

1 Introduction

This fact sheet provides an overview of site characterization for PFAS and media-specific occurrence of PFAS. Additional information is available in the Guidance Document.

The intent is not to present general site characterization principles, but to highlight unique considerations for this family of emerging contaminants. The general principles of site characterization are similar for PFAS as for any contaminant, in that the physical setting, release specifics, proximity to receptors, and fate and transport characteristics will determine the sampling locations and requirements. PFAS toxicity, persistence, mobility, ubiquity, the large number of compounds in this family of chemicals, the variability and uncertainty of specific compounds and their criteria being regulated, and the emerging nature of PFAS, must be considered in PFAS site characterization efforts. Section 10 of the Guidance Document focuses on groundwater and source identification, and also covers other media.

Media-specific occurrence of PFAS in the environment is an active area of research. The material presented in this fact sheet and in Section 6 of

ITRC has developed a series of fact sheets that summarizes recent science and emerging technologies regarding PFAS. The information in this and other PFAS fact sheets is more fully described in the *ITRC PFAS Technical and Regulatory Guidance Document (Guidance Document)* (https://pfas-1.itrcweb.org/).

This fact sheet describes:

- Considerations for completing site characterizations where PFAS has been released to the environment
- Briefly provides information about media-specific occurrence of PFAS

the Guidance Document is not the result of an exhaustive literature review but is included to provide a general understanding of PFAS concentrations. Because analytical methods are still being optimized and standardized, it is often difficult to compare results between studies and the reported data may offer various levels of confidence. Media types considered in a site investigation include air, soil and sediment, groundwater, surface water, and biota.

2 Site Characterization Issues Relevant to PFAS

Evolving Science and Regulations

The state of the science and regulatory environment continue to evolve. Therefore, many aspects of PFAS, such as toxicology and behavior in the environment, sampling and analytical methodologies, and the regulatory environment including regulatory limits and compounds of interest need to be reevealuated during the course of characterizing a PFAS site. Section 11 of the Guidance Document covers sampling and analysis for PFAS and Section 8 includes information about regulations.

Source, Fate and Transport Properties and Other Considerations

Sources: An overview of PFAS sources is discussed in Section 2.6 of the Guidance Document. In addition to the main sources that include industrial discharges and AFFF releases, there are also "secondary sources," such as sources created through movement of contaminated media into an area that was previously uncontaminated, or an area where physical or chemical processes have concentrated PFAS (for example, atmospheric deposition or multi-media interfaces).

Historical view: Historical investigations may not have assessed PFAS contamination because it was not regulated, not a contaminant of concern during the original investigation, or because the analytical methods were unavailable at the time. Plumes may be extensive, having years to develop before being discovered or addressed.

Pathways: PFAS may be present or migrate via pathways that are not typically observed with other compounds. For example, PFAS may be present in groundwater at a site via air deposition from an off-site source and there may be no direct on-site release.

Complex transitions between media: The behavior of PFAS in the environment may deviate greatly from typical contaminants. Transitions between media may be complex because of specific characteristics of these compounds.

Partitioning: PFAS will bind to soils and sediments through hydrophobic, electrostatic, and interphase partitioning. However, binding of PFAS to organic carbon is not as strong as traditional hydrophobic compounds (for example, PCBs, PAHs), so Koc alone is likely a poor predictor of binding of PFAS to organic carbon. Care should be taken to obtain the most up to date Koc values. Given the significance of electrostatic or interphase partitioning mechanisms, it may be

Site Characterization Considerations and Media-Specific Occurrence for Perand Polyfluoroalkyl Substances (PFAS) *continued*

appropriate to evaluate those for many sites. Section 4 of the Guidance Document includes information about PFAS physical and chemical properties.

Balancing priorities: Because drinking water is a major pathway of exposure for PFAS and established plumes may have spread to downgradient areas, priority might be given to managing drinking-water exposure pathways over site characterization.

Monitoring-point construction: Investigators must be mindful of ways that monitoring-point construction may influence sampling results such as historical wells with fluoropolymer tape.

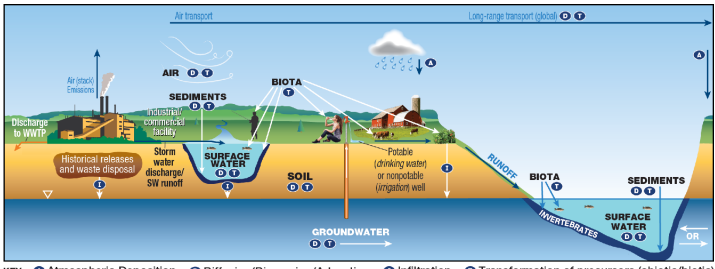
Widespread use: Because use of PFAS-containing products is widespread, there may be multiple sources in a given area that may act as additional or alternative sources of PFAS at a site. In addition, PFAS are often detected in low levels in samples from locations without an obvious source; there may be a need to evaluate site-specific anthropogenic ambient background to determine contributions in groundwater or soils that are not due to an on-site release. See, for example, Strynar et al. (2012).

Compound suite: Selection of a broad suite of compounds may prove useful for applications such as fingerprinting (if multiple sources are suspected) or for understanding potential effects of precursor degradation. Note that with currently available analytical methods, even a broad compound suite will not detect all PFAS.

Geologic heterogeneity: Because of the low regulatory limits that are used for delineation of groundwater impacts and the mobile nature of certain PFAS, extra emphasis should be placed on understanding the effects of hydrogeologic heterogeneity on the groundwater plume.

Initial Conceptual Site Model (CSM)

CSMs are useful tools for the presentation and evaluation of site characteristics, releases, contaminant fate and transport, and exposure pathways and can help focus a site investigation. An example generalized industrial CSM is presented in Figure 1.



KEY 🚯 Atmospheric Deposition 💿 Diffusion/Dispersion/Advection 🕕 Infiltration 🕕 Transformation of precursors (abiotic/biotic)

Figure 1. CSM for industrial sites.

Source: Adapted from figure by L. Trozzolo, TRC. Used with permission. PFAS-1, Figure 2-20.

The CSM presents most known and potential PFAS source areas, transport mechanisms, and pathways on a simplified physical setting. The CSM also illustrates exposure routes and receptors. CSM development is an iterative process over the duration of the project that will incorporate information obtained during site investigation, risk assessment, remedy design, and remedy implementation and optimization.

Some specific challenges related to identifying the nature and extent of impacts for CSM development for PFAS sites are cases of surface water creating very large dilute groundwater plumes through recharge (ATSDR 2008) and identification of potential ecological and human receptors in the context of risk-based evaluations for PFAS.

Site Characterization Considerations and Media-Specific Occurrence for Perand Polyfluoroalkyl Substances (PFAS) *continued*

Site Investigation

Development of Site Investigation Work Plan

PFAS investigation work plans should take into consideration the initial CSM and a comprehensive site preliminary assessment and incorporate applicable information and concepts from the Guidance Document. A few such items are sampling procedures and equipment to prevent cross-contamination, analytical methods, compounds to be reported, quality assurrance/quality control, geographically variable and changing regulatory requirements and criteria, site-specific environmental setting, human health and/or ecological risk assessment, and potential treatment technologies.

In addition, evaluation of the geologic and hydrogeologic framework associated with the site is critical, and sometimes that framework may need to be addressed on a regional basis, as the PFAS impacts may extend significant distances from the site.

Investigative techniques to characterize source soils and determine the three-dimensional extent of soil and groundwater contamination should be considered. High-resolution site characterization (HRSC) technologies, while providing near real-time subsurface lithological and hydrologic information, can be used to efficiently obtain discrete groundwater samples in the intervals of interest, such as the high transmissivity zones and lithological interfaces.

Key factors to consider in development of a work plan include: leaching from the vadose zone to the saturated zone; matrix diffusion; desorption; nonaqueous phase liquids (NAPL) dissolution; non-site sources; atmospheric deposition; overland runoff; groundwater discharge to surface water or surface water recharge of groundwater; suburface features, including utility lines; multicomponent mixtures; and the PFAS precursors that may be present.

As with other compounds, a site investigation for PFAS relies upon understanding the nature of sources as well as the lateral and vertical extents. At some sites, plumes of more mobile, shorter chain PFAS have been observed to be relatively more extensive than longer chain PFAS due to less retardation in groundwater. The partitioning behavior of PFAS is discussed in Section 5.2 of the Guidance Document. Further, comingling of contaminants has a potential to impact PFAS extent.

Data Analysis and Interpretation

Examples of factors and processes that need to be assessed by available approaches, methods, and tools that may be relevant to PFAS sites discussed in the Guidance Document are: retardation coefficients and travel time, mass flux/mass discharge, contributions from different sources, atmospheric deposition, vadose zone percolation, matrix diffusion, upgradient site contributions, transformation pathways and rates, assessing plume stability, modeling PFAS fate and transport, and visualization methods.

Forensics and Source Identification

Source identification is one of the challenges of PFAS investigations. Where there are no documented releases or there are possibly several comingled releases, multiple lines of evidence may be needed for source identification (also referred to as PFAS forensics). Source identification seeks to use the evaluation of both typical and advanced chemical analyses to potentially differentiate among contaminant sources and age-date release events. Advanced techniques continue to be developed including: chemical fingerprinting; signature chemicals; isotopic fingerprinting; contaminant transport models; molecular diagnostic ratios; radionuclide dating; and microscopic analysis.

3 Media-Specific Occurrence

PFAS occurrence in various environmental media is an active area of research. The material presented in the Guidance Document is not the result of an exhaustive literature review but is included to provide a relative understanding of PFAS concentrations typically found in environmental media. Section 6 of the Guidance Document includes figures summarizing the observed concentrations of PFAS that have been reported in the literature. Section 17.1 includes tables summarizing important details concerning each study used in developing the figures.

Air

Certain PFAS are found in ambient air, with elevated concentrations observed or expected in urban areas nearest to emission sources, such as manufacturing facilities, WWTPs, fire training facilities, and landfills (Barton et al. 2006 Ref#241; Ahrens et al. 2011; Liu et al. 2015 Ref#340). Although outdoor air containing PFAS can enter buildings, the presence of indoor sources can cause indoor air concentrations of certain PFAS to be higher than outdoor air concentrations (Fromme et al. 2015; Shoeib et al. 2011).

Site Characterization Considerations and Media-Specific Occurrence for Perand Polyfluoroalkyl Substances (PFAS) *continued*

Precipitation

PFAS have also been observed in precipitation and other related samples, including rainwater, surface and subsurface snow, sea ice, and meltwater. Concentrations of PFAS in precipitation vary over many orders of magnitude (Casas et al. 2021; Pike et al. 2020; Xie et al. 2020; Chen et al. 2019 Ref#2413; MacInnis et al. 2019; Wang et al. 2019 Ref#2415).

Soil, Sediment, and Biosolids

PFAS are found in soil and sediment due to atmospheric deposition, exposure to impacted media (for example, landfill leachate or biosolids), and direct releases. PFAS distribution in soil is complex, reflecting several site-specific factors such as total organic carbon (TOC), particle surface charges, and phase interfaces. Properties of individual PFAS, such as alkyl chain length and ionic functional group, are also important factors. PFOS, PFOA, and other long-chain PFCAs are typically the predominant PFAS identified in surface soils and sediments (Zhu et al. 2019; Rankin et al. 2016; Strynar et al. 2012). PFAS occurrence in biosolids is reported to be prevalent across the United States at a wide range of concentrations, and may be from municipal, residential, or industrial sources, or a combination of any of those sources (see ITRC's Biosolids and PFAS fact sheet found at https://pfas-1.itrcweb.org/fact-sheets/).

Groundwater

USEPA has assembled an extensive data set of the occurrence of six PFAAs in public drinking water through the monitoring of large drinking water systems under the UCMR3 program (USEPA 2017 Ref#933). One or more PFAAs were detected in 4% of the reporting public water systems (USEPA 2017 Ref#920). Groundwater sources had approximately double the detection rate of surface water sources (Hu et al. 2016). Groundwater occurrence data collected as part of domestic and international studies have also characterized the range of PFAS concentrations associated with release sites.

Surface Water

Freshwater, marine water, and stormwater PFAS concentrations usually depend on proximity to the point of release and source concentrations. In addition to releases associated with identified sources, stormwater runoff from nonpoint sources may contribute significant loads of PFAS to surface water (Wilkinson et al. 2017; Zushi and Masunaga 2009). The sorption of PFAS to suspended solids may affect surface water PFAS concentrations. Suspended microplastics may also influence PFAS in surface water (Llorca et al. 2018). In addition to PFOS and PFOA, many other PFAS have been observed in surface waters, including compounds other than PFAAs.

Biota

Because PFAS are widespread and have a propensity to bioconcentrate, they have often been found in fish, wildlife, and humans. PFAAs, particularly PFOS, are frequently the dominant PFAS detected in biota (Houde et al. 2011). PFAA concentrations in biota are influenced by uptake and elimination of both PFAAs and their precursors, as well as biotransformation rates of PFAA precursors (Asher et al. 2012; Gebbink, Bignert, and Berger 2016). Therefore, concentrations of PFAAs observed in biota at one location may not reflect concentrations in other environmental media.

4 References and Acronyms

The references cited in this fact sheet and further references can be found at <u>https://pfas-1.itrcweb.org/references/</u>. Reference numbers are included in this fact sheet for non-unique citations in the Guidance Document reference list. The acronyms used in this fact sheet and in the Guidance Document can be found at <u>https://pfas-1.itrcweb.org/acronyms/</u>.



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