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Bacterial transformation of per- and poly-fluoroalkyl substances: a review for the field of bioremediation

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Per- and polyfluoroalkyl substances (PFAS) have received growing attention as more research on their potential risk and widespread extent of contamination has become available. Remediation schemes are currently being developed to help mitigate exposure to PFAS, especially in aquifers underlying sites where extensive application of aqueous film forming foam (AFFF) for fire suppression and/or fire training has resulted in widespread PFAS contamination. To-date, microbial remediation (bioremediation) has not been considered for PFAS treatment due to the recalcitrant nature of these compounds, and the lack of identified organisms that can completely degrade PFAS to non-toxic by-products. However, based on recent literature, the role of microorganisms may be important to our understanding of processes occurring at contaminated sites where transformation of PFAS precursors to terminal end-products has been observed. This paper reviews the current state of knowledge on the biotransformation of PFAS by microorganisms and includes topics of particular interest to bioremediation schemes. First, we review studies linking the exposure of PFAS to microbial toxicity at the cellular level and then expand the viewpoint to observed changes at the microbial community level. Second, we review the different classes of PFAS observed to biotransform and provide an overview of reactions that are likely to occur under different environmental conditions. Third we evaluate the role of co-metabolism and specific enzyme classes in the observed biotransformation of larger PFAS to terminal perfluoroalkyl acids (PFAAs). Lastly, we identify knowledge gaps in the understanding of PFAS biotransformation and provide suggestions for future research that may result in new strategies that include bioremediation as a solution for PFAS in environmental matrices.

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1. Introduction

The past two decades have seen a growing awareness and interest in remediation schemes for compounds classified as per- and polyfluoroalkyl substances, collectively known as PFAS. PFAS are anthropogenic compounds synthesized and mass produced for their unique and advantageous properties including hydro- and lipo-phobicity, extreme heat resistance, and high surface-activity.¹ Their basic structure consists of a functional group that is typically charged ("head") attached to a fluoroalkyl backbone ("tail"), and this combination of the "head" and "tail" structure yields molecules that are surface active (*i.e.* surfactant), particularly at the air-water interface. Surfactant PFAS lower the surface tension of the media which allows ready formation of foams, leading to extensive use of PFAS in fire-fighting foams.² In addition to foams, PFAS have been used in a range of industrial applications and consumer

products. For example, they have been used as components in food packaging liners, stain resistant textiles and water-resistant goods (since PFAS repel both water and oil), as well as in cables, wires, gaskets and hoses, due to their stability under high temperatures. They are also used in engines as lubricants and coatings to protect parts from oxidation.³

PFAS are structurally diverse. For example, the head can be a range of functional groups including carboxylates, sulfonates, sulfonamides, phosphonates, and other larger moieties consisting of combinations of the previously listed groups and compounds such as hydrocarbons and amines, and the tail varies by length and degree of fluorination.⁴ It is also possible to have tails that include both fluoroalkyl and ether moieties, and these PFAS are commonly referred to as per- and polyfluoroalkyl ether acids (PFEAs; *e.g.*, GenX).⁵ Due to the extreme stability of the carbon-fluorine bond, many PFAS have been shown to be recalcitrant in the environment even if transformation of the head group is observed.⁶ Widespread use and transport *via* the ocean and atmosphere has led to global PFAS contamination.⁷⁻⁹ This, along with rising health concerns has led to incorporation of some PFAS in the Stockholm Convention list for Persistent

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Organic Pollutants and to voluntary phaseouts by major manufacturers.¹⁰ Recently, the United States Environmental Protection Agency (EPA) proposed stringent drinking water maximum contaminant levels (MCLs) for perfluorooctanoic acid (PFOA) and perfluorooctanesulfonic acid (PFOS) (4 ng L⁻¹) and a composite of perfluorononanoic acid (PFNA), perfluorohexanesulfonic acid (PFHxS), perfluorobutanesulfonic acid (PFBS) and GenX calculated based on their relative hazard indices.¹¹ There is a pressing need for viable methods of treatment and remediation as there are documented PFAS impacts to thousands of water supplies across the US and abroad.¹¹

To-date there has not been an all-encompassing solution for PFAS remediation, particularly with regards to legacy PFAS releases, such as those used for fire-training with aqueous film forming foam (AFFF), where decades of AFFF use have led to extensive soil and groundwater contamination. However, in recent years there have been promising advances in destructive technologies, such as photocatalytic systems with boron nitride and titanium dioxide, showing mineralization capabilities and more recently a non-photocatalytic system using dimethyl sulfoxide (DMSO) proving capable of mineralizing PFOA and GenX at near room temperature through decarboxylation reactions.¹²⁻¹⁵ Other destructive processes such as ball milling, super critical water oxidation, plasma, and thermal decomposition as well as non-destructive technologies such as adsorption onto activated carbon and removal *via* ion exchange resin have also shown promising results with a potential to be applied *ex situ*, and in a few cases *in situ* (e.g., adsorption to injected colloidal activated carbon or clay media).¹⁶⁻²⁰ In addition to physical and chemical removal and destruction technologies, recent studies suggest that biological processes may be just as important to consider for the overall remediation strategy of impacted sites and therefore are the focus of this review.

Bioremediation has been applied for treatment of a wide variety of compounds and is often the primary *in situ* remedial approach.²¹ However, the unique chemistry of PFAS makes enzymatic transformation to non-toxic end-products difficult. That being said, detoxification may only be one component of an overall remediation strategy. In many cases, PFAS must first be extracted from the subsurface before any remedial technique can be effectively applied. Increasing extraction efficiency of specific PFAS through transformation of these compounds, often called “perfluoroalkyl acid (PFAA) precursors” to more mobile, and sometimes smaller, anionic PFAAs could be a more suitable role for bioremediation in PFAS remediation schemes. This is similar to PFAS flushing approaches using oxidative approaches that have been studied at the lab scale.²² Understanding the role of biodegradation and bioremediation in tackling the PFAS problem is critical to effectively remediate sites in the future; therefore, understanding the current literature and knowledge gaps related to PFAS biotransformation is necessary. There have been several literature reviews on microbial transformation of PFAS²³⁻²⁶ and one on microbial mechanisms involved in transformation;²⁷ however, this review will uniquely focus on topics applicable to bioremediation schemes including:

- (1) The impact of PFAS on bacteria (*i.e.* observed toxicity and community shifts).
- (2) Currently established bacterial degradation pathways for different classes of PFAS.
- (3) Discussion on the role of co-metabolism in observed degradation including prominent enzymes thought to be responsible.
- (4) Future work based on identified knowledge gaps.

The goal of this review is to provide a concise update on the status of bacteria–PFAS interactions as it pertains to bioremediation and therefore aid in the dissemination of studies that show potential usefulness for bacteria in remediation schemes.

PFAS are a large class of compounds currently encompassing 12 000 different fluorinated molecules in the EPA CompTox master list at the time of writing. The large number of PFAS and the tendency of naming conventions to vary among different chemical companies and research groups has prompted us to provide a shortened list of PFAS relevant to discussions within the body of this review (Table 1), for acronyms from specific studies mentioned in tables, the reader is referred to the original text.

2. Effects of PFAS on microorganisms

When applying *in situ* bioremediation approaches, it is important to develop a conceptual site model. For example, understanding redox profiles, available electron donors (*i.e.* organic material in the soil, hydrocarbons from petroleum products, methane, inorganic donors such as ammonia), and relevant electron acceptors at a given site (such as oxygen, nitrate, sulfate, iron, and carbon dioxide) can help define the dominant bacterial communities and the degradative processes that these communities can mediate. Understanding toxicity impacts may play an important role for some sites where the targeted enzyme(s) or microbial process (e.g., nitrification, reductive dehalogenation) may be inhibited by PFAS contamination. The diversity of both microorganisms and PFAS make it impossible to study exact processes for every situation. However, observations on toxicity mechanisms and overall shifts in community structures due to PFAS exposure can help improve conceptual models of impacted systems. First PFAS toxicity effects on the cellular level will be discussed followed by more generally observed community shifts.

2.1 Cellular effects

The effects of PFAS on individual cells are relevant from a practical standpoint for understanding potential impacts on pollutant biodegradation (e.g., on reductive dehalogenation of chlorinated solvents) and from a theoretical standpoint for understanding which cellular characteristics contribute to PFAS toxicity or lack thereof. There have been several recent studies investigating the effects of PFAS on bacterial cells (Table 2). These effects include changes in cell morphology, surface charge, and membrane fluidity.

The current studies look only at a small subset of known PFAS and do not include any zwitterionic, fluorotelomer, or



Table 1 Primary PFAS discussed in the review

| PFAS structure | Full name | CAS | Acronym used |
|----------------|---|-------------|--------------|
| | Perfluorobutanoic acid | 375-22-4 | PFBA |
| | Perfluoropentanoic acid | 2706-90-3 | PFPeA |
| | Perfluorohexanoic acid | 307-24-4 | PFHxA |
| | Perfluorooctanoic acid | 335-67-1 | PFOA |
| | Perfluorooctanesulfonic acid | 1763-23-1 | PFOS |
| | 6:2 fluorotelomer sulfonate | 425670-75-3 | 6:2 FTS |
| | 6:2 fluorotelomer sulfonamide alkylamine | N/A | 6:2 FTAA |
| | 6:2 fluorotelomer sulfonamide alkylbetaine | N/A | 6:2 FTAB |
| | 6:2 fluorotelomer thioether amido sulfonate | 88992-47-6 | 6:2 FtTAoS |
| | 6:2 fluorotelomer unsaturated carboxylic acid | 70887-88-6 | 6:2 FTUAC |



Table 1 (Contd.)

| PFAS structure | Full name | CAS | Acronym used |
|----------------|--|-------------|--------------|
| | 8:2 fluorotelomer acrylate | 74049-08-4 | 8:2 FTAC |
| | 8:2 fluorotelomer methacrylate | 1996-88-9 | 8:2 FTMAC |
| | N-Ethyl perfluoroctane sulfonamido ethanol | 1691-99-2 | EtFOSE |
| | N-Ethylperfluoro-1-octanesulfonamide | 4151-50-2 | EtFOSA |
| | 2-(N-Ethyl-perfluoroctanesulfonamido) acetate | 909405-49-8 | EtFOSAA |
| | Perfluoroctanesulfonamide | 754-91-6 | FOSA |
| | Perfluoroctane sulfonamide acetic acid | 2806-24-8 | FOSAA |
| | Sodium p-perfluorous nonenoxybenzene sulfonate | 0829-87-7 | OBS |

sulfonamide compounds, which greatly limits their field applicability given the prevalence of these groups of compounds in historic AFFF used at impacted sites.³⁵ However, they do provide preliminary insight into mechanisms of toxicity, with the most commonly observed mechanism being membrane disruption. Membrane disruption has been observed through visual methods such as transmission electron microscopy (TEM) as well as using luminescence to detect the transfer of dye through the membrane in cell cultures.^{30,32} Naumann *et al.* took this a step further and specifically looked at the membrane interactions using isolated membranes from *Alcanivorax borkumensis* and model phospholipid compounds of both mono- and bi-layer membranes.²⁸ They performed compressibility tests and found that while PFAS did cause lipid

condensation, (*i.e.* caused the phospholipids to become more tightly packed), it still increased membrane fluidity in all cases. This finding is interesting as it implies that intramolecular forces may be disrupted by PFAS with some areas of the membrane being condensed while the others have increased fluidity. They also found a chain length and functional group dependent relationship where longer fluorinated chains were more likely to insert themselves and maintain association with the membrane as were sulfonates.

PFAS have also been observed to trigger stress responses in cells such as the production of extracellular polymeric substances (EPS).³³ EPS production is the key component in the development of biofilms and is thought to protect the cells from harmful conditions.^{36,37} Weathers *et al.*, noted stress induced



Table 2 Studies reporting PFAS linked to cellular toxicity^a

| Source | PFAS | Bacteria | Toxicity effect | Method of observation |
|--------|--|--|---|--|
| 28 | PFOA, PFNA, POS, PFHxS (0.01–50 mg L ⁻¹) | <i>Alcanivorax borkumensis</i> , synthetic mono-layer and bi-layer phospholipids | Increased membrane fluidity | Compressibility measurements |
| 29 | PFHxS (0.02 mg L ⁻¹) | <i>Pseudomonas</i> PS27 and PDMF10 | Morphology changes (rod to round) | TEM |
| 30 | PFBA, PFOA, PFNA, PFBS, PFHxS, PFOS (0.1–50 mg L ⁻¹) | <i>Aliivibrio fischeri</i> | Quorum sensing Increased cell membrane permeability | Luminescence measurements |
| 31 | PFBA, PFOA, PFNA, PFBS, PFHxS, PFOS (0.01–50 mg L ⁻¹) | <i>Staphylococcus epidermidis</i> , <i>Aliivibrio fischeri</i> DC43 | Partitioning into membrane | QCM-D, zeta potential |
| 32 | PFOA, PFOS (0.1–500 mg L ⁻¹) | <i>Escherichia coli</i> | PFOA-DNA oxidative stress PFOS-membrane disruption | TEM, FTIR, viability assays, RT-PCR, SOD assay, zeta potential |
| 33 | PFBA, PFPcA, PFHxA, PFHpA, PFOA, PFNA, PFDA, PFUnA, PFBS, PFHxS, PFOS (0.1, 0.2, 0.5, 1, 2, 5, 10 mg L ⁻¹) | <i>Rhodococcus jostii</i> RHA1 | Increased EPS production, increase in 2 stress-related genes (<i>sigF3</i> , <i>pfmA</i>) | qRT-PCR, optical density EPS assay, light microscopy |
| 34 | PFOS (0.1, 1, 5 mg L ⁻¹) | Thermophilic anaerobic sludge | Increased EPS production, increased ROS levels, increased membrane permeability | EPS assay, ROS assay, luminescence measurements |

^a TEM, Transmission Electron Microscopy; QCM-D, Quartz Crystal Microbalance with Dissipation Monitoring; qRT-PCR, Quantitative Reverse Transcription Polymerase Chain Reaction; RT-PCR, Reverse Transcription Polymerase Char Reaction; FTIR, Fourier transform infrared spectroscopy; SOD, Superoxide Dismutase; EPS, Extracellular Polymeric Substances; ROS, Reactive Oxygen Species.

changes in *Rhodococcus* RHA1 when exposed to a mixture of eleven PFAAs, including a doubling of cell length, likely due to either EPS production causing cells to remain together after division or incomplete cell division, as well as enhanced biofilm production.³⁸ The specific mechanism of toxicity was studied more in-depth by Liu *et al.* who differentiated between the effects of PFOA and PFOS on *Escherichia coli* cells.³² Exposure to PFOS increased membrane fluidity to a greater extent than PFOA which is consistent with the above discussion of membrane affects in which longer chain sulfonates were linked to greater membrane disruption.³² PFOA's main route of toxicity appears to be oxidative damage, especially to DNA, and overall PFOA had a higher observed toxicity than PFOS for *E. coli*.³² In addition to toxicity effects, PFOS and PFOA may alter the hydrophobicity of the cells themselves: at the lowest dose tested (10 mg L⁻¹) these PFAS appear to stimulate production of porins and lipopolysaccharides, making cell surfaces more hydrophobic.³² It is worth noting here that the lowest doses tested in this study are orders of magnitude higher than typical environmental concentrations in groundwater which average around 100 ng L⁻¹ with high values typically below 10 µg L⁻¹.³⁹ At higher doses (PFOS: 500 mg L⁻¹; PFOA 15 mg L⁻¹), PFAS can bind to the membrane's hydrophobic sites, leaving the hydrophilic head to interact with the surroundings, and potentially changing the cell charge.³² Significant zeta potential changes were not observed until PFAS exceeded 50 mg L⁻¹, which is consistent with Fitzgerald *et al.* who found no zeta potential differences in live cells (of both gram-positive and gram-negative bacteria) at concentrations up to 50 mg L⁻¹.^{31,32} However,

Fitzgerald *et al.* did observe differences in zeta potential with their phospholipid model membrane (separate from actual cells) and theorized that the cell wall prevented the surface charge changes since the PFAS associated with the membrane and the cell wall remained unchanged.³¹ Exploring this idea of cell wall and PFAS interactions, a preliminary study on the difference between gram-negative (lacking an outer layer of peptidoglycan) and gram-positive bacteria (having an outer layer of peptidoglycan) in relation to sorption of PFOA and PFOS indicated that the gram-positive bacteria sorbed more PFAS.⁴⁰ This leads to some interesting questions on the role of the cell wall and the presence of peptidoglycan in interactions with PFAS. More research is needed to better understand these interactions.

While these studies cannot be directly applied to all types of cells and PFAS that may be present at a contaminated site, they do provide a good framework for understanding the effect of some PFAS on specific strains of bacteria. These studies also demonstrate that toxicity varies by compound, and it is likely to also vary by microorganism, potentially leading to appreciable shifts in microbial community structure as discussed in the following section.

2.2 Microbial community effects

As PFAS contamination is widespread and may affect natural systems with diverse microbial communities, it is of interest to know how microbial ecology changes with PFAS exposure. These changes may reflect toxicity of specific PFAS (*i.e.*, with some species, and those that depend on them, declining in



density), but they also may lead to a better understanding and identification of strains with the ability to biotransform PFAS. These PFAS-transforming organisms may increase in the affected communities, particularly if they gain carbon, energy or nutrients during PFAS biotransformation. To understand community shifts, several studies have performed 16S rRNA surveys on samples from a variety of different sites (Table 3). These studies have shown several consistent trends in community shifts. This section first discusses bacteria observed to increase in numbers after PFAS exposure and then those observed to decline with PFAS exposure.

2.2.1 Bacteria increasing with PFAS exposure. Overall, the phylum Proteobacteria (gram-negative), one of the most genetically varied and widely present phyla, increased in nearly all cases with PFAS exposure.^{41–43,47,48,54} More specifically, Gammaproteobacteria was found as a dominant class in many systems with PFAS. This corresponds with findings from Kim *et al.* where strains found to transform 6:2 FTOH were all Gammaproteobacteria. In general, microbes that participate in the sulfur cycle, such as the classes Alphaproteobacteria, Verrucomicrobia, and Gammaproteobacteria, show an increase in abundance with sulfur-based PFAS present.^{47,51} Studies have shown possible metabolic desulfonation with the fluorotelomer sulfonates and their precursors indicating sulfur cycling could

be positively correlated with biotransformation of 6:2 fluorotelomer sulfonate (6:2 FTS).^{47,55} This will be addressed further in the biodegradation section of this review.

2.2.2 Bacteria declining with PFAS exposure. In contrast to the sulfur cycle, where the number of sulfur cycling organisms increased with the presence of PFAS, the nitrogen cycle may be inhibited as the numbers of nitrifiers decreased after introduction of PFAS in wetland sediments (6:2 FTS) and activated sludge (PFOS).^{4,43,54,56} Ke *et al.* studied a community of ammonia oxidizing archaea and ammonia oxidizing bacteria and found that the archaea dominated when exposed to sodium *p*-perfluorous nonenoxybenzene sulfonate (OBS).⁵² The overall bacterial community and nitrification rates were adversely affected by OBS.

It is important to put these studies into context of environmental sites and recognize that although laboratory studies have shown impacts on microbial communities, PFAS may be significantly less important than general environmental parameters such as pH and oxidation-reduction potential (ORP) at sites with long term exposure.⁵⁷ Further analysis of the metadata generated by these studies is needed before any firm conclusions are possible on the effect of PFAS on community structure. It is also important to recognize that 16S rRNA gene surveys have inherent bias that selects for microorganisms that are already

Table 3 Community shifts due to PFAS exposure^a

| Source | System | PFAS | Dominated | Decreased |
|-------------------|-------------|--|---|---|
| (41) ¹ | SBR | PFOS | ^a Proteobacteria, Bacteroidetes | Chloroflexi, Actinobacteria |
| (41) ¹ | SBR | PFOS | ^b Gammaproteobacteria, Bacteroidia, Alphaproteobacteria | Anaerolineae, Actinobacteria, and Chloroflexia |
| (42) ³ | AS | PFOA (20 mg L ⁻¹) | ^a Bactroidetes, Proteobacteria, Acidobacteria | Planctomycetes |
| (43) ² | WS | 6:2 FTS | ^a Proteobacteria, Acidobacteria, Gammaproteobacteria | Planctomycetes, bacteroidetes |
| (43) ² | WS | 6:2 FTS | ^d (<i>a</i>)Methylocaldum (Gammaproteobacteria) | SM1A02 (Planctomycetes), Ohtaekwangia, Taibaiella (Bacteroidetes) |
| (44) ² | AD | AFFF (DCP + PFOS) | N.D. | Decrease in methanogenic activity |
| (45) ² | AD | PFOS (500 mg L ⁻¹) | N.D. | No decrease in methanogenic activity |
| (46) ² | WAS, AS, FC | PFOA (10 mg L ⁻¹) | ^b Burkholderiaceae | Hyphomicrobium |
| (46) ² | WAS, AS, FC | PFOA (10 mg L ⁻¹) | Acidimicrobium sp. Strain A6 | N.D. |
| (46) ² | WAS, AS, FC | PFOA (10 mg L ⁻¹) | ^d Dechloromonas, Desulfobulbus, Desulfosporosinus, Desulfovibrio | N.D. |
| (47) ¹ | AW | PFOS/PFOA (2–600 ng L ⁻¹) | ^a Bacteroidetes, Chloroflexi | N.D. |
| (47) ¹ | AW | PFOS/PFOA (2–600 ng L ⁻¹) | ^b Gammaproteobacteria, Flavobacteriia | N.D. |
| (47) ¹ | AW | PFOS/PFOA (2–600 ng L ⁻¹) | ^c Rhodobacterales | N.D. |
| (48) ¹ | FWP | PFOA (0.45–5 mg L ⁻¹) | ^a Proteobacteria and Chloroflexi | Actinobacteria, baceroidetes, Verrucomicrobia |
| (48) ¹ | FWP | PFOA (0.45–5 mg L ⁻¹) | ^b Betaproteobacteria | N.D. |
| (49) ³ | RS | PFOA (8–465 ng L ⁻¹) | ^d Thiobacillus, Sulfurimonas | N.D. |
| (50) ¹ | LL | 6:2 FTS (650 µg L ⁻¹) | ^a Acidobacteria, actinobacteria | N.D. |
| (51) ¹ | AS | PFOS (0.01–1 mg L ⁻¹) | ^b Bactroidetes, Verrucomicrobia | Betaproteobacteria |
| (51) ¹ | AS | PFOS (0.01–1 mg L ⁻¹) | ^d Comamonas | Zooglea, Pseudomonas, |
| (52) ¹ | GS | OBS (1–100 mg kg ⁻¹) | Ammonia oxidizing archaea | Ammonia oxidizing bacteria |
| (53) ¹ | WS | PFBA, PPFA, PFHxA, PFHpA, PFOA, PFHxS, PFBS, PFOS (0.2–4.2 µg/L) | ^a Proteobacteria, Verrucomicrobia Cyanobacteria | Actinobacteria |

^a 1. Dose-response, 2. Exposure-response, 3. Existing contaminated system; a. phylum, b. class, c. order, d. genus; SBR sequencing batch reactor (wastewater treatment), AS activated sludge, WS wetland sediments, AD anaerobic digester, WAS waste activated sludge, FC filter cake, AW Antarctic waters, FWP freshwater pond, RS river sediment, LL landfill leachate, GS grassland soil. N.D. no data reported.



characterized both in the sample preparation steps and in the bioinformatics analysis. So, although PFAS does appear to induce community structure change, especially at a lab-scale, more in-depth studies linking community composition, PFAS concentration, and other environmental factors are needed.

3. Microbial transformation of PFAS

The next section of this review focuses on observed biotransformation of PFAS by different bacterial communities and individual strains. Identifying degradation pathways can help inform our understanding of what occurs once PFAS is released into the environment from both a forensic and predictive standpoint. Our understanding of microbial transformation of PFAS has evolved over the past twenty years, as is often the case for emerging contaminants.⁵⁸ Initially there were conflicting reports on whether or not PFAS biotransformed at all. In the early 2000s, studies showed the expected environmental half-life varying from 10 to 1000 years for the same fluoroacrylate polymer.^{59,60} One study indicated that PFOA was biologically inert, while others showed significant transformation of the functional head groups of perfluorinated compounds produced by electrochemical fluorination (ECF) and polyfluorinated compounds produced from the fluorotelomerization process (FT) leading to the idea that only transformation of the functional group, and not the fluorinated tail, was likely.^{61,62} In more recent studies, instances of defluorination for PFOA and the fluorotelomer alcohols (FTOHs) and transformation of FT-precursors has been observed as well as increased reports of a number of ECF-precursor functional groups undergoing bacterial transformation. Further discussions of FT-PFAS degradation pathways, ECF-PFAS degradation pathways, and terminal PFAA defluorination follow and are based on the studies cited in Table 4.

3.1 FTOH biotransformation

One of the first classes of PFAS found to biotransform, and therefore a commonly studied one, is the N:2 FTOHs, specifically 6:2 and 8:2 FTOH, which have partially fluorinated chains with a CH_2 group at the end of the fluorinated tail. N:2 FTOHs, such as 6:2 FTOH (Fig. 1), generally transform through a series of intermediates into either PFOA, shorter chain carboxylates (PFBA, PFPeA, PFHxA) or saturated acids, through what was termed the “one carbon removal pathway”.^{97–101} During N:2 FTOH biotransformation, the unsaturated carboxylic acid (N:2 FTUCA) appears to be a key intermediate from which different pathways branch.⁹⁷ From N:2 FTUCA, one pathway produces acids and the other produces ketones.^{100,102} The acid pathway releases two CF_2 while the ketone pathway only releases one.^{100,102} It has been noted that the pathway used can be influenced by the addition of specific co-substrates (e.g., lactate), opening possibilities for influencing the type of end-products that are produced from N:2 FTOH degradation and possibly allowing for selection of acid products instead of ketones.^{100,102} There remains some ambiguity concerning the proposed metabolic pathway from 6:2 FTOH to PFBA, which

may reflect differences in bacterial culture; but overall the acid pathway continues to produce acids until it branches off into a ketone pathway, which produces terminal end-products.^{70,103} It is also interesting to note that the soil culture used in Liu *et al.* appeared to add an amide group to 5:2 FTUCA producing 5:3 Uamide, a reaction that currently appears unique to that study.¹⁰⁴

3.2 FTOH precursor degradation

There are many PFAS identified that have a FT structure (*i.e.* partially fluorinated carbon chain) and larger complex functional groups. These “FT-precursors”, especially those with sulfonamide functional groups, can undergo cleavage of the ester bond and transform to FTOHs (Fig. 2), and then proceed along the established degradation pathway for FTOHs into the terminal short chain perfluorinated carboxylic acids (PFCAs) and other ketone and acid intermediates discussed above.^{43,69,70} These precursor studies have included soils (unsaturated), sediments (saturated), activated sludge, and pure cultures. Each of these microbial systems will be discussed in more detail below.

3.2.1 Soils. Aerobic soils typically host a large variety of microorganisms and are commonly contaminated with PFAS at areas with AFFF use. Aerobic soils have been shown to be particularly conducive to FT-precursor transformation, although the responsible organisms are largely unknown. Recently a large FT precursor, 6:2 Fluorotelomer thioether amido sulfonate (6:2 FtTAoS), was observed to completely transform within a week to PFCAs in aerobic soils under aerobic conditions and with an enriched soil culture where previously it had been shown to transform within 120 days under nitrate-reducing conditions and over 270 days for sulfate-reducing conditions.^{65,71,72} Other amine containing FT-precursors, 6:2 FTAA and 6:2 FTAB showed slow but significant biotransformation in aerobic soils collected from a biopile constructed for remediation of petroleum hydrocarbon contamination.⁷⁴ Lastly, smaller FT-precursors such as 8:2 fluorotelomer acrylate (8:2 FTAC) and 8:2 fluorotelomer methacrylate (8:2 FTMAC) were found to biotransform with differences in rates based on the functional group and soil type (*i.e.* 8:2 FTAC degraded slower than 8:2 FTMAC and degradation was faster in forest soil than grassland soil).⁶⁴

3.2.2 Sediments. Subsurface sediments in lakes, rivers, and wetlands have also been shown to have bacterial communities capable of transforming FT-precursors. Several studies have evaluated 6:2 FTS transformation in sediments under both anaerobic and aerobic conditions. Significantly higher levels of 6:2 FTS transformation have been reported under aerobic conditions, especially when sulfur is limiting^{43,69} (Fig. 2). One possible explanation for this increased transformation is that monooxygenases (which add a singlet oxygen to an organic substrate) are partly responsible for the observed reactions. This hypothesis is supported by increased abundance of the genus *Methanomethylovorans* (a methanotroph) under aerobic conditions after exposure to 6:2 FTS, possibly implicating methane monooxygenase in the transformation of 6:2 FTS.⁴³ However, in



Table 4 Studies of microbial PFAS biotransformation (n.d. = no data; R = room temperature)

| Category | Source | Starting compound | Organism | Bio-transformation products | Transformation efficiency | Experiment length/temp (°C) | Half-life reported | Starting concentration |
|---------------|--------|---------------------------------|---|---|-------------------------------------|-----------------------------|-------------------------------------|--|
| FT precursors | 63 | FTU and HMU | Aerobic soils (both agricultural land and forest) | PFOA, PFHxA, and PFHxA, 7:2 s FTOH, 8:2 FTOH | Forest soil (24–27%) | 180d (22) | FTU: 126–148d, HMU: 478–667d | 100 µg g ⁻¹ |
| | 64 | 8:2 FTAC and FTMAC | Aerobic soils | PFOA, FTOH | 10.3% mol production of PFCA | 105d (R) | 8.2 FTAC <5d and FTMAC 15 d | 75–100 µg g ⁻¹ |
| | 65 | 4:2 6:2 and 8:2 FTTAOs, FTTAOs, | Aerobic soil from AFFF site | 4:2, 6:2, 8:2 FTs, 6:2 FTUCA, 5:3 FTCA, C4–C8 PFCA. Newly characterized: 6:2 and 8:2 FTSOAs and FTSO2As (4 total) | >95% yield to intermediates | 60d (30) | n.d. | 26 µM 6:2 FTTAOs, 0.04 µM 8:2 FTTAOs and 0.005 µM 4:2 FTTAOs |
| | 66 | 6:2 FTAA and 6:2 FTAB | Activated sludge | 6:2 FTOH, 6:2 FTCA, 6:2 FTUCA, 5:3 FTCA, PFHxA, PFPA, PFBA | 16 and 6 mol% quantifiable products | 109d (R) | 419d FTAA and 1817d FTAB | 600 µg/400 mL |
| | 55 | 6:2 FTAB, 6:2 FTAA | <i>Gordonia</i> sp. strain NB4-1Y | *degradation was not tracked, gene expression instead was measured | NR (30) | NR (30) | 60 µM | |
| | 67 | 6:2 FTOH, 4:2 FTOH, TFMAs | Mixed landfill soil culture (aerobic) | 5:3 FTCA, PFHxA, PFPA, PFBA, TFA, TFA | ~100% for FTOHs and >7% for TFMAs | 32d (R) | 1.5d 6:2 FTOH and 300 ppb for TFMAs | 10 ppb for FTOHs and 300 ppb for TFMAs |
| | 43 | 6:2 FTS | Aerobic and anoxic wetland sediments | 5:3 acid, PFHxA, PFPeA | Aerobic 8.9%. anaerobic, 0.7 mole% | 120d (30) | n.d. | 160 µg/20 mL for aerobic and 100 µg/20 mL for anoxic |
| | 50 | 6:2 FTS, 8:2 FTOH | Landfill leachate | PFBA, PFPeA, PFHxA, PFHxA, PFOA | 60d (20) | n.d. | 20 mL for anoxic | |
| | 68 | 6:2 FTSA | <i>Rhodococcus jostii</i> RHA1 | 6:2 FTCA, 6:2 FTUCA, α -OH 5:3 FTCA | >3 log reduction | 6d (30) | 650 µg L ⁻¹ | 40 µM |
| | 69 | 6:2 FTSA | Sediments (aerobic and anaerobic) | 5:3 acid, PFPeA, PFHxA | 98.9% aerobic, 0% | 100d (R) | <5d aerobic | 1.3 mg L ⁻¹ |
| | 70 | 6:2 FTSA and 6:2 FTAB | <i>Gordonia</i> sp. strain NB4-1Y | PFBA, PFPeA, PFHxA, 4:3 FTCA, 5:3 FTCA, 6:2 FTCA, 6:2 FTUCA, 5:2 FT ketone and 6:2 FTOH | 99.9% FTSA and 70.4% FTAB | 7d (30) | n.d. | 60 µM FTSA and 60 µM FTAB |
| | 71 | 6:2 FTTAOs | Anaerobic AFFF impacted soil | 6:2 FTTP | 75% | 270d (30) | n.d. | 20 µM |
| | 72 | 6:2 FTTAOs | Aerobic soil culture | 6:2 FTs, PFHxA, PFPeA, PFBA | 100% | 20d (R) | 3.81d to 6.65d | 12–16 µM |



Table 4 (Contd.)

| Category | Source | Starting compound | Organism | Bio-transformation products | Transformation efficiency | Experiment length/temp (°C) | Half-life reported | Starting concentration |
|----------|---|---|---|---|---------------------------|------------------------------|---|------------------------|
| 45 | PFOs, 6:2 FTSA, PFBS, TFA, PFPA, PFPeA, 3 H-PFPtA, trifluoropropionic acid, 2,2-bis(trifluoromethyl)propanoic acid, 5H-PFPtA, trifluoropentanoic acid | Aerobic and anaerobic wastewater sludge | n.d. | n.d. | 1241d (R) | n.d. | 200–500 ppm | |
| 73 | 6:2 PAPS | <i>Pseudomonas oleovorans</i> , <i>Pseudomonas butanovora</i> , and <i>Pseudomonas fluorescens</i> DSM 8341 | 6:2 FTtOH, 6:2 FTtCA, 6:2 FTtUCA, 5:2 ketone, 5:2 SFtOH, PFHxA, 5:3 acid, 5:3 uacid | 8.6 mol% per gram of cells | 30d (30) | n.d. | 6.9 ppm of 6:2 FTtOH equivalents | |
| 74 | AFFF, 6:2 FTtAB | Soil from site contaminated from AFFF and from a crude oil spill | PFtCS, PFHxA, PFHtPA, PFPtPeA, PFtBA, 5:3 FTtCA, 6:2 FTtUA, 5:2 ketone, 6:2 FTtSA | Reduction in total measured PFAS: 24% | 60d (R) | 31d | PFtOS: <2100 ng g ⁻¹ , PFtOA: <850 ng g ⁻¹ , 6:2 FTtAB: 10 µg g ⁻¹ | |
| 75 | 6:2 FTtAOtS | Aerobic and anoxic soils | 6:2 FTtS, PFHxA | 84% removal | 282d (30) | n.d. | 35 µM | |
| 76 | Cationic quaternary ammonium compounds (QACs): PFOAAmS and PFOSAmS | Non-impacted area close to AFFF site | PFOA | 30 mol% yield to PFOA. | 180d (22) | PFOAAmS: 142d, PFOSAmS: n.d. | 1 µg g ⁻¹ | |
| 77 | EtFOtSA | Non-AFFF-impacted soil | PFtOS, FOSA, FOSAA, EtFOtSAA | 4% yield to PFtOS | 182d (22) | 13.9 ± 2.1 d | 3.12 nmol g ⁻¹ | |
| 78 | EtFOtSA | Carrot microcosm | FOSAA, FOSA, PFtOS | 81% depletion EtFOtSA | 80d (25) | 35.8 ± 3.7 d to 40 ± 7.8d | 3.8 mg kg ⁻¹ | |
| 79 | EtFOtSE | Aerobic soils (acidic, alkaline) | EtFOtSAA, EtFOtSA, FOSAA, FOSA, PFtOS, EtFOtSA, EtFOtSAA, FOSA, PFtOS | (1.06, 5.49) mol% >99% | (180, 210)d (22) | (26, 31)d | 3.7 µg g ⁻¹ | |
| 80 | EtFOtSE | St. Bernard soil | FOSA, PFtOS | | 105d (22) | 8.7d branched, 9.6d linear | 4.72 nmol g ⁻¹ of EtFOtSE, sum of branched and linear. | |
| 81 | MeFBSE and EtFOtSE | Anaerobic digester communities | MeFBtSAA and PFBtS; EtFOtSAA and PFtOS | 75% depletion MeFBtSAA and 3% depletion EtFOtSE | 108d (20–26) | 35.8d MeFBtSAA 1860d EtFOtSE | 1 ppm | |



Table 4 (Contd.)

| Category | Source | Starting compound | Organism | Bio-transformation products | Transformation efficiency | Experiment length/temp (°C) | Half-life reported | Starting concentration |
|---|--------|--|---|---|---|-----------------------------|--|---|
| | 82 | PFOANO and PFOSNO | Aerobic soils | PFOA and PFOS | PFOA 15–21 mol%, PFOS: 2 mol% | 90d (25) | PFOANO: 3–7d, PFOSNO: 15d | PFOANO: 800 ng g ⁻¹ , PFOSNO: 2000 ng g ⁻¹ ~2 µg g ⁻¹ |
| | 83 | PFOANO, PFOAAmS, PFOAB, PFOSNO, PFOSAmS, PFOSB | Aerobic soils | FOSAA, FOSA, PFOS, PFOA | 0.06–32.6% | 90 d–150d (22) | 14–675d | |
| | 84 | AmPr-FHxSA | Aerobic soil enrichments | FHxSA-Pra, FHxSA, PFHxS | 30% | 70d (30) | n.d. | 0.2 µM |
| PFAAs transformation in complex systems | 85 | PFOA | Activated sludge-enriched, photocatalytic, synergistic system | n.d. | 79.7% removal | n.d. (25) | 4d | 500 µg L ⁻¹ |
| | 47 | PFOS | Antarctic ocean | n.d. | 50% | 6d (R) | 43h for PFOS | 2–600 ng L ⁻¹ |
| | 86 | PFOS and PFOA | Chemorganoheterotrophic bacteria and total yeast and molds | n.d. | 46–69% PFOS depletion and 16–36% PFOA depletion | 28d (28) | 20.3d PFOS and 38.9d PFOA | 1 µg mL ⁻¹ for total PFAAs |
| | 87 | PFOS and PFOA | Activated sludge | n.d. | 15–42% | 1d (22) | 1.2d | 100, 250, 500 ng L ⁻¹ |
| Complex precursor mixes | 88 | PFAS released from carpet | Methanogenic community | Unable to conclude. | n.d. | 552d (37) | n.d. | n.d. |
| | 57 | AFFF | 13.7 times higher PFAS measured in biotic than abiotic treatments | n.d. | n.d. | n.d. (n.d.) | n.d. | n.d. |
| | 89 | AFFF impacted soil | Soil samples taken from impacted site for community analysis | 8:2 FTS with increase of PFOA, PFHxA, PFAAs | n.d. | 1.5d (R) | n.d. | n.d. |
| Terminal end-product transformation | 90 | PFOS | Aerobic and anaerobic soil columns | <i>Pseudomonas</i> n.d. | 75% | 6d (28) | 3d PFOS pure culture, 3 months in soil | 0.1 w/v% |
| | 91 | PFOA and PFOS | <i>plecoglossicida</i> 2.4-D | PFBA, PFPeA, PFHxA, PFHpA | 60% | 100d (30) | n.d. | 0.24 mM PFOA and 0.2 mM PFOS |
| | 46 | WWTP biosolids (with PFOA) | <i>Acidimicrobium</i> sp. strain A6 | PFBA, PFPeA, PFHxA, PFHpA | 68% | 150d (25) | n.d. | 0.2–10 mg L ⁻¹ PFOA |
| | 92 | PFOA | <i>Acidimicrobium</i> sp. strain A6 | n.d. | 48.1% | 2.5d (30) | n.d. | 500 mg L ⁻¹ |
| | 93 | PFOA | <i>Pseudomonas</i> <i>parafilosa</i> YAB1 | n.d. | 58.6% | 4d (30) | n.d. | 500 mg L ⁻¹ |
| | | | <i>Pseudomonas</i> <i>parafilosa</i> YAB1 – gene shuffled | | | | | |

Table 4 (Contd.)

| Category | Source | Starting compound | Organism | Bio-transformation products | Transformation efficiency | Experiment length/temp (°C) | Half-life reported | Starting concentration |
|----------|------------------|---|--|-----------------------------|---------------------------|--|---------------------|------------------------|
| 94 | PFCAs | <i>Dehalococcoides</i> , KB-1, mixed culture | Defluorinated PFCAs | ~100% | 150d (R) | n.d. | 75 μ M | |
| 95 | PFMeUPA, FTMeUPA | <i>Dehalococcoides</i> , KB-1, mixed culture | Hydrogenated and defluorinated transition products | >90% | n.d. (34) | 130d | 75 μ M | |
| 96 | PFOS and PFOA | <i>Ensifer adhaerens</i> strain M1 | n.d. | 100% | 6d (26-28) | 3d for PFOS and 2d PFOA | 1 g L ⁻¹ | |
| 90 | PFOS | <i>Pseudomonas</i> sp. <i>plecoglossicida</i> 2.4-D | n.d. | 75% | 6d (28) | 3d PFOS pure culture, 3 months in soil | 0.1 w/w% | |

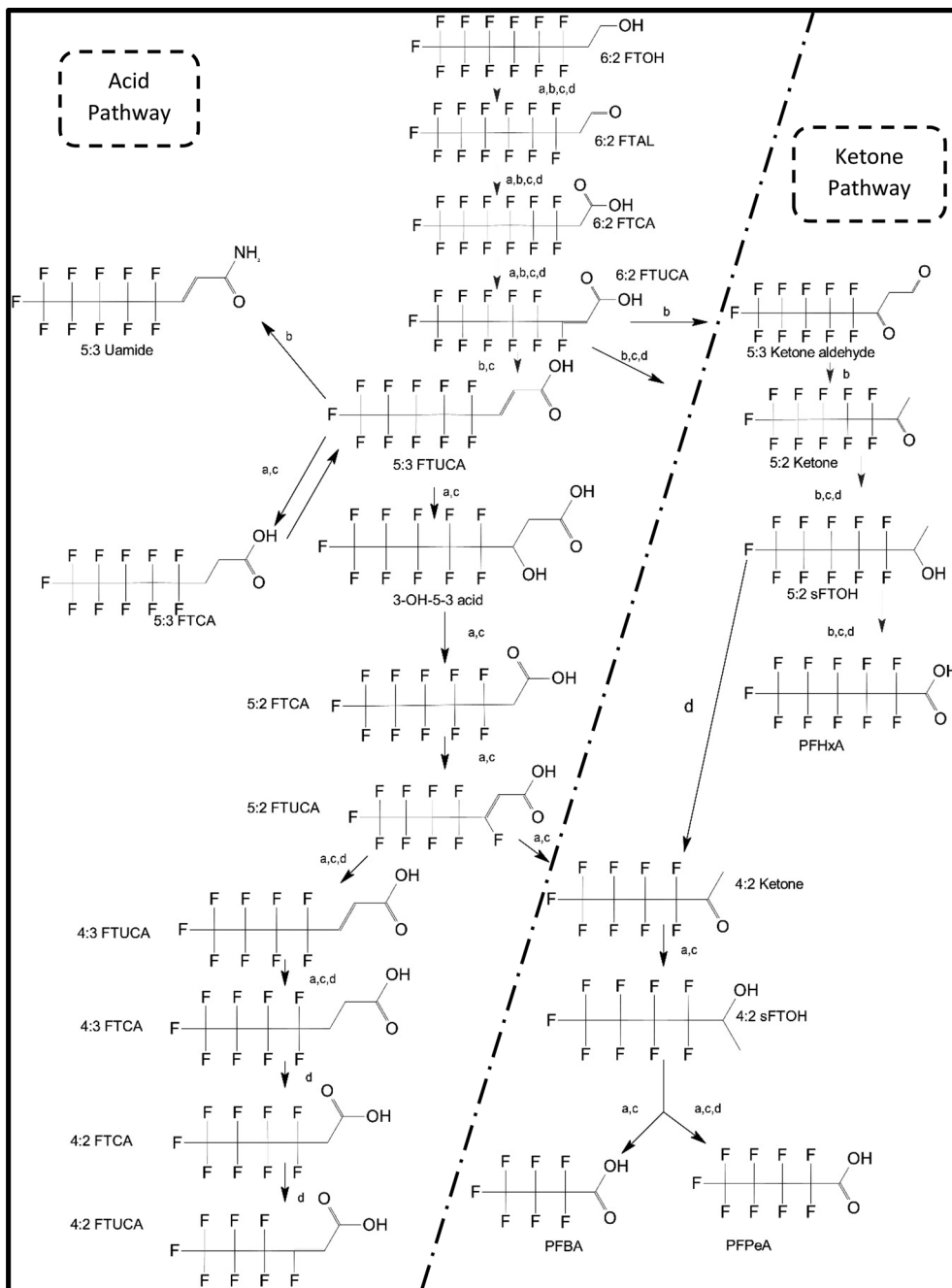
separate studies under anaerobic conditions, Yin *et al.* observed biotransformation indicating enzymes other than monooxygenases are able to transform 6:2 FTS, especially under sulfur-limiting conditions.⁴³ Overall, both aerobic and anaerobic sediments were shown to support transformation of these FT-precursors. This transformation was slower in all cases studied than initial transformation of 6:2 FTOH.⁶⁹ The rate differences were attributed to 6:2 FTS possessing a strong electron withdrawing group (sulfonate) while 6:2 FTOH possesses an electron donating group (hydroxyl).^{43,105}

3.2.3 Activated sludge. Activated sludge from wastewater treatment plants is a popular matrix for microbial studies as this environment selects for bacteria that are able to degrade diverse substrates. Therefore, when the biodegradability of PFAS was first explored, many studies looked at activated sludge systems.^{103,106,107} While activated sludge is not strictly relevant to most bioremediation schemes, the reactions, and the bacteria that perform them, can provide insights that may be useful for other environments. Moreover, in that sludges from wastewater treatment frequently contain PFAS and are often spread onto agricultural or other lands, transformation reactions in this environment are important in their own right. In recent years, biotransformation reactions of 6:2 fluorotelomer sulfonamide alkylamine (6:2 FTAA) and 6:2 fluorotelomer sulfonamide alkylbetaine (6:2 FTAB), which were previously discussed in the soils section, have also been observed in sludges; they generally transform at comparable rates to those observed in soils.^{66,108}

3.2.4 Pure cultures. Lewis *et al.* examined the transformation of larger FTOH precursors known as polyfluoroalkyl phosphates (PAPs) using activated sludge and three bacterial strains (*Pseudomonas oleovorans*, *P. butanovora*, and *P. fluorescens* DSM 8341) that had previously been shown to transform FTOHs.¹⁰² All were able to transform PAPs to different degrees, and various substrates were found to induce different pathways. Other pure culture studies evaluated the ability of *Gordonia* sp. NB4-1Y, isolated from vermicompost (compost degraded using worms), to transform 6:2 FTS and 6:2 FTAB.⁷⁰ Biotransformation reactions with this strain occurred at a much faster rate than observed in soils and activated sludge (7 days *versus* over 60 days).^{66,70,74,109} It was observed that both 6:2 FTSA and 6:2 FTAB did produce some of the same metabolites but at different rates (6:2 FTAB metabolized slower than 6:2 FTSA).⁷⁰ 6:2 FTS was also degraded by *Rhodococcus jostii* RHA1 under sulfur-reducing conditions, and it was found that defluorination followed desulfonation as suggested in earlier studies.^{68,108} Overall, FTOH precursor transformation seems to be performed by both bacteria with monooxygenases and bacteria with sulfur-reducing capabilities under varied environmental conditions.

3.3 ECF precursor degradation

Unlike FT compounds, ECF compounds have a fully fluorinated carbon chain making them significantly more resistant to biological transformation. They have only been observed to undergo transformation of the functional head group and not defluorination (Fig. 3). Of these functional head transformations, the most studied pathway starts with *N*-ethyl

Fig. 1 6:2 FTOH biotransformation pathway. a¹⁰² b¹⁰⁴ c¹⁰⁰ d⁷⁰

perfluorooctane sulfonamidoethanol (EtFOSE). A common daughter product produced from EtFOSE is *N*-ethylperfluoro-1-octanesulfonamide (EtFOSA), the main ingredient in Sulframid, an ant killer frequently used in Brazil to reduce leaf cutting ant populations on agricultural lands.¹¹⁰ The application of this insecticide has been linked to high levels of PFOS (a potential daughter product of EtFOSA) in Brazilian surface water.^{78,110} In addition to ant killer, these sulfonamide-ECF compounds are common ingredients in AFFF formulations, especially in many 3M formulations from 1988–1998, lending an even greater relevance to understanding their biotransformation pathways for impacted soils and groundwaters.³⁵

Biotransformation pathways have been proposed for several of the larger sulfonamide ECF compounds shown in Fig. 3. Many of these pathways converge at EtFOSA and then proceed to FOSAA, FOSA, and finally, PFOS. Cook *et al.* recently reported that the C6 homologue to the C8 sulfonamide PFOSAm, AmPr-FHxSA, transformed to PFHxS in aerobic soil enrichments amended with carbon substrates, including diethyl glycol monobutyl ether (DGBE), benzene, toluene, ethylbenzene, and xylene (BTEX), acetate, and methane.⁸⁴ AmPr-FHxSA was observed to follow a similar transformation path as the C8 compounds, but with fewer identified intermediates. Demethylation, acetylation, and hydroxylation are all proposed

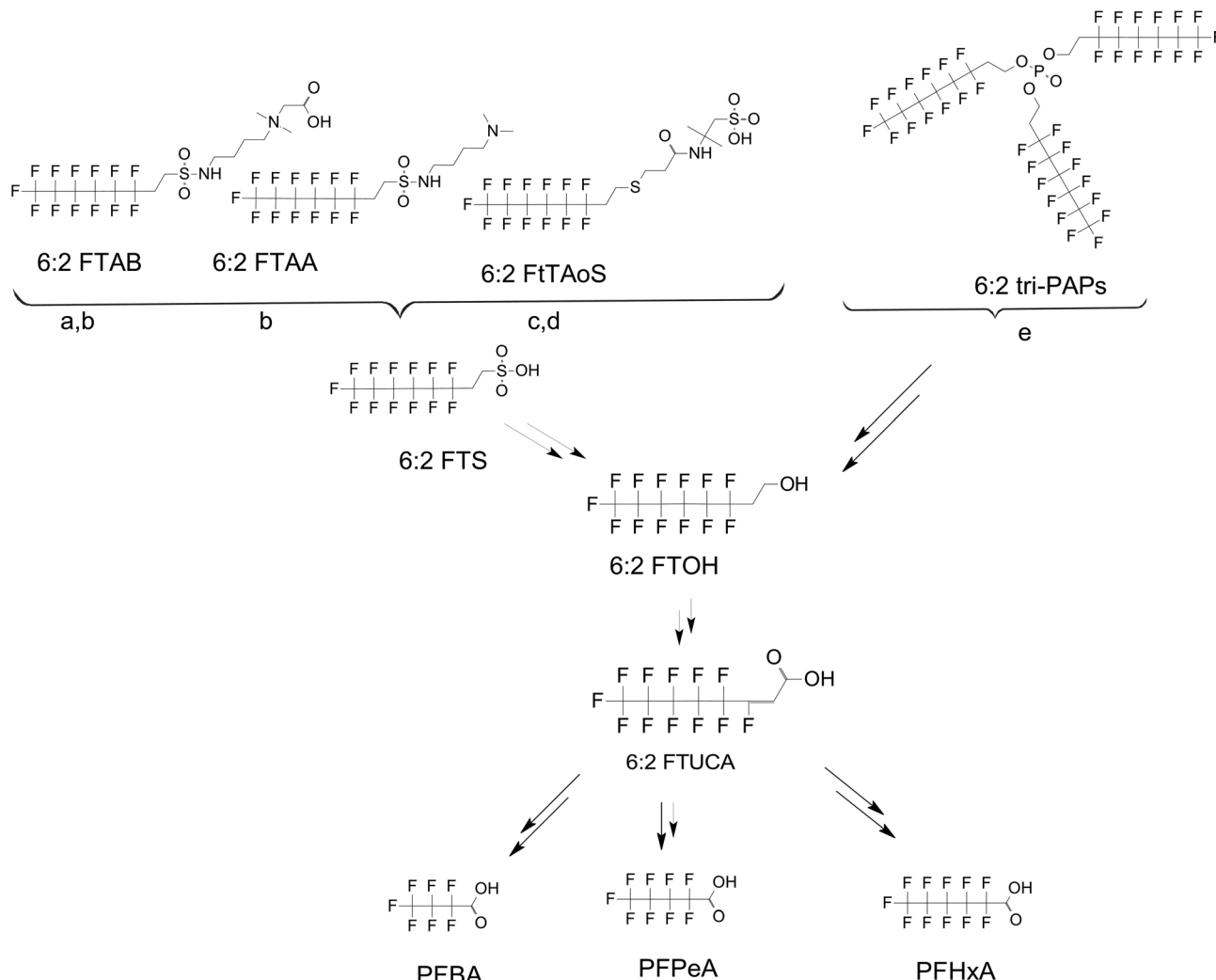


Fig. 2 Transformation of 6:2 FTS and precursors a,⁶⁶ b,⁷⁰ c,⁷² d,⁷¹ e⁷³ double arrows indicate multiple transformation steps.

enzymatic mechanisms for transformation of these compounds. Interestingly, several of these studies have proposed a direct transformation step from larger precursors (EtFOSE, PFOSAm, PFOSB intermediates formed by hydroxyl and demethylation) to PFOS, and this has subsequently been supported with modeling.^{80,111}

The biotransformation pathways that result in terminal PFAAs have been of particular interest because of the regulatory focus on PFOA and PFOS specifically, and the high concentrations of these compounds found at contaminated sites.¹¹³ When looking at the transformation pathways, it is worth noting that only the linear isomers are shown here but there are typically two isomers of EtFOSE, and likely the other larger precursors as well, one linear (n) and one branched (br), and therefore all of the transformation products also have two isomers: linear isomers degrade to linear transformation products and branched to branched products.⁸⁰ It was initially thought that the branched isomers would biotransform faster; however, several studies have reported the opposite: the linear EtFOSE isomer and transformation products, with the exception of

perfluorooctanesulfonamide (FOSA), are more readily transformed by bacteria.^{78,80} This is opposite of the metabolic preference in high-order organisms such as fish.¹¹⁴ Although progress has been made over the past decade, the biotransformation pathways of ECF compounds need significantly more study. We need a better understanding of transformation reactions to terminal PFAAs and knowledge concerning reactions that may result in defluorination.

3.4 PFOS and PFOA biotransformation

Most of the current literature focuses on the biotransformation of precursors to terminal end-products in environmental samples and microbial consortia. However, specific strains of bacteria have also been reported to defluorinate perfluorinated moieties such as PFOA and PFOS.^{90,92,115,116} The first strain to be identified as capable of breaking the carbon-fluorine bond in PFAS was *Pseudomonas fluorescens* D2, which was able to use the recalcitrant polyfluorinated H-PFOS as a source of sulfur; however, it was unable to transform PFOS itself (*i.e.*, fully saturated with fluorine).¹¹⁶ Alpha oxidation was theorized to be

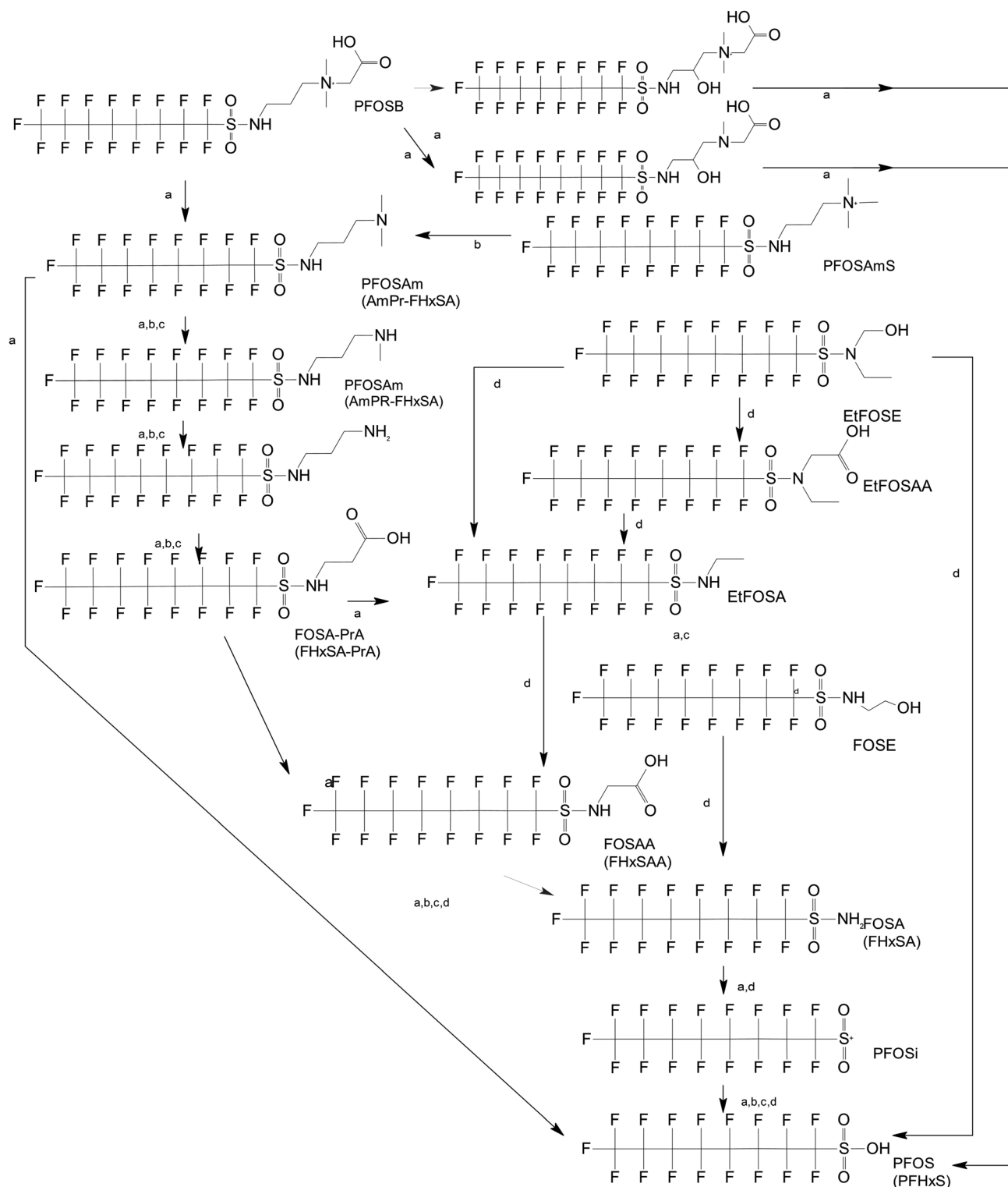


Fig. 3 ECF Pathway. a⁸³ b⁷⁶ c¹¹² d¹¹¹ names in parentheses indicate C6 homologues observed in ref. 84.

the transformation mechanism of H-PFOS.¹¹⁶ The researchers studying *P. fluorescens* D2 were also the first to propose a link between sulfur metabolism and defluorination.¹¹⁶

A few bacteria have been reported to be capable of using PFOS and/or PFOA as substrates resulting in degradation of

these PFAAs. Such strains were isolated from contaminated material such as soil from old factory sites or activated sludge from wastewater treatment plants with high PFAS concentrations. These environmental sources were used as the original inocula followed by a series of isolations using PFOS as the

substrate to isolate *Pseudomonas aeruginosa* strain HJ4, *Ensifer adhaerens* strain M1, and *Pseudomonas plecoglossicida* strain 2.4-D.^{90,96,115} Only one strain (*Pseudomonas parafulva* strain YAB1) to-date has been isolated using PFOA instead of PFOS.⁹² YAB1 was subsequently put through a series of gene shuffling and re-isolation procedures to modify the strain to have improved degradation efficiency.⁹³ Although these papers show promising results, they are the only ones to report organisms able to grow solely on PFOA and/or PFOS.

A strain not isolated originally on PFOS or PFOA, but showing evidence of defluorination of both of these PFAAs is *Acidimicrobium A6*. This bacterium was originally isolated based on its capability to grow using the feammox reaction, oxidizing ammonium (NH_4^+) to nitrite (NO_2^-) while simultaneously reducing ferric iron ($\text{Fe}(\text{III})$) to ferrous iron ($\text{Fe}(\text{II})$).^{46,91,117} The degradation of PFOA and PFOS observed with A6 is thought to be co-metabolic (see next section), similar to its degradation of other contaminants such as perchloroethylene (PCE).⁹¹ In order to confirm defluorination, higher concentrations of PFAS were used than would commonly be found in the environment (*i.e.*, 0.24 mM and 0.2 mM for PFOA and PFOS respectively).⁹¹ It is unclear yet whether this organism will be relevant for *in situ* treatment of PFAS. However, recent work by this group has shown that PFOA in biosolids could be partially defluorinated when incubated with both $\text{Fe}(\text{III})$ and strain A6.⁴⁶ The enzymatic mechanism for defluorination of PFOA and PFOS has yet to be elucidated, however a model was published linking defluorination to ammonia oxidation.¹¹⁸ It is also worth noting that PFOS desulfurization was theorized to have occurred as well as defluorination, but the source of sulfate in samples could not be positively identified as originating from PFOS/PFOA.⁹¹

In addition to isolating individual strains, three studies have reported the transformation of PFOS and PFOA in mixed culture or in combination with other technologies.⁸⁵⁻⁸⁷ In these studies, a decrease in PFAA concentration in mixed communities was observed, however, evidence for microbial transformation was not confirmed. Other possible mechanisms for the observed mass reduction could be sorption or, in the case of Beskoski *et al.*, the use of dimethyl sulfoxide (DMSO) as a solvent for PFOA, as DMSO has recently been shown to degrade PFOA at low temperatures.^{15,86}

4. Co-metabolic degradation of PFAS

The majority of studies seem to implicate a co-metabolic degradation mechanism for the biotransformation of PFAS. This section discusses co-metabolism in the context of bioremediation and evaluates its role in the observed biotransformation of PFAS.

4.1 What is co-metabolism?

Co-metabolic reactions are most commonly catalyzed by “promiscuous” enzymes that have binding sites with broad substrate specificity such that they perform both growth-linked and non-growth-linked reactions.¹¹⁹ Historically, many different oxygenase enzymes (*e.g.*, methane monooxygenase, ammonia

monoxygenase, toluene dioxygenase) have been observed to co-metabolize a range of xenobiotic compounds, which differ based on the organism/enzyme.

During co-metabolic transformation of a pollutant, microbial growth is typically dependent on the primary growth substrate (*e.g.*, methane or propane) rather than the contaminant. As such, this process can sometimes be applied as a bioremediation strategy to treat contaminants that are present at low environmental concentrations (*e.g.*, $\mu\text{g L}^{-1}$) but above relevant regulatory standards or toxicity thresholds.

For this reason, co-metabolism research has recently focused on a variety of contaminants that occur at low concentrations in environmental matrices, such as 1,4-dioxane (stabilizer in chlorinated solvents), *n*-nitrosodimethylamine (disinfection and liquid rocket fuel byproduct), 1,2,3-trichloropropane (solvent and pesticide manufacturing byproduct), and a broad array of pharmaceuticals, estrogens, and personal care products found in wastewater.¹²⁰⁻¹²² Often, these compounds are present in the environment at concentrations too low to support microbial growth causing co-metabolic processes to be the primary path of removal. Since these and other recalcitrant, and often halogenated, compounds are broken down *via* co-metabolic pathways, the possibility to use co-metabolism to remediate PFAS should be explored.

4.1.1 Current evidence of PFAS co-metabolic biotransformation. As previously discussed, microbial biotransformation has been most widely explored and reported for fluorotelomer alcohols, (4:2, 6:2 and 8:2 FTOH in particular). In many cases, observed transformation mechanisms of FTOHs are likely to be co-metabolic in nature, but there remains a lack of understanding of the enzymes involved.^{23,73,99,100,102} For example, many of the FTOH studies have observed that communities and cultures could not grow solely on FTOHs suggesting the process to be co-metabolic.^{99,100} However, Lewis *et al.* observed that polyfluoroalkyl phosphates (PAPs), a class of FTOH precursors, were used as primary substrates for the initial degradation step and therefore concluded the degradation of PAPs was not purely co-metabolic.⁷³

In an effort to find useful patterns among the studies that have provided insight for how PFAS are biotransformed (Table 5), several trends are evident and mapped here: (I) defluorination has been observed only through co-metabolic processes, (II) larger precursors with complex functional groups have been observed to transform potentially through both metabolic and co-metabolic pathways, and (III) desulfonation appears to be a driving factor for the metabolic transformation of precursors in sulfur-limited environments.

Expanding on each of these three claims, first there is no convincing evidence of defluorination proceeding metabolically. Yu *et al.* observed defluorination and theorized a *Dehalobacter* strain may be responsible but were unable to identify a specific organism or show growth on PFAS.⁹⁴ The same holds true for the feammox process as it requires the primary substrates of ammonium (or hydrogen) as the electron donor and ferric iron as the electron acceptor in order to observe PFOA and PFOS defluorination.^{46,91} Defluorination by the FTOH one-carbon-removal pathway, observed in activated sludge and



Table 5 Co-metabolism potential reported in literature

| Source | PFAS | Notes on Co-metabolism in study |
|--------|-----------------------|--|
| 62 | 8:2 FTOH | Replenishment of organic carbon enhanced microbial mineralization of multiple $-CF_2-$ groups in the fluorocarbon chain of ^{14}C -8:2 FTOH. |
| 102 | 4,6,8:2 FTOH | Presence of formate increased FTOH transformation 6-fold, cannot grow on just FTOH |
| 99 | 8:2 FTOH | Ethanol, octanol, 1,4 dioxane used as carriers and carbons sources, could not grow on just FTOH |
| 100 | 6:2 FTOH | Supports co-metabolism, all strains studied used other carbon sources |
| 99 | 8:2 FTOH | Used 1,4, dioxane, ethanol and octanol. FTOH could not support growth by itself, concluded it is a co-metabolic process |
| 68 | 6:2 FTOH | Defluorination observed and alkane monooxygenases implicated as possible co-metabolic enzymes responsible |
| 73 | 6:2 FTOH | Showed dependency on co-substrate supporting co-metabolic mechanism |
| 72 | 6:2 FtTAoS | Found that BTEX amendments increased transformation of PFAS |
| 123 | HPFOS | Used glucose/acetate to partially degrade H-PFOS |
| 124 | 6:2 FTSA | Different carbon sources used and defluorination observed, haloacid dehalogenase implicated for defluorination activity |
| 70 | 6:2 FTAB and 6:2 FTSA | Used acetate as the carbon source, observed defluorination after desulfonation in media where PFAS were the only sulfur source. |
| 91 | PFOA | Feammax, NH_4^+ and Fe(III) needed for defluorination |
| 95 | PFCAAs | Unable to isolate defluorinating organism, TCE as substrate |
| 73 | PAPs | Used same organisms as. ¹⁰⁰ The enzyme inducers resulted in less biotransformation of PAPs implying initial transformation is metabolic. |
| 84 | AmPr-FHxSA | Studied soil enrichments ability to transform AmPr-FHxSA with different carbon substrates, methane, acetate, DGBE, BTEX, methane and acetate produced highest amounts of transformation products |

aerobic soils, was shown to be dependent on primary substrates in pure culture studies, additionally supporting co-metabolic defluorination.^{101,102}

Second, the functional groups of PFAS have been observed to transform both metabolically and co-metabolically with the latter appearing more commonly in the literature.^{72,100,124} Instances of metabolic transformation appear to be limited to very large functional groups such as 6:2 PAPs, where the initial transformation step is metabolically mediated and consecutive transformations appear to switch back to either co-metabolism, where the organism is not able to grow on the compound, or to metabolic desulfonation as described below⁷³

Third, desulfonation appears to be the main mechanism for metabolic transformation of sulfur-containing precursors under sulfur-limiting conditions (Fig. 1). Several studies have observed that 6:2 FTSA is desulfonated only when there is no other source of sulfur, strongly suggesting that 6:2 FTSA is used as the sole sulfur source for some bacterial strains.^{70,124} In addition to precursor transformation, the sulfur cycle has also been linked to PFOS desulfonation in Antarctic ocean communities, further strengthening the importance of sulfur cycling to PFAS transformation.¹²⁵ Overall, metabolic sulfur use and co-metabolism both appear to play a large role in PFAS biotransformation, and studies are ongoing to isolate the relevant enzymes and mechanisms, so these transformation reactions can be more fully understood. Some of these studies are reviewed in the subsequent sections.

4.2 Potentially responsible enzymes

There have been several reviews on the enzymes potentially responsible for PFAS biotransformation reactions and more generally cleavage of the carbon–fluorine bond.^{27,126,127} Current

research has focused on two main areas: co-metabolic and metabolic degradation using oxygenase enzymes and reductive dehalogenation using dehalogenase enzymes which are typically dependent on reduced corrinoid cofactors.

4.2.1 Co-metabolic degradation using oxygenase enzymes.

There has been a growing interest in oxygenase enzymes and their ability to transform PFAS by catalyzing reactions with PFAS functional groups.^{25,124} As discussed previously, the primary reason for the interest in oxygenases is their high *promiscuity index*, which indicates they often have the ability to catalyze reactions with compounds other than their primary substrates.¹¹⁹ Oxygenase enzymes linked to the co-metabolic defluorination of 6:2 FtTAoS include cytochrome P450, toluene dioxygenase and alkane monooxygenase.^{68,72}

In addition to oxygenase enzymes carrying out co-metabolic reactions, they have also been linked to metabolic degradation *via* desulfonation.^{55,68,109} In particular, the alkanesulfonate monooxygenase system has been upregulated in desulfonation of 6:2 FTSA and 6:2 FTAB with strains *Gordonia* sp. strain NB4-1Y and *Rhodococcus jostii* RHA.^{55,68,109} Within the class of alkanesulfonate monooxygenases, nitrilotriacetate monooxygenases and flavin-dependent monooxygenases (FMN) have been implicated for carrying out the desulfonation of PFAS.^{55,57} Additionally the *alkB* gene, which codes for oxygenases, was found to be a marker correlated with the PFAS transformation potential of systems.⁷³ There have been a number of studies that have theorized on the role of oxygenase enzymes but none have explicitly studied them.^{62,70,90}

4.2.2 Reductive defluorination using dehalogenase enzymes.

There has been widespread use of the reductive dechlorination pathway for the remediation of chlorinated solvents.¹²⁸ Since chlorine and fluorine are both halogens, it was



originally theorized that reductive defluorination might be a viable process to remediate PFAS; however, initial results showed that perfluorinated compounds were highly resistant to such transformation reactions.¹²⁹ The difficulty with using fluorocarbons for reductive defluorination can be attributed, in part, to the strong electronegativity of fluorine resulting in a redox potential on the edge of what is considered energetically favorable for microorganisms (-450 mV).^{27,130} Because of this, adenosine triphosphate (ATP) is consumed in order to break the C-F bond rather than generated as with chlorinated compounds.²⁷ Until recently, there were few studies that observed any reductive defluorination. Instead, it was observed that PFAS hindered the reductive dechlorination process.³⁸ However, a recent study by Yu *et al.* reported that reductive defluorination of unsaturated C₆ per- and polyfluorinated compounds was observed in a mixed dechlorinating culture, but that *Dehalococcoides mccartyi*, the key species responsible for the complete reductive dechlorination of chlorinated ethenes, was not responsible.⁹⁴ *Dehalobacter*, in conjunction with other unknown bacterial genera, was theorized to catalyze the defluorination, but difficulty isolating defluorinating strains points to a co-metabolic mechanism.⁹⁴ Many of the enzymes involved in dehalogenation reactions (dechlorination and debromination) have been found to be dependent upon reduced low-potential cobalt corrinoid cofactors.²⁷ It is likely that a reduced corrinoid, specifically B12, as well as a reducing agent, such as Ti(III) or HS⁻, are required during the reported enzymatic defluorination of branched poly-fluorinated compounds.^{131,132} The same enzymatic reactions do not defluorinate similar linear polyfluorinated PFAS.^{131,132} Overall, the enzymatic mechanism(s) of PFAS defluorination are not well understood and remain the subject of intense investigation. This research will likely determine the long-term potential for PFAS bioremediation.

5. Conclusions and recommendations for future work

Over the past two decades understanding of microbial degradation of PFAS has progressed from the conclusion that most PFAS were biologically inert to the idea that biologically mediated biotransformation and even biodeflourination are possible. While progress has been made in documenting PFAS biotransformation and understanding processes already occurring at contaminated sites, the potential for bioremediation to serve as a viable technology for treating PFAS in the environment remains unclear. There are many data gaps that must be filled to better understand this potential. Importantly, with few exceptions, the observed biotransformation reactions of PFAS to date do not result in the complete defluorination and/or detoxification of the parent molecules. In many instances, precursor compounds are transformed to terminal PFAAs, which are likely to persist based upon current knowledge. As noted in the introduction to this review, the microbial conversion of precursors to PFAAs could be part of an overall

remedial strategy at sites as these compounds tend to be more readily extractable from groundwater and other environments.

One of the more critical needs is to both reproduce and develop a clearer understanding of PFAS defluorination, as reported for strain A6 and a handful of other organisms and microbial enrichments. The development of *in situ* bioremediation strategies for PFAS destruction and detoxification depends largely on the capacity of bacteria (or other organisms including archaea and fungi) to defluorinate PFAS. Isolating a variety of organisms that carry out these reactions under different geochemical conditions, and then understanding the reactions themselves, is likely the key to determining whether bioremediation can be a viable strategy for PFAS treatment as it is now for so many other contaminants of concern. When defluorination reactions are better understood, studies can then be conducted to determine potential approaches to enhance this activity at contaminated sites (e.g., adding amendments, changing REDOX, bioaugmentation) and to assess the true potential for this process. Defluorination may also be influenced by both the relatively low concentrations of PFAS at most sites (*i.e.*, limiting growth-linked reactions but perhaps supporting co-metabolic ones) and the diverse array of PFAS often present, as compounds may undergo different reactions. Once these reactions are more clearly understood, there is also the potential for genetically engineering strains specifically for PFAS defluorination.

Author contributions

Jessica LaFond: Writing-original draft. Kayleigh Millerick: Writing-review and editing. Paul Hatzinger: SERDP funding, writing-review and editing. Jennifer Guelfo: Writing-review and editing. Andrew Jackson: Conceptualization, Writing-review and editing.

Conflicts of interest

The authors declare no conflict of interest associated with this review.

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