** Diffusion/Dispersion/Advection

PFAS-1, Figure 2-20, CSM for landfills and WWTPs. Figure Adapted from figure by L. Trozzolo, TRC, used with permission.

# Source-specific considerations

#### AFFF

- Diverse mix of PFAS
- Manufacturing methods determine PFAS composition
- Fate and Transport affected by co-contaminants and historical remediation

#### **Industrial**

- **PFAS mixture may be less diverse**
- Site-specific process determines PFAS composition
- Air transport may be important (may cause diffuse soil sources from deposition)

#### Landfill

- 5:3 FTCA as "signature compound"
- May include volatile PFAS (FTOHs and PFBA; possible short-range deposition)
- Types of waste accepted determine PFAS composition and concentration

#### WWTP & Biosolids

- Short chains and terminal products may dominate (not always)
- Types of waste accepted AND treatment process determine PFAS mix



chate release from lined landfills could occur in:



PFAS-1, Section 2.6 Releases to the Environment.

### Site Characterization: General Considerations

### Conceptual Site Model

#### Review existing data and information

Site history, PFAS use/types, release locations, mixtures released, co-contaminants

Nature and location of potential receptors

#### Hydrogeologic setting

- Topography/Drainage
- Soil type
- Depth to water
- Nature of aquifer system & characteristics – unconfined/confined; permeability, porosity, velocity
- Nature of GW discharge seeps, tributaries, rivers, lakes, supply wells

Surface Water Relevant regulations, permits, standards

# Site Characterization: General Considerations

### **Transport Pathways**

- Release to GW
	- Vadose Zone Depth to water; soil type (sands vs. clays; clay layering); organic carbon content; pH; ion exchange capacity
	- Air-Water Interface Grain size; moisture content
	- $GW$  System Soil type; pH; TDS/salts/divalent cations, cocontaminants; confining units
- Release to SW
	- **GW** discharge
	- **Storm runoff**



# What to Analyze

### **VADOSE ZONE**

- Soil type/particle size/porosity
- Moisture content
- Cation Exchange Capacity (CEC)
- **Example 2** Surface charge
- Multivalent cations
- Soil pH
- Organic Carbon
- Depth to GW

\* Focus on air-water interface and clay lenses and layers



#### **SATURATED GROUNDWATER SYSTEM**

- pH, TDS, salts, turbidity, multivalent cations, co-contaminants
- GW gradient and velocity
- Depth to basement (unconfined vs confined conditions)
- Presence and depth to fractured rock, if applicable

\* Focus on potential influences to GW movement (nearby pumping wells, GW-SW interactions, etc)



# High Potential for PFAS Sample Contamination

Low PFAS screening or regulatory criteria

Inefficient decontamination procedures in source areas

Possible contamination from, or sorption to, equipment and materials





## PFAS-Containing Foam – Four Layers

Foam – contains higher concentrations of PFAS than the underlying layers; as foam is formed, it removes PFAS from the water column.

Surface micro layer  $-$  about 50  $\mu$ m thick - includes the airwater interface. Likely highest concentration of PFAS in water column

Neuston Layer – zone directly below surface micro layer. Rich in aquatic organisms

Underlying water column





# Analytical Method Differences



PFAS-1, Section 11.2.1.3 Sample Analysis and Table 11-3.

### TOP Assay (qualitative)



- **Estimate concentrations of oxidizable** precursors in sample.
- **Precursors can transform to** measurable PFAAs. TOP Assay oxidation forces transformation.
- **Predominant precursor transformation** products are perfluorocarboxylic acids (PFCAs).
- Increased concentrations of PFCAs after oxidation provide estimate of oxidizable precursors.
- Potential low biases:
	- Incomplete oxidation
	-



# Data Visualization: PFAS Chemical Profiles

#### Many different methods to visualize PFAS profiles (all based on the same data)



#### **Bar Charts**

- Grouped by PFAS class and ordered by MW
- Y-axis allows consideration of concentrations
- 

#### **Pie Charts**

- Grouped by PFAS class and ordered by MW
- Limited space to label compounds with large compound list
- Can be difficult to compare samples
- Easier to interpret with 10 or fewer compounds



- **Radar/Spider Charts** • Ordered by PFAS class ordered by MW PFMPA NEtFOSAA Limited space to label **NEtFOSE** PFUnA NMeFOSAA PEDoA compounds with a large NMeFOSE PFTrDA compound list **NEtFOSA** PFTeA • Easy to compare 3-5 samples, NMeFOSA PFHxDA 10.2FTS PFODA gets cluttered with more 8.2FTS 6.2FTS samples in a single plot
- Easier to interpret with 10 or fewer compounds



■ Can get cluttered with many samples | | Sar Chart figures adapted from PFAS-1 Figure 10-2. Source M. Benotti, NewFields. Used with 50 permission. Figures source: Michael Bock, Verdantas. Used with permission.

## Data Visualization: Spatial Coordinates



## Visual Representations of Data: Case Study 1

Example is from a site in Sweden where multiple fire training areas impacted groundwater

Rather than percent abundance, axes display a log-based 10 scale

Total PFCAS and PFSAS shown instead of individual compounds, to identify general group patterns

6:2 FTS differentiated from other precursors due to sitespecific considerations

Patterns in shape pertain to precursors vs. PFCA/PFSA relationships





### Data Visualization: Case Study 1

- All chemical-specific precursors non-detect in main aquifer channel, attenuated near fire training areas.
- **PFSA and PFCA concentrations decline** along the main flow channel (for example, downgradient of G8 and G9).
- Concentrations of precursors in the vicinity of the fire training areas shown in the inset map are much higher than in the main groundwater flow channel.
	- Sorption
	- Biotransformation of 6:2 FTS and other precursors
	- Possibly groundwater  $-$  surface water interactions
- Elevated PFCAs, PFSAs, and 6:2 FtS near at least two fire training areas: AFFFs produced using telomerization used at one point.



PFAS-1, Figure 15-3. PFAS composition in groundwater. Source: G. Carey, Porewater Solutions. Used with permission. 53

## Data Visualization: Case Study 2

- Comparison of source zone concentrations (dark blue outline) vs. downgradient samples (light blue fill) show changes due to
	- **•** Migration
	- Dissolved Oxygen infusion wells
- Introduction of oxygen  $\rightarrow$  transformation of precursor compound FHxSA.
- Some differential transport of short and long chains.



# Questions

#### **Site** Characterization

- Concerns for different source types
- Hydrogeological setting
- Sampling and Analysis
- Data interpretation and visualization

### Fate and Transport

- Small-scale processes
- Movement in and between environmental media
- Differential transport of PFAS species (environmental fractionation)
- Precursor transformation

**Conceptual** 

Site Model

#### Forensics

- PFAS fingerprinting
- Identification of unknown sources
- Techniques for distinguishing between multiple or overlapping sources

ECO<sub>S</sub>

<https://pfas-1.itrcweb.org/> <sup>55</sup>





# Understanding PFAS Sources



## Learning Objectives

### Site Characterization

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### Fate and Transport

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#### **Forensics**

- PFAS fingerprinting
- Identification of unknown sources
- Techniques for distinguishing between multiple or overlapping sources

**Conceptual** Site Model

## What is Environmental Forensics?

Environmental forensics is the application of the scientific method to answer questions about contamination events and their origin.

#### Link contamination to a source

Distinguish between multiple sources

**Estimate** the contribution from multiple individual sources to overlapping plumes

Evaluate effects of remediation or natural attenuation Predict or model sitespecific fate and transport of contaminants



# PFAS Forensics Applications

- PFAS Forensics: application of environmental forensics to PFAS
	- Arguably in its nascent stages
- Applications so far include:
	- Source identification
	- Differentiation of overlapping sources
	- Evaluation of environmental fractionation
	- Identifying the types of AFFF used at a site
	- Identifying transport-related signatures
- Setting and history are very important





PFAS-1, Figure 15-2 Radial diagrams illustrating PFSA trends at an AFFF release site. Source: G. Carey, Porewater Solutions. Used with permission. PFAS Figure Source: M. Olson, Trihydro. Used with permission.

### Source Identification: process of finding or identifying a source of contamination.

• Often applied when contamination is discovered without prior knowledge of source

### Forensic methods can also be applied to source differentiation

- Two plumes from different sources overlap
- Anomalous data within known plume points to other undiscovered sources
- Distinguish between point and nonpoint (i.e., diffuse) sources – use caution



# PFAS Forensics Techniques

- **Diagnostic ratios of PFAS mixtures** (e.g., PFOS:PFOA)
- Data visualization techniques (e.g., radial diagrams, bar charts)
- Specialized analytical techniques (e.g., non-target analysis, linear/branched speciation)
- Advanced multivariate statistics
	- (e.g., PCA, cluster analysis, receptor models)





PFAS-1, Figure 10-4 PCA scores plot of the analysis of a PFAS data set. Source: M. Benotti, NewFields. Used with permission. PFAS Figures Source: M. Olson, Trihydro. Used with permission. 61

#### PCA scores plot

Table 10-1. Example diagnostic ratios to compare PFAS signatures of Sample A or Sample B





# PFAS Profiles: Bar Charts



- Comparison of PFAS signatures in individual samples using "unstacked" bar graphs
- Bars are scaled to show relative contributions, but y-axis can be used to ascertain concentrations
- Can be used as a quick way to compare PFAS profiles between one or several environmental samples



PFAS-1, Section 10.5.1.2 Figures adapted from PFAS-1 Figure 10-2. Source M. Benotti, NewFields. Used with permission. Figure source: Michael Bock, Verdantas. Used with permission.

### Diagnostic ratios of PFAS mixtures: Stacked Bar Charts

- Stacked bar charts show compositional differences in PFAS.
- Removes the effect of total concentration, permitting similar fingerprints (e.g., potential sources) to plot together regardless of magnitude.
- Can be used as a quick way to compare PFAS profiles between few environmental samples











PFAS-1, Figure 15-1. Comparison of shallow and deep soil samples at an AFFF release site. Source: G. Carey, Porewater Solutions. Used with permission.

# Data Visualization: Radial Diagrams

- Radial diagrams (a.k.a., radar charts, spider plots) – useful for quick visual reference
	- Can be useful beyond source differentiation.
	- Section 15.1 of ITRC PFAS-1 has two case studies showing environmental fractionation.
- Comparison of PFAS signatures in hypothetical Sample A and B using Radial Diagrams.
- **Plotting position is PFAS percent relative** abundance (concentration divided by total PFAS).
- Can be used as a quick way to compare PFAS profiles between few environmental samples on one graph, or many on a map.





# Multivariate statistics

- In **PCA** correlated variables are transformed into a small number of uncorrelated variables. These principal components are used to simplify the visual representation of the variability in the dataset
- In **factor analysis/receptor models**, end member/factors that characterize the underlying structures in the dataset are identified. End/memebers/factors are used to simplify the visualization of the variability in the dataset.
- In **cluster analysis**, the similarity/dissimilarity between samples are calculated and used to define clusters of similar samples in a larger dataset.





# Advanced multivariate statistics: Principal Component Analysis (PCA)





# Advanced multivariate statistics: Clustering Analysis (CA)

Groups samples based on similarity/dissimilarity (i.e., chemical profile similarity)

- Not all clusters are sources. They may also reflect:
	- Site-specific fate-and-transport processes
	- Background/ambient
	- A mixture of sources found at similar proportions in large groups of samples





# Advanced multivariate statistics: Factor Analysis (FA) and receptor models

#### Blind Source Separation

- **Estimate number of factors/end members**
- Estimate profiles for each factor/end member
- Estimate contribution of each factor/end member to each sample
- Not all end members are sources some may be weathering or transformation patterns, etc.





# Advanced multivariate statistics: Limitations

No method, including multivariate statistics, is a silver bullet

- Factor analysis and clustering typically require user to specify number of factors or clusters.
- Iterative process to find a realistic solution.
- Practitioner must interpret results (clusters or factors don't inherently "mean" anything).









PFAS-1, Section 10.5 Source Identification. Illustration of generalized clustering. PFAS-1, Figure 10-4 PCA scores plot of the analysis of a PFAS data set. Source: M. Benotti, NewFields. Used with permission. Bar Chart figures adapted from PFAS-1 Figure 10-2. Source M. Benotti, NewFields. Used with permission. Bar charts and radar charts figure source: Michael Bock, Verdantas. Used with permission. bar charts and radar charts hydre source. Michael DOCK, verdantas. Osed with permission.<br>PFAS molecule diagrams. Christopher Olivares, UC Irvine. Used with permission.

## Advanced multivariate statistics: Limitations

#### Site conditions often produce many non-source signatures:

- Diffuse (ambient) signatures or stable mixtures found in large numbers of samples
- Patterns due to environmental fractionation or differential transport
- Chemical (i.e., precursor) transformations

#### Always use multiple lines of evidence

- Other lab analyses
- Spatial / temporal trends
- Site-specific history and analytical records
- etc.



 $(\ )$ 



# Environmental Factors affecting PFAS Forensics

Sites typically contain mixtures of PFAS that exhibit wide range of physical and chemical properties

- **PFAS transported from source zones are redistributed** downgradient
- Groundwater transport depends on:
	- Soil characteristics (OM, pH, minerology, etc)
	- **PFAS characteristics (chain length, functional groups, molecular structure,** etc.)


# PFAS Forensics Examples

Examples of fingerprinting studies for source identification include:

HRGCMS was used to assign the identity of 14 major compounds that accounts for a majority of the detectable PFAS in contemporary AFFF. (Ruyle et al 2021)

PFAS profiles have been developed for different sources, including various AFFF formulations. (Dasu et al 2022)



## Review and Wrap Up

### Site Characterization

- Concerns for different source types
- •Hydrogeological setting
- •Sampling and Analysis
- Data interpretation and visualization

#### Fate and Transport

- •Small-scale processes
- •Movement in and between environmental media
- •Differential transport of PFAS species (environmental fractionation)
- Precursor transformation

#### Forensics

- PFAS fingerprinting
- Identification of unknown sources
- Techniques for distinguishing between multiple or overlapping sources

**Conceptual** Site Model



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# Questions



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