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Linear perfluoroalkyl carboxylate reduction dynamics with solvated electrons from ferrocyanide and sulfite†

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The initial reduction dynamics of perfluoroalkyl substances (PFASs) by the hydrated electron, e_{aq}^- , is a topic of great interest and importance due to the pervasive environmental presence of PFAS in soil and waters and the need to remediate the contamination. Understanding how PFAS behaves in water, including the potential effect of aggregation on the apparent PFAS reduction rate constant (k_{PFAS}) is, therefore, of paramount importance. In this publication we re-examine the proposition that submicellar aggregation decreases the apparent k_{PFAS} for sodium perfluorocarboxylates of varying chain lengths (NaPF_xA, x = number of carbons in the PFAS backbone, ranging from 4 to 8) using transient absorption spectroscopy. We compare the dynamics for e_{aq}^- quenching by NaPF_xA in aqueous solutions of ferrocyanide, $Fe(CN)_6^{4-}$, and sulfite, SO_3^{2-} . The results demonstrate that the apparent rate constant depends on the choice of e_{aq}^- precursor. We demonstrate that the ionic strength of the solution and the counterion of the PF_xA salt both affect the measured rate constant of PFAS reduction by e_{aq}^- . The results presented here help to better understand PFAS degradation by advanced reduction processes.

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Environmental significance

Advanced reduction processes (ARP) have proven to be quite effective at the destruction of PFAS and other contaminants in tainted waters. However, the processes underlying the reduction of PFAS by hydrated electrons (formed by the UV-excitation of appropriate ions capable of carrying out ARP, like sulfite) are still poorly understood; particularly, the early processes related to the initial reduction of PFAS. Until now, studies utilizing fast spectroscopic techniques like nanosecond transient absorption spectroscopy to elucidate the dynamics between the hydrated electron and PFAS have used ferrocyanide instead of sulfite as the electron donor for a variety of reasons with the assumption that the electron precursor has little effect on the reaction between the hydrated electron and PFAS. In this study we demonstrate that this assumption may not hold true and explore ionic strength effects, as well as counterion effects, to try to pinpoint the factors that make the fast reaction occurring in solutions of ferrocyanide and sulfite so vastly different. We argue that perfluoroalkyl carboxylates (PF_xA) tend to aggregate in water at concentrations well below the critical micelle concentration. Furthermore, the concentration of sulfite needed to effectively carry out ARP for practical degradation of contaminated waters shifts the PF_xA equilibrium towards aggregation. The implication is that the [PF_xA] used in typical degradation studies found in the literature are primarily in some aggregated state. Therefore, the degradation models proposed thus far require revision to include the aggregation of PF_xA and the role of aggregation in modulating reactivity with hydrated electrons.

Introduction

Groundwater contamination by poly- and perfluoroalkyl substances (PFAS), such as perfluorooctanoic acid (PFOA in the literature, PF8A here), has garnered significant attention due to their potential adverse health effects.¹ Historically, PFASs were used as particularly effective components in firefighting foams,² however their ubiquitous use now extends beyond fire suppression³ for their exceptional thermal and electrochemical

stability arising from the strength of the C–F bond (*e.g.* ~ 127 kcal mol⁻¹, 5.5 eV for C₂F₆).⁴ Not coincidentally, decontamination efforts aimed at removing PFAS from the environment have proven challenging due to these same chemical properties.⁵

UV-irradiation of solutions containing sulfite (SO_3^{2-}) under alkaline conditions (pH > 8) efficiently degrades PFAS by producing hydrated electrons, e_{aq}^- , that are photodetached from SO_3^{2-} .^{6–8} This process has become the prototype of so-called ‘Advanced Reduction Processes’ (ARP) for high energy reductive decontamination of tainted waters. The large standard reduction potential (-2.9 V)⁹ of the e_{aq}^- is nearly isoenergetic to the potentiometric biases required for the defluorination reaction of many perfluoroalkanes (*e.g.* $E_{PF8A}^\circ \sim 2.5$ V vs. SHE and $E_{PF8S}^\circ < 3.2$ V vs. SHE).^{10–14}

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† Electronic supplementary information (ESI) available. CCDC 2279639 contains the supplementary crystallographic data for the crystal structure of NaPF8A in Fig. S7 in the supplemental. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d3va00223c>



However, most of the work reporting on the fast reaction dynamics of the reduction of PF8A by e_{aq}^- has been done in solutions of ferrocyanide ($\text{Fe}(\text{CN})_6^{4-}$) rather than SO_3^{2-} .^{15–18}

$\text{Fe}(\text{CN})_6^{4-}$ is a well-studied precursor for generating solvated electrons *via* photodetachment.^{19–23} It is an attractive alternative to SO_3^{2-} for modelling the behaviour of e_{aq}^- quenching by PFAS, particularly in the absence of Brønsted acids. Unlike SO_3^{2-} ($\text{p}K_{\text{b}} = 6.8$), $\text{Fe}(\text{CN})_6^{4-}$ is aprotic at $\text{pH} > 4.3$ and has a 100-fold larger molar extinction coefficient (ϵ) at the 254 nm excitation wavelength (SO_3^{2-} $\epsilon_{254\text{nm}} \sim 50 \text{ M}^{-1} \text{ cm}^{-1}$; $\text{Fe}(\text{CN})_6^{4-}$ $\epsilon_{254\text{nm}} \sim 5000 \text{ M}^{-1} \text{ cm}^{-1}$).^{24–28} Ferrocyanide also has a high photodetachment quantum yield, even in the near UV.²³ In particular, the quantum yield of e_{aq}^- formation (Φ_e) from $\text{Fe}(\text{CN})_6^{4-}$ approaches unity as the excitation wavelength decreases (SO_3^{2-} $\Phi_e \sim 0.1$ at 254 nm excitation, $\text{pH} 8$; $\text{Fe}(\text{CN})_6^{4-}$ $\Phi_e \sim 0.65$ at 254 nm excitation, $\text{pH} 8$).²⁸

SO_3^{2-} and $\text{Fe}(\text{CN})_6^{4-}$ are not unique in their ability to undergo photoinduced ionization to produce e_{aq}^- . Other photochemically active anions also produce e_{aq}^- and have been used in ARP for the degradation of PFAS. Examples include ethylenediaminetetraacetate (EDTA^{4-}),²⁹ ionic indole derivatives,³⁰ as well as iodide (I^-) solutions that produce e_{aq}^- in both the absence and presence of SO_3^{2-} .^{31–34}

We previously reported that the apparent rate constant measured for e_{aq}^- quenching by PF8A can be underestimated at high concentrations (above $\sim 1 \text{ mM}$) due to surfactant aggregation¹⁸ including micellization above $\sim 30 \text{ mM}$. Aggregation of PF8A results in a decrease of the monomeric quencher concentration and appears to reduce the apparent e_{aq}^- quenching rate constants.¹⁸

As mentioned above, the majority of transient studies probing PFAS reduction dynamics by the e_{aq}^- have been carried out in solutions of $\text{Fe}(\text{CN})_6^{4-}$ due to the reasons already outlined. This assumes that the initial reduction of PFAS by e_{aq}^- is independent of the source of the e_{aq}^- . However, this assumption omits other factors that could potentially modulate e_{aq}^- reactivity with PFAS. Due to the differences in the photophysical properties of the various electron donors and the solution conditions required to optimize the degradation of contaminants by ARP, it is imperative to characterize the reduction dynamics of PFAS in each case. In this report, we challenge the supposition that the dynamics of ARP are independent of the e_{aq}^- donor, find clear differences between processes carried out in solution of $\text{Fe}(\text{CN})_6^{4-}$ *versus* SO_3^{2-} , and present evidence that suggest that these differences are due to differences in the aggregation state of the PFAS induced by the respective solution conditions. The evidence presented here suggests that these differences are, in part, due to ionic strength effects that affect not only the apparent quenching rate constants resulting from well-known screening effects, but also promote submicellar aggregation of PFAS at low concentrations. In addition to the transient absorption studies used to characterize the quenching rate constants, concentration-dependent UV-visible absorption spectra were measured and provide corroborating evidence of aggregation.

Experimental

All chemicals were used as received without further purification. Sodium sulfite (Na_2SO_3 , 98%) was obtained from Alfa Aesar. Potassium ferrocyanide trihydrate was obtained from Sigma Aldrich and Alfa Aesar at different levels of purity ($>99.95\%$, $>99.5\%$, and $>98.5\%$). Sodium perfluorooctanoate (NaPF_8A , 95%) was obtained from Strem. Sodium perfluoroheptanoate (NaPF_7A , 95%) and sodium perfluoropentanoate (NaPF_5A , 95%) were obtained from Matrix Scientific. Sodium perfluorohexanoate (NaPF_6A , 95%) and sodium perfluorobutanoate (NaPF_4A , 95%) were obtained from Combi-Blocks, Inc.

Sample preparation

Stock solutions of Na_2SO_3 and $\text{K}_4\text{Fe}(\text{CN})_6$ were freshly prepared before each analysis. Samples used for transient absorption experiments were prepared in a standard 1 cm pathlength quartz cuvette and consisted of either 10 mM Na_2SO_3 or 40 μM $\text{K}_4\text{Fe}(\text{CN})_6$ in 40 mM borate buffer¹⁵ ($\text{pH} \sim 10$) by the appropriate dilution of the stock solutions. Dissolved O_2 was removed by sparging with N_2 and samples were kept de-aerated by maintaining a dynamic headspace of N_2 . Stock solutions of NaPF_xA , where x is the number of carbons comprising the perfluoroalkyl backbone, were prepared at a concentration of either 80 mM or 500 mM.

UV-visible absorption

Steady-state UV-visible absorption spectra were obtained using a PerkinElmer Lambda 1050 UV-vis spectrometer. Spectra were collected between 190 nm and 400 nm either in neat water or aqueous 1 M NaCl solutions. No $\text{K}_4\text{Fe}(\text{CN})_6$ or Na_2SO_3 were added in these experiments. Aliquots of the NaPF_xA stock solutions were added to 2 mL neat water or 1 M NaCl and the absorbance at 230 nm was monitored.

Nanosecond transient absorption

Nanosecond transient absorption (nsTA) measurements were performed on an Edinburgh LP980K where the pump and probe were quasi-collinear. Samples were irradiated with a 254 nm laser pulse (pulse width $< 7 \text{ ns}$) using an Ekspla NT340B optical parametric oscillator. Each transient shown represents the average of at least 5 shots. Sample solutions were continuously stirred at 800 rpm using a Quantum Northwest Luma 40 thermostatted cuvette sample holder while maintaining a dynamic headspace of N_2 during the acquisition of the e_{aq}^- lifetimes. All transient data obtained were analyzed using OriginPro software.

Ionic strength correction

The observed decay rates obtained from monoexponential fits of the kinetic traces were corrected for the ionic strength (μ) of the solution,

$$\mu = \frac{1}{2} \sum c_i Z_i^2$$



where c_i is the concentration of ion i and Z_i is the charge of the ion, with each addition of PFAS according to^{35–38}

$$\log k_{\text{obs}} = \log k_{\mu=0} + 1.02 Z_{\text{PFAS}} Z_e \frac{\mu^{1/2}}{1 + a\mu^{1/2}}$$

where k_{obs} is the observed e_{aq}^- decay rate, $k_{\mu=0}$ is the e_{aq}^- decay rate at zero μ , Z_{PFAS} and Z_e are the charge on the PFAS and e_{aq}^- , respectively, and a is a parameter related to the contact distance (r) between the interacting species; if $r < 3 \text{ \AA}$ $a = 1$, whereas $a = 3$ for $r > 3 \text{ \AA}$.^{38,39} The contact distance r between PFAS and e_{aq}^- is estimated as the sum of their hydrodynamic radii. The radii have been estimated from their diffusion coefficients, D_i , according to

$$r_i = \frac{kT}{6\pi\eta D_i}$$

The diffusion coefficient for e_{aq}^- is reported to be $2.5 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ ($r = 1.5 \text{ \AA}$),⁴⁰ and determined by us for PF8A to be $4 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$ ($r = 4 \text{ \AA}$) using ¹⁹F DOSY NMR (see ESI, Fig. S1 and

S2†). We, therefore, chose $a = 3$ to correct the observed decay rates (Fig. S3†).

Results and discussion

Irradiation of aqueous solutions containing either Na_2SO_3 or $\text{K}_4\text{Fe}(\text{CN})_6$ with a 254 nm, <7 ns pulse produces a positive absorption on the microsecond timescale at 720 nm, which corresponds to the formation and decay of the e_{aq}^- .^{41–43} In 40 mM borax (pH ~ 10), the lifetimes obtained for the e_{aq}^- produced from a 10 mM solution of Na_2SO_3 or a 40 μM solution of $\text{K}_4\text{Fe}(\text{CN})_6$ in the absence of NaPFxA (τ_0) are $28 \pm 6 \mu\text{s}$ and $11 \pm 3 \mu\text{s}$ (Fig. 1), respectively.

e_{aq}^- quenching by NaPF8A in $\text{Fe}(\text{CN})_6^{4-}$ and SO_3^{2-}

NaPF8A was added stepwise to aqueous borate solutions containing either 40 μM $\text{Fe}(\text{CN})_6^{4-}$ or 10 mM SO_3^{2-} and the transient decays at 720 nm were recorded upon excitation at 254 nm to probe the interaction of e_{aq}^- with NaPF8A. The transients

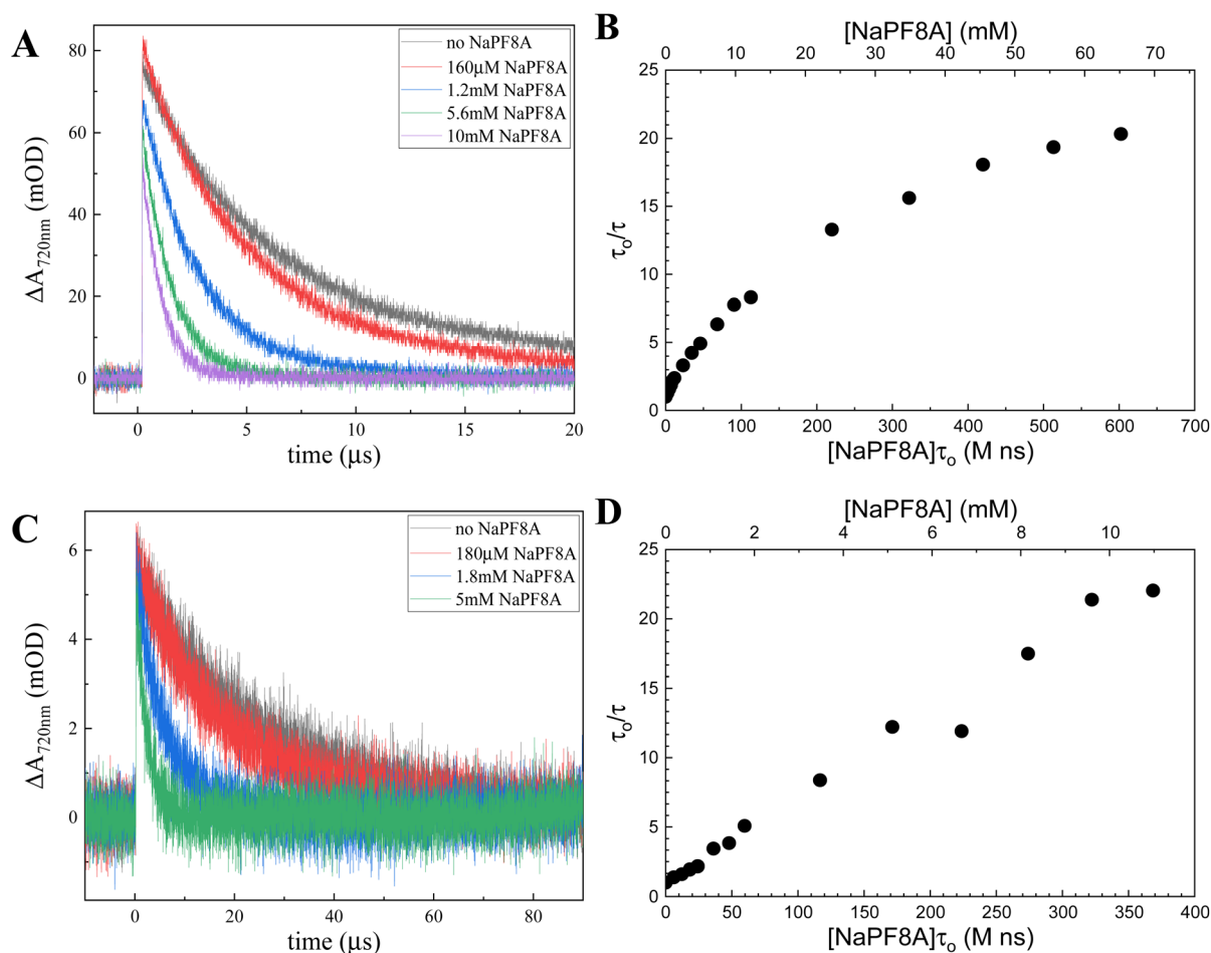


Fig. 1 (A) Transient lifetime decay of the e_{aq}^- in 40 mM borate solutions of 40 μM $\text{K}_4\text{Fe}(\text{CN})_6$ in the absence (black) and presence of 160 μM (red), 1.2 mM (blue), 5.6 mM (green), and 10 mM (purple) NaPF8A. (B) Modified Stern–Volmer plot of the e_{aq}^- lifetime rate as a function of the [NaPF8A]. (C) Transient lifetime decay of the e_{aq}^- in a 40 mM borate solution of 10 mM Na_2SO_3 in the absence (black) and presence of 180 μM (red), 1.8 mM (blue), and 5 mM (green) NaPF8A. (D) Modified Stern–Volmer plot of the e_{aq}^- lifetime data presented in panel (C).



obtained at 720 nm with each addition of NaPF8A were fit to a monoexponential decay function. In the presence of NaPF8A, an increase in the e_{aq}^- decay rate was noted in solutions of both SO_3^{2-} and $\text{Fe}(\text{CN})_6^{4-}$ (Fig. 1). The apparent bimolecular rate constant, k_{NaPF8A} , for the initial interaction of NaPF8A and the e_{aq}^- was determined from the decrease in the e_{aq}^- lifetime with increased [NaPF8A]. The rate expression corresponding to the diffusional quenching of e_{aq}^- by NaPF8A monomers resulting in the reduction of NaPF8A according to reaction 1,



is:

$$-\frac{d[e_{\text{aq}}^-]}{dt} = k_0 + k_{\text{NaPF8A}}[\text{NaPF8A}] \quad (2)$$

where k_0 is the e_{aq}^- decay rate in the absence of NaPF8A, and corresponds to the reciprocal of the lifetime of the e_{aq}^- ($k_0 = \tau_0^{-1}$) in the absence of a quencher, and k_{NaPF8A} is the apparent rate constant for the reduction of NaPF8A according to reaction 1. The data are presented according to the Stern–Volmer formalism

$$\frac{\tau_0}{\tau_{\text{NaPF8A}}} = 1 + k_{\text{NaPF8A}}[\text{NaPF8A}]\tau_0 \quad (3)$$

where τ_0 has the same meaning given above and τ_{NaPF8A} is the lifetime of the e_{aq}^- in the presence of NaPF8A and assumes diffusional oxidative quenching of e_{aq}^- by NaPF8A according to eqn (1).

In $\text{Fe}(\text{CN})_6^{4-}$, the e_{aq}^- decay rate depends linearly on [NaPF8A] for [NaPF8A] < 1 mM with a k_{NaPF8A} of $\sim 3.3 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$. However, the e_{aq}^- decay rate becomes nonlinear for [NaPF8A] > 1 mM. These results are consistent with our previous results,¹⁸ which we attributed to the submicellar aggregation of NaPF8A. We will note that others have reported on the propensity of PF8A to form pre-micellar aggregates. For example, Sarmiento, *et al.*^{44,45} invoked this process to explain changes in the partial molar volume in aqueous solutions of LiPF8A at submicellar concentrations.

Szajdzinska-Pietek and Gebicki⁴⁶ reported radiolysis data that also exhibited two distinct linear regimes in the dependence of the e_{aq}^- decay rate as a function of the concentration of ammonium perfluorooctanoate, $\text{NH}_4\text{PF8A}$, in water at [NH₄PF8A] between 5 mM and 40 mM. In their work, they reported a discontinuity that occurs near the critical micelle concentration (~ 30 mM).

We previously developed a kinetic model for the e_{aq}^- decay rate dependence on [NaPF8A]¹⁸ that included e_{aq}^- quenching terms by both NaPF8A monomers according to reaction 4 and aggregates assuming that the only configuration of the aggregates are dimers according to reaction 5.



The fits reported were restricted to [NaPF8A] < 2 mM to isolate the dynamic concentration range in which monomers

and dimers are presumed to dominate the population of NaPF8A. Although adequate fits of the data < 1 mM NaPF8A were obtained using the model, it was stressed that the PF8A aggregation is likely complex and involves a wide range of aggregation states. We, therefore, will refrain from invoking the dimerization model here and, instead, report k_{NaPF8A} based on linear fits over regions in which the concentration dependence is nearly linear (*e.g.* [NaPF8A] < 1 mM, 1 mM < [NaPF8A] < 10 mM, and [NaPF8A] > 10 mM). From this point on, [NaPF8A] refers to the total concentration of NaPF8A rather than distinguishing between monomers and aggregates; as a result, the reported k_{NaPF8A} are likely lower than the actual values for the bimolecular rate constants.

A linear fit of the data in 40 mM borate at [NaPF8A] < 1 mM yields a $k_{\text{NaPF8A}} = (2.0 \pm 0.9) \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$, which is approximately a factor of 3 smaller, but on the same order of magnitude as the result previously reported at concentrations below 200 μM in neat water, $(7.1 \pm 0.6) \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$.¹⁷ The reaction conditions here differ from those of our previous reports due to the presence of borate, which buffered the sample solutions at pH 10. Repeating the experiment where aliquots of NaPF8A were added to a solution of $\text{Fe}(\text{CN})_6^{4-}$ in neat water (Fig. S3B†) yields a $k_{\text{NaPF8A}} = (5.5 \pm 1.6) \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ at [NaPF8A] < 1 mM, which is in excellent agreement with our previous results.^{17,18} The reason for the near 3-fold decrease in k_{NaPF8A} in borate is not readily apparent and outside the scope of this study.

Linear fits at 1 mM < [NaPF8A] < 10 mM yield a k_{NaPF8A} of $(5.9 \pm 2.0) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$, which is in agreement with previous results reported by us¹⁸ and by Szajdzinska-Pietek and Gebicki⁴⁶ (*e.g.*, $5.1 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$) for [NH₄PF8A] < 30 mM, but still about five times greater than the $(1.7 \pm 0.5) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ value reported by Huang, *et al.*¹⁶ Linear fits of the data at [NaPF8A] > 10 mM yield a k_{NaPF8A} of $(2.4 \pm 0.7) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$, which is within experimental error of the aforementioned results reported by Huang *et al.*¹⁶ and in good agreement with the $1.3 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ value reported by Szajdzinska-Pietek and Gebicki⁴⁶ for [NH₄PF8A] > 30 mM, above its critical micelle concentration, CMC.

The reduction of PFxA by e_{aq}^- is believed to proceed by one of three mechanisms: the associative (reaction 6), concerted dissociative (reaction 7), or stepwise dissociative mechanism (reaction 8). Due to experimental limitations, the dominant mechanism driving PFAS reduction and defluorination has yet to be positively identified. As such, a good deal of theoretical work has been dedicated to disentangling these fast initial reactions.^{12,47–51}



Daily and Minakata⁴⁸ used density functional theory to calculate the one electron reduction potentials corresponding



Table 1 Summary of k_{NaPFxA} obtained in aqueous solutions of ferrocyanide and sulfite

	$k_{\text{NaPFxA}} \text{ (M}^{-1} \text{ s}^{-1}\text{)}$			Literature	
	$[\text{NaPFxA}] < 1 \text{ mM}$	$1 \text{ mM} < [\text{NaPFxA}] < 10 \text{ mM}$	$[\text{NaPFxA}] > 10 \text{ mM}$	$[\text{NaPFxA}] > 5 \text{ mM}$	Ref.
40 $\mu\text{M Fe(CN)}_6^{4-}$					
NaPF8A	$(2.0 \pm 0.9) \times 10^8$	$(5.9 \pm 2.0) \times 10^7$	$(2.4 \pm 0.7) \times 10^7$	$(1.7 \pm 0.5) \times 10^7$	16
NaPF7A	$(7.7 \pm 2.0) \times 10^7$	$(2.5 \pm 0.4) \times 10^7$	$(1.9 \pm 0.4) \times 10^7$		
NaPF6A	$(9.8 \pm 2.0) \times 10^7$	$(3.3 \pm 0.3) \times 10^7$	$(2.4 \pm 0.3) \times 10^7$		
NaPF5A	$(6.7 \pm 1.6) \times 10^7$	$(1.5 \pm 0.3) \times 10^7$	$(1.3 \pm 0.1) \times 10^7$		
NaPF4A	$(1.0 \pm 0.2) \times 10^7$	$(3.7 \pm 0.2) \times 10^7$	$(2.5 \pm 0.3) \times 10^7$	$(7.1 \pm 0.3) \times 10^6, (1.3 \pm 0.1) \times 10^7$	15 and 16
10 mM SO_3^{2-}		$k_{\text{NaPFxA}} \text{ (M}^{-1} \text{ s}^{-1}\text{)}$			
NaPF8A		$(5.6 \pm 0.2) \times 10^7$			
NaPF7A		$(2.0 \pm 0.1) \times 10^7$			
NaPF6A		$(2.9 \pm 0.2) \times 10^7$			
NaPF5A		$(1.0 \pm 0.1) \times 10^7$			
NaPF4A		$(2.0 \pm 0.2) \times 10^7$			

to reactions 6–8 for PF4A, PF6A, and PF8A among other organic compounds. From the theoretical reduction potentials the authors then calculated the predicted rate constants for reduction and found that for PF8A the associative and concerted dissociative mechanisms are expected to proceed with a rate constant of $5.8 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ and $1.7 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$, respectively. These predicted values are in excellent agreement with our results in neat aqueous solutions of Fe(CN)_6^{4-} at $[\text{NaPF8A}] < 1 \text{ mM}$ and $> 10 \text{ mM}$, respectively, and in very good agreement with solutions of Fe(CN)_6^{4-} in borax (Table 1) in the same concentration ranges.

We note that a recent study has disputed the nonlinearity observed by us for e_{aq}^- quenching in solutions of Fe(CN)_6^{4-} at PFAS concentrations $< 1 \text{ mM}$.¹⁵ In order to verify that our observed trend is not due to impurities in our commercially obtained $\text{K}_4\text{Fe(CN)}_6$, we performed the same experiment where aliquots of NaPF8A were incrementally added to aqueous solutions (in the absence of borate) of $\text{K}_4\text{Fe(CN)}_6$ obtained from two different suppliers and three different degrees of purity. We found the trend and results, particularly at $[\text{NaPF8A}] < 1 \text{ mM}$, to be virtually indistinguishable between the three data sets (Fig. S3B†).

Interestingly, the interactions between e_{aq}^- and NaPF8A are quite different in 10 mM SO_3^{2-} solutions compared to Fe(CN)_6^{4-} over similar $[\text{NaPF8A}]$. Upon addition of NaPF8A to 10 mM SO_3^{2-} , negligible e_{aq}^- lifetime quenching was observed below $200 \mu\text{M}$. At $[\text{NaPF8A}] > 200 \mu\text{M}$, however, the e_{aq}^- decay rate increases linearly with $[\text{NaPF8A}]$ up to 10 mM and has a k_{NaPF8A} of $(5.6 \pm 0.2) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$, which correlates well with the values obtained here for the k_{NaPF8A} in $1 \text{ mM} < [\text{NaPF8A}] < 10 \text{ mM}$ solutions of Fe(CN)_6^{4-} . A nonlinear dependence of the e_{aq}^- decay rate on $[\text{NaPF8A}]$ is also observed in solutions of 1 mM KI (Fig. S4†) in neat water; however, the dependence of the e_{aq}^- decay rate on $[\text{NaPF8A}]$ becomes increasing linear when the concentration of KI is increased. The k_{NaPF8A} between $1 \text{ mM} < [\text{NaPF8A}] < 10 \text{ mM}$ was also observed to decrease from $5.9 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ in 1 mM KI to $3.5 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ in 10 mM KI . Moreover, a significant difference in the k_{NaPF8A} obtained between $1 \text{ mM} <$

$[\text{NaPF8A}] < 10 \text{ mM}$ was noted when comparing results in 10 mM KI and $80 \mu\text{M indole}$ to $40 \mu\text{M Fe(CN)}_6^{4-}$ and 10 mM SO_3^{2-} .

Effect of NaPF8A on the absorption spectrum of e_{aq}^-

We recorded the e_{aq}^- spectrum in Fe(CN)_6^{4-} and SO_3^{2-} solutions both with and without 4 mM NaPF8A (Fig. 2). The e_{aq}^- spectrum is unaffected by the presence of NaPF8A suggesting that the solvent properties are unperturbed. It is known that the transition energy of e_{aq}^- depends on the solvent and is, therefore, sensitive to the differences in its local environment.^{52–54} These differences manifest in a shift of the wavelength of the e_{aq}^- absorption maximum. For example, excitation of aqueous Triton tx-165 at concentrations above its CMC results in the formation of two distinct populations of e_{aq}^- evidenced by the presence of two absorption maxima located at 720 nm and 630 nm that respectively correspond to e_{aq}^- in water and in the less polar micellar interior.⁵⁴ Here, the e_{aq}^- displays an absorption maximum at 719 nm and 717 nm in solutions of Fe(CN)_6^{4-} and SO_3^{2-} , respectively, in the absence of NaPF8A. In the presence of 4 mM NaPF8A , no shifts were observed for the e_{aq}^- absorption maxima in solutions of Fe(CN)_6^{4-} and SO_3^{2-} suggesting that perturbations to the physicochemical properties of the solvent are negligible up to 4 mM NaPF8A .

Salt effect on the NaPF8A reduction rate constant

Increased salt concentration and ionic strength are known to promote aggregation and lower CMC values.^{55,56} To determine the effect of ionic strength on the measured rates constants, we incrementally added aliquots of NaPF8A into solutions of Fe(CN)_6^{4-} and SO_3^{2-} containing 1 M NaCl (Fig. 3). In solutions of Fe(CN)_6^{4-} , the presence of 1 M NaCl results in a more pronounced deviation from a linear dependence of the e_{aq}^- decay rate with $[\text{NaPF8A}]$ and the k_{NaPF8A} values at $[\text{NaPF8A}] < 1 \text{ mM}$ and $1 \text{ mM} < [\text{NaPF8A}] < 10 \text{ mM}$ are $(1.3 \pm 0.3) \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ and $(4.1 \pm 0.5) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$, respectively. The k_{NaPF8A} obtained at $[\text{NaPF8A}] < 1 \text{ mM}$ in the absence and presence of 1 M NaCl are found to agree within experimental error. However, at $1 \text{ mM} < [\text{NaPF8A}] < 10 \text{ mM}$, addition of NaCl induces a nearly two-fold



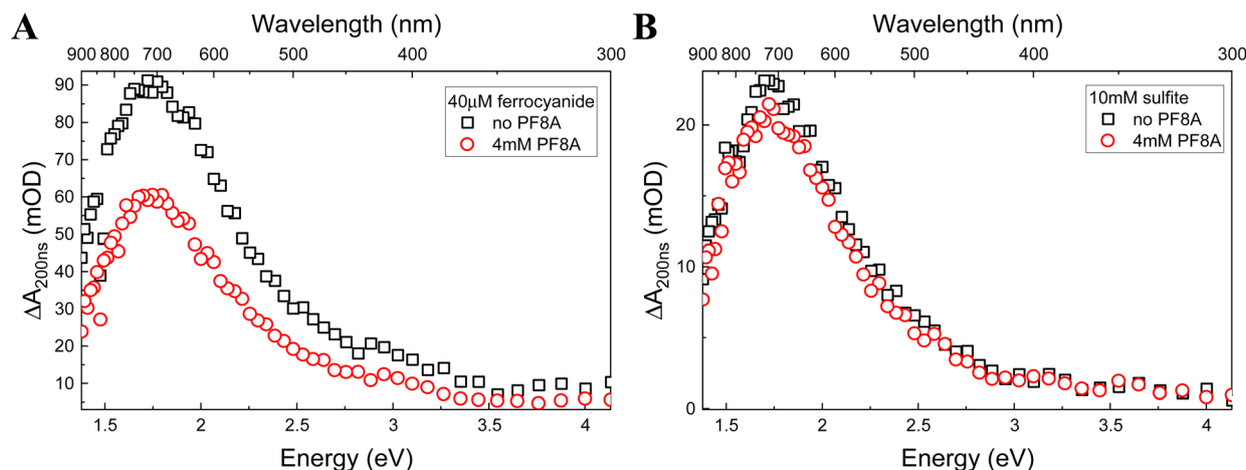


Fig. 2 Transient absorption spectra of the solvated electron at 200 ns after excitation in the absence (black) and presence (red) of 4 mM NaPF8A in 40 mM borate solutions of (A) 40 μM $\text{Fe}(\text{CN})_6^{4-}$ and (B) 10 mM SO_3^{2-} .

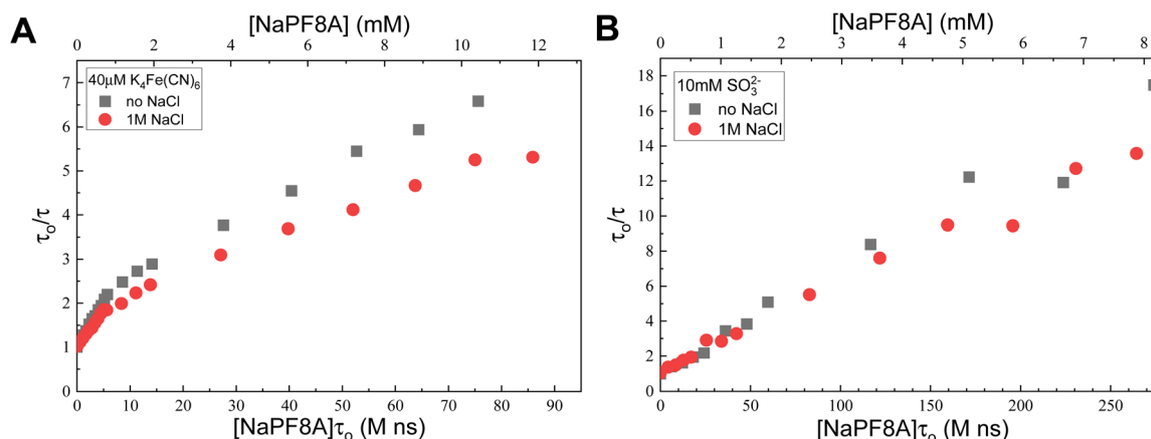


Fig. 3 Modified Stern–Volmer plots of the e_{aq}^- lifetime dependence on [NaPF8A] in the absence (black) and presence of (red) 1 M NaCl in 40 mM borate solutions of (A) 40 μM $\text{Fe}(\text{CN})_6^{4-}$ and (B) 10 mM SO_3^{2-} .

decrease in the k_{NaPF8A} . Interestingly, the addition of 1 M NaCl to solutions of SO_3^{2-} resulted in a nearly negligible decrease in k_{NaPF8A} to $(4.2 \pm 0.8) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ (Fig. 3).

There is less of an effect from adding NaCl to the 10 mM SO_3^{2-} because it has a higher initial ionic strength than that of the 40 μM $\text{Fe}(\text{CN})_6^{4-}$ so the former is less susceptible to additional increases in the ionic strength. Increasing the NaCl concentration promotes NaPF8A aggregation but the effect is reduced in sulfite solution because the ionic strength is already high. This interpretation is corroborated by the trend observed in varying concentrations of KI with NaPF8A (Fig. S4[†]) in which the non-linearity of the data decreases with increasing [KI] and, as a result, the solution ionic strength (*vide supra*). The results obtained at high [NaCl] lend further support to this proposal. As noted earlier, the critical micelle concentration is known to decrease with increasing ionic strength.^{55,56} Analogously, we believe that the interactions between PFAS monomers increase with increasing ionic strength shifting the equilibrium from monomers to aggregates.

Effect of PF8A counterion on e_{aq}^- quenching

Counterions are known to affect surfactant aggregation.^{57,58} For example, Lunkenheimer, *et al.* demonstrated a decrease in the CMC of various PF8A salts following the series $\text{Li}^+ > \text{Na}^+ > \text{K}^+ > \text{NH}_4^+ > \text{Rb}^+ > \text{Cs}^+$.^{57,58} We, therefore, investigated whether there were differences between Na^+ and NH_4^+ cations on the e_{aq}^- quenching behavior of PF8A in $\text{Fe}(\text{CN})_6^{4-}$ and SO_3^{2-} solutions. Results of $\text{NH}_4\text{PF8A}$ and NaPF8A quenching using $\text{Fe}(\text{CN})_6^{4-}$ and SO_3^{2-} as precursor are shown in Fig. 4. In $\text{Fe}(\text{CN})_6^{4-}$ solutions the dependence of the e_{aq}^- decay rate was lower for NaPF8A than for $\text{NH}_4\text{PF8A}$ at 1 mM $< [\text{XPF8A}] < 10$ mM which is consistent with more aggregation in the former. We found that the k_{XPF8A} (where X is either Na^+ or NH_4^+) was 3–4 times greater for $\text{NH}_4\text{PF8A}$, $(2.0 \pm 0.1) \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ than for NaPF8A, $(5.9 \pm 0.2) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$. We believe the difference in the k_{XPF8A} observed between Na^+ and NH_4^+ are related to differences in the interaction strength between the carboxylate head group of PF8A and the counter ion. We should note that the radius of hydration differ between Na^+



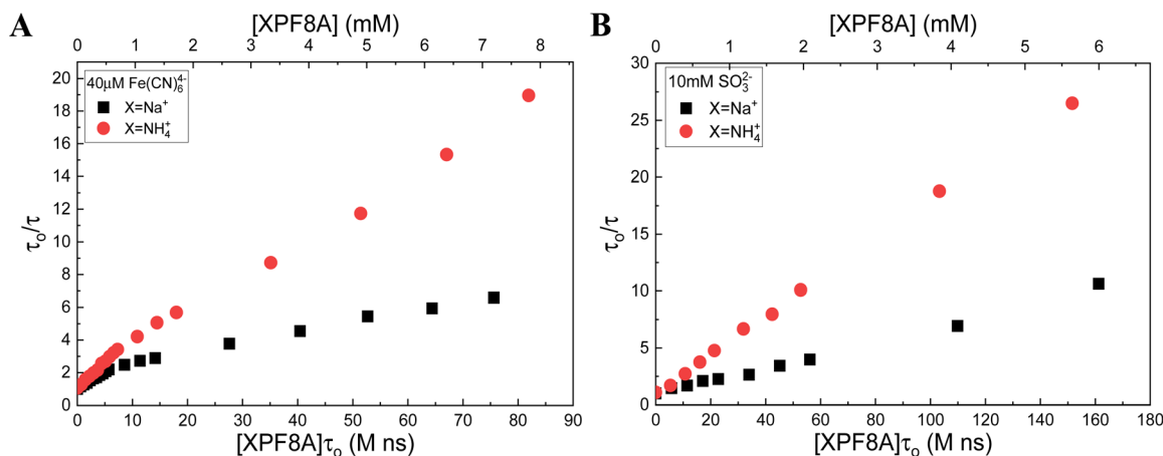


Fig. 4 Modified Stern–Volmer plots of the e_{aq}^- lifetime dependence on the [NaPF8A] (black) and [NH₄PF8A] (red) in 40 mM borate solutions of (A) 40 μM $\text{Fe}(\text{CN})_6^{4-}$ and (B) 10 mM SO_3^{2-} .

and NH_4^+ which may also contribute to the differences in the observed trends;⁵⁸ however, the nature of the contribution is uncertain at this time. Interestingly, in solutions of $\text{Fe}(\text{CN})_6^{4-}$ the dependence of the e_{aq}^- decay rate on [NH₄PF8A] displays a larger deviation from linearity in the presence of 1 M NaCl (Fig. S5†) likely due to cation exchange in excess Na⁺. In solutions of SO_3^{2-} , we found that the k_{XPF8A} was similarly larger by a factor of 3–4 for NH₄PF8A, $(1.8 \pm 0.1) \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ compared to NaPF8A, $(5.6 \pm 0.2) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$.

e_{aq}^- quenching by NaPFxA

In addition to NaPF8A, we also examined NaPFxA of other chain lengths, where x corresponds to the number of carbons comprising the PFAS backbone ($x = 8, 7, 6, 5,$ and 4). In each case we found a nonlinear dependence of the e_{aq}^- decay rate on [NaPFxA] in $\text{Fe}(\text{CN})_6^{4-}$ solutions (Fig. 5). Although the data are nonlinear in $\text{Fe}(\text{CN})_6^{4-}$, suggesting aggregation with increasing [NaPFxA] in all cases, we will direct our attention to the k_{NaPFxA} at $1 \text{ mM} < [\text{NaPFxA}] < 10 \text{ mM}$ for comparison to the data obtained in SO_3^{2-} . In both $\text{Fe}(\text{CN})_6^{4-}$ and SO_3^{2-} a general

decreasing trend is observed for the k_{NaPFxA} in the range $1 \text{ mM} < [\text{NaPFxA}] < 10 \text{ mM}$ with decreasing chain length, which is in good agreement with the trend that has been reported in the literature for the initial reduction of PFxA.¹⁶ We will note that although the k_{NaPF8A} obtained by us in $\text{Fe}(\text{CN})_6^{4-}$ ($6.6 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$) and SO_3^{2-} ($5.6 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$) at $[\text{NaPF8A}] > 1 \text{ mM}$ are 3–4 times greater than the $1.7 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ value reported by Huang, *et al.*¹⁶ they are in very good agreement with the $5.1 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ value reported by Szajdzinska-Pietek and Gebicki⁴⁶ in the concentration range of $5 \text{ mM} < [\text{NH}_4\text{PF8A}] < 30 \text{ mM}$. We have previously¹⁸ pointed out that the rate constant reported by Huang, *et al.*¹⁶ is in very good agreement with the $1.3 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ rate constant reported by Szajdzinska-Pietek and Gebicki⁴⁶ at $[\text{NH}_4\text{PF8A}]$ above the CMC $\sim 30 \text{ mM}$.

Steady-state absorption spectrum of NaPFxA

A number of experimental techniques have traditionally been used to monitor aggregation in solution, particularly *vis-à-vis* surfactant aggregation and micelle formation, including surface tensiometry, conductivity, steady-state absorbance and

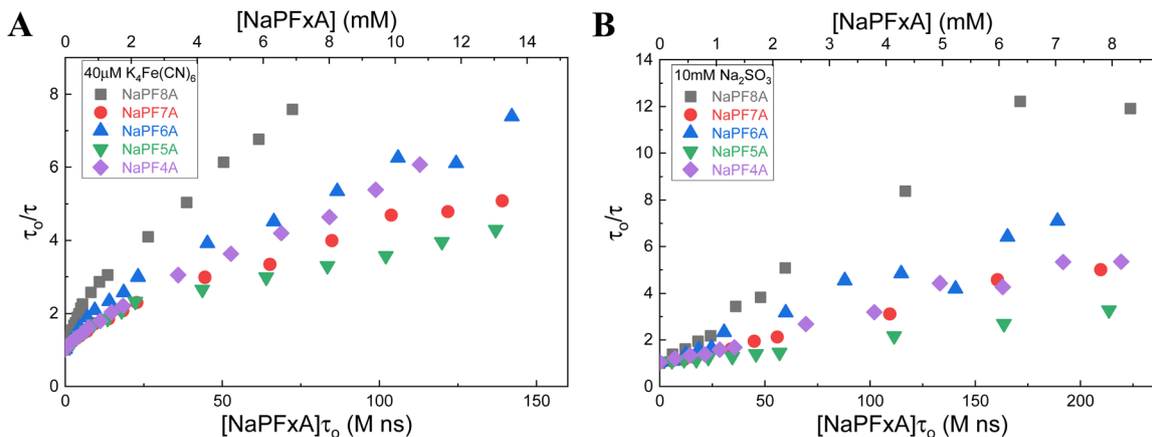


Fig. 5 Modified Stern–Volmer plots of the e_{aq}^- lifetime dependence on the [NaPFxA] in 40 mM borate solutions of (A) 40 μM $\text{Fe}(\text{CN})_6^{4-}$ and (B) 10 mM SO_3^{2-} .



fluorescence spectroscopy. In the case of absorbance and fluorescence spectroscopy, due to the low molar extinction and fluorescence quantum yields of many surfactant molecules, surfactant aggregation is typically estimated with the help of solvatochromic reporters or hydrophobic chromophores that preferentially partition within the hydrophobic core of micelles. However, despite the low molar extinction of many surfactants, we believe that deviations from linearity of the absorbance of the surfactants in water may be observed and would be indicative of aggregation (Fig. S6†).⁵⁹

As a demonstration that surfactant absorption can be used to identify aggregation, steady-state spectra were measured for aqueous solutions with increasing amounts of NaPFxA up to 40 mM. At low concentrations of NaPF8A (<100 μ M), the spectrum obtained is relatively featureless at wavelengths > 200 nm (Fig. 6). A shoulder centered at \sim 210 nm emerges at \sim 100 μ M NaPF8A and increases in prominence with increasing [NaPF8A]. At [NaPF8A] > 2 mM, another shoulder is observed centered at \sim 255 nm; at much higher concentrations another feature is observed at \sim 288 nm. A small deviation from linearity in the

steady-state absorption is observed between 5 mM and 40 mM (as shown in the inset of Fig. 6). The first derivative plot of the absorbance at 230 nm with respect to the [NaPF8A] displays an inflection point between 30 mM and 38 mM; this is consistent with the reported CMC values between 33 mM and 36 mM for NaPF8A.^{60–66} When adding $\text{NH}_4\text{PF8A}$ to neat water up to 10 mM, we observed only a feature at 210 nm up to 10 mM.

Results similar to those for NaPF8A were observed for other NaPFxAs up to 10 mM. In all cases, a prominent inflection point in the steady-state absorbance data of aqueous NaPFxA solutions was observed at [NaPFxA] < 0.5 mM, well below the respective CMCs, which we attribute to the formation of pre-micellar aggregates (*vide infra*).^{18,44,45} It is interesting that this inflection feature occurs at nearly the same concentration for each NaPFxA when the CMCs are very much chain length dependent and vary in magnitude by a considerable amount suggesting that pre-micellar aggregation is less influenced by the chain length and dominated by some other property.^{60,66} It is also noteworthy that, although absent in the transient absorption data in solutions of SO_3^{2-} , deviations from linearity in the

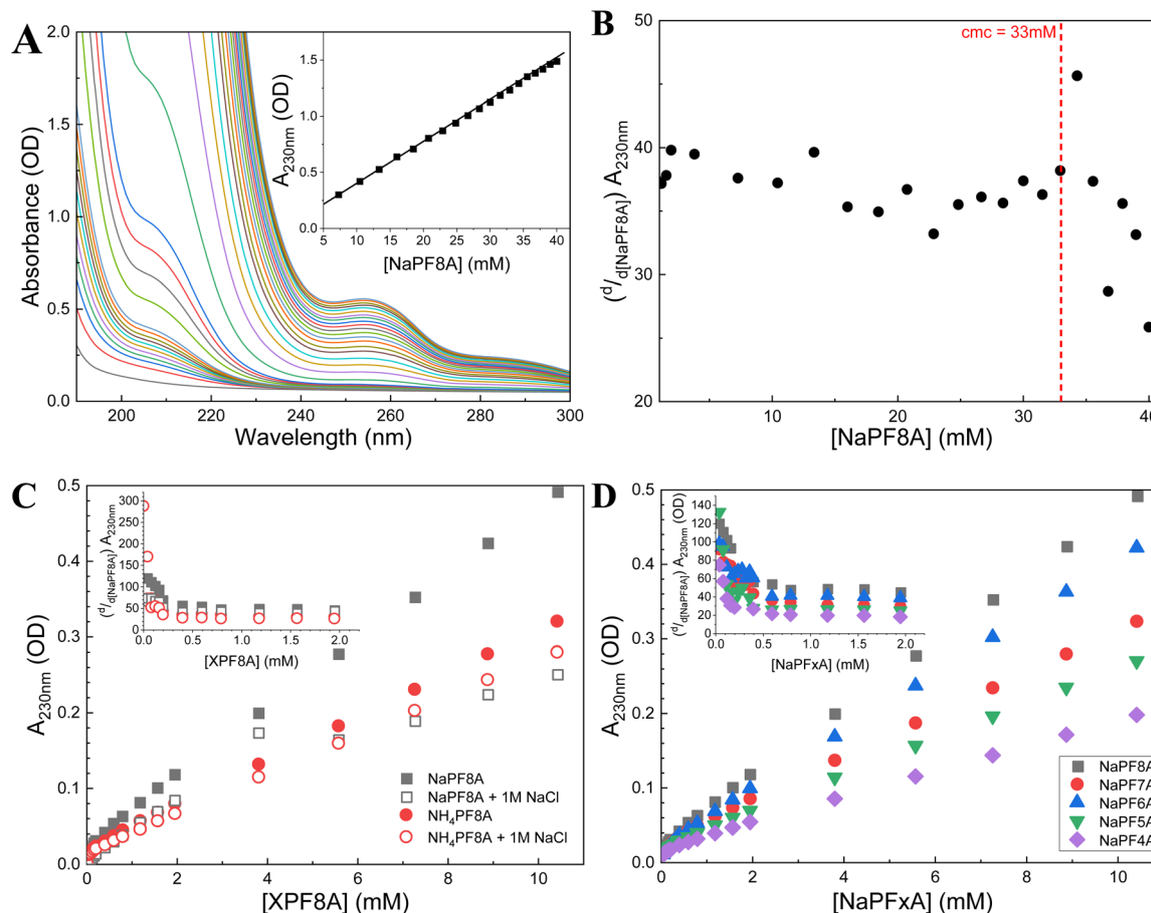


Fig. 6 (A) Steady-state absorbance of aqueous solutions (in the absence of borate, $\text{Fe}(\text{CN})_6^{4-}$, and SO_3^{2-}) at concentrations of NaPF8A increasing from 40 μ M to 40 mM. (B) First derivative of the steady-state absorbance at 230 nm as a function of the concentration of NaPF8A between 5 mM and 40 mM. The dotted red line is the CMC value reported for NaPF8A in ref. 60. (C) Steady-state absorbance at 230 nm as a function of the concentration of NaPF8A (black) and $\text{NH}_4\text{PF8A}$ (red) up to 10 mM; the inset shows the first-derivative of the same data between 0 mM and 2 mM. (D) Steady-state absorbance at 230 nm as a function of the concentration of NaPF8A (black), NaPF7A (red), NaPF6A (blue), NaPF5A (green), and NaPF4A (purple) up to 10 mM; the inset shows the first-derivative of the same data between 0 mM and 2 mM.



observed e_{aq}^- decay rates as a function of $[NaPFxA]$ in solutions of $Fe(CN)_6^{4-}$ occur at nearly the same concentrations lending more weight to the argument that the trends observed in the transient absorption data in solutions of $Fe(CN)_6^{4-}$ are due to aggregation effects. We believe that the latter is not coincidental and that the physical process inducing the nonlinearity in the steady-state absorbance is the same process that underly the trend in the transient absorption data in solutions of $Fe(CN)_6^{4-}$; namely, submicellar aggregation of $NaPFxA$.

Conclusions

Although the biomolecular rate constants for the quenching of the e_{aq}^- by PFAS were previously assumed to not depend on the identity of the e_{aq}^- photodetachment precursor, in actuality the measurement of the reaction kinetics are complicated by several factors. Here, nanosecond transient absorption spectroscopy was used to measure apparent rate constants, k_{NaPFxA} , for $NaPFxA$ quenching of the e_{aq}^- with ferrocyanide and sulfite precursors in water. The results herein demonstrated that ionic strength and cation identity effects carry significant ramifications for PFAS reactivity with the e_{aq}^- due to their influence on submicellar surfactant aggregation.

We found that the k_{NaPFxA} for e_{aq}^- quenching by $NaPFxA$ is not constant over the quencher concentration range for ferrocyanide. Additional studies were carried out to demonstrate the aggregation of $NaPFxA$, even below the CMC. Ionic strength is known to not only influence the measured rate constants for e_{aq}^- due to electrostatic screening, which we accounted for, but it can also influence aggregation, as shown here. The results of other measurements further corroborate our interpretation that aggregation explains the anomalous, concentration-dependent rate constant values. The e_{aq}^- decay rate was also found to vary when the $PFxA$ counterion was changed in a manner consistent with aggregation. Steady-state UV-visible spectra of $NaPFxA$ without ferrocyanide or sulfite (or borate) deviate significantly from the Beer-Lambert law well below their CMC, which is yet another indication of submicellar aggregation of $PFxA$. The results here reinforce our previous interpretation that $PFxA$ exhibit strong aggregation tendencies, which should be taken into account to get accurate rate constants when measuring solvated electron quenching kinetics.

Author contributions

WAM: conceptualization, methodology, investigation, formal analysis, writing – original draft. JAR: investigation, formal analysis, writing – review & editing. BLC: investigation, formal analysis, writing – review & editing. AE: supervision, writing – review & editing. JCO: supervision, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 Y. Li, T. Fletcher, D. Mucs, K. Scott, C. H. Lindh, P. Tallving and K. Jakobsson, *Occup. Environ. Med.*, 2018, **75**, 46–51.
- 2 X. C. Hu, D. Q. Andrews, A. B. Lindstrom, T. A. Bruton, L. A. Schaidler, P. Grandjean, R. Lohmann, C. C. Carignan, A. Blum, S. A. Balan, C. P. Higgins and E. M. Sunderland, *Environ. Sci. Technol. Lett.*, 2016, **3**, 344–350.
- 3 G. F. Peaslee, J. T. Wilkinson, S. R. McGuinness, M. Tighe, N. Caterisano, S. Lee, A. Gonzales, M. Roddy, S. Mills and K. Mitchell, *Environ. Sci. Technol. Lett.*, 2020, **7**, 594–599.
- 4 C. D. Vecitis, H. Park, J. Cheng, B. T. Mader and M. R. Hoffmann, *Front. Environ. Sci. Eng.*, 2009, **3**, 129–151.
- 5 B. Ji, P. Y. Kang, T. Wei and Y. Q. Zhao, *Chemosphere*, 2020, **250**, 126316.
- 6 Y. R. Gu, T. Z. Liu, H. J. Wang, H. L. Han and W. Y. Dong, *Sci. Total Environ.*, 2017, **607**, 541–548.
- 7 Y. R. Gu, W. Y. Dong, C. Luo and T. Z. Liu, *Environ. Sci. Technol.*, 2016, **50**, 10554–10561.
- 8 Y. R. Gu, T. Z. Liu, Q. Zhang and W. Y. Dong, *Chem. Eng. J.*, 2017, **326**, 1125–1133.
- 9 R. A. Marcus, *J. Chem. Phys.*, 1965, **43**, 3477–3489.
- 10 J. A. Marsella, A. G. Gilcinski, A. M. Coughlin and G. P. Pez, *J. Org. Chem.*, 1992, **57**, 2856–2860.
- 11 A. A. Pud, G. S. Shapoval, V. P. Kukhar, O. E. Mikulina and L. L. Gervits, *Electrochim. Acta*, 1995, **40**, 1157–1164.
- 12 D. J. Van Hoomissen and S. Vyas, *Environ. Sci. Technol. Lett.*, 2019, **6**, 365–371.
- 13 B. Guan, J. Zhi, X. Zhang, T. Murakami and A. Fujishima, *Electrochem. Commun.*, 2007, **9**, 2817–2821.
- 14 V. Ochoa-Herrera, R. Sierra-Alvarez, A. Somogyi, N. E. Jacobsen, V. H. Wysocki and J. A. Field, *Environ. Sci. Technol.*, 2008, **42**, 3260–3264.
- 15 C. K. Amador, D. J. Van Hoomissen, J. Q. Liu, T. J. Strathmann and S. Vyas, *Chemosphere*, 2023, **311**, 136918–136926.
- 16 L. Huang, W. B. Dong and H. Q. Hou, *Chem. Phys. Lett.*, 2007, **436**, 124–128.
- 17 W. A. Maza, V. M. Breslin, J. C. Owrutsky, B. B. Pate and A. Epshteyn, *Environ. Sci. Technol. Lett.*, 2021, **8**, 525–530.
- 18 W. A. Maza, B. D. Etz, T. C. Schutt, B. L. Chaloux, V. M. Breslin, B. B. Pate, M. K. Shukla, J. C. Owrutsky and A. Epshteyn, *Environ. Sci. Technol. Lett.*, 2022, **9**, 226–232.
- 19 P. Airey and F. S. Dainton, *Proc. R. Soc. London, Ser. A*, 1966, **291**, 340–352.
- 20 M. Shirom and G. Stein, *J. Chem. Phys.*, 1971, **55**, 3372–3378.
- 21 M. Shirom and M. Tomkiewi, *J. Chem. Phys.*, 1972, **56**, 2731–2736.
- 22 G. Stein, *Isr. J. Chem.*, 1970, **8**, 691–697.
- 23 G. M. Sando and J. C. Owrutsky, *J. Phys. Chem. B*, 2006, **110**, 9586–9592.



- 24 W. A. Maza, V. M. Breslin, N. T. Plymale, P. A. DeSario, A. Epshteyn, J. C. Owrutsky and B. B. Pate, *Photochem. Photobiol. Sci.*, 2019, **18**, 1526–1532.
- 25 A. E. Martell and R. M. Smith, *Critical Stability Constants*, Plenum Press, New York, N.Y., 1974.
- 26 E. Hayon, A. Treinin and J. Wilf, *J. Am. Chem. Soc.*, 1972, **94**, 47–57.
- 27 L. Dogliotti and E. Hayon, *J. Phys. Chem.*, 1968, **72**, 1800–1807.
- 28 M. C. Sauer, R. A. Crowell and I. A. Shkrob, *J. Phys. Chem. A*, 2004, **108**, 5490–5502.
- 29 P. F. Gu, C. J. Zhang, Z. Y. Sun, H. Z. Zhang, Q. Zhou, S. J. Lin, J. Y. Rong and M. R. Hoffmann, *Chem. Eng. J.*, 2020, **379**, 122338.
- 30 Z. H. Chen, Y. Teng, N. Mi, X. Jin, D. S. Yang, C. Wang, B. Wu, H. Q. Ren, G. X. Zeng and C. Gu, *Environ. Sci. Technol.*, 2021, **55**, 3996–4006.
- 31 H. Park, C. D. Vecitis, J. Cheng, N. F. Dalleska, B. T. Mader and M. R. Hoffmann, *Photochem. Photobiol. Sci.*, 2011, **10**, 1945–1953.
- 32 H. Park, C. D. Vecitis, J. Cheng, W. Choi, B. T. Mader and M. R. Hoffmann, *J. Phys. Chem. A*, 2009, **113**, 690–696.
- 33 Y. Qu, C. J. Zhang, F. Li, J. Chen and Q. Zhou, *Water Res.*, 2010, **44**, 2939–2947.
- 34 K. E. Yu, X. C. Li, L. W. Chen, J. Y. Fang, H. L. Chen, Q. B. Li, N. P. Chi and J. Ma, *Water Res.*, 2018, **129**, 357–364.
- 35 J. Rabani, in *Solvated Electron*, ed. R. F. Gould, American Chemical Society, Washington, D.C., 1965, vol. 50, ch. 17, p. 242.
- 36 M. Matheson, in *Solvated Electron*, ed. R. F. Gould, American Chemical Society, Washington, D.C., 1965, vol. 50, ch. 5, p. 45.
- 37 S. R. Logan, *Trans. Faraday Soc.*, 1967, **63**, 3004–3008.
- 38 G. Czapski and H. A. Schwarz, *J. Phys. Chem.*, 1962, **66**, 471–474.
- 39 S. Gordon, E. J. Hart, M. S. Matheson, J. Rabani and J. K. Thomas, *J. Am. Chem. Soc.*, 1963, **85**, 1375–1377.
- 40 M. Anbar, in *Solvated Electron*, ed. R. F. Gould, American Chemical Society, Washington, D.C., 1965, vol. 50, ch. 6, p. 55.
- 41 G. V. Buxton, C. L. Greenstock, W. P. Helman and A. B. Ross, *J. Phys. Chem. Ref. Data*, 1988, **17**, 513–886.
- 42 L. M. Dorfman, F. Jou and R. Wageman, *Ber. Bunsen-Ges.*, 1971, **75**, 681–685.
- 43 E. J. Hart, B. Michael and K. H. Schmidt, *J. Phys. Chem.*, 1971, **75**, 2798–2805.
- 44 E. Blanco, P. Messina, J. M. Ruso, G. Prieto and F. Sarmiento, *Mol. Phys.*, 2005, **103**, 3271–3281.
- 45 A. Gonzalez-Perez, J. M. Ruso, G. Prieto and F. Sarmiento, *J. Surfactants Deterg.*, 2004, **7**, 387–395.
- 46 E. Szajdzinska-Pietek and J. L. Gebicki, *Res. Chem. Intermed.*, 2000, **26**, 897–912.
- 47 M. J. Bentel, Y. Yu, L. Xu, Z. Li, B. M. Wong, Y. Men and J. Liu, *Environ. Sci. Technol.*, 2019, **53**, 3718–3728.
- 48 R. Daily and D. Minakata, *Environ. Sci.: Water Res. Technol.*, 2022, **8**, 543–574.
- 49 S. Biswas, S. S. R. K. C. Yamijala and B. M. Wong, *Environ. Sci. Technol.*, 2022, **56**, 8167–8175.
- 50 S. Biswas and B. M. Wong, *ACS EST Engg.*, 2023, DOI: [10.1021/acsestengg.3c00216](https://doi.org/10.1021/acsestengg.3c00216).
- 51 S. S. R. K. C. Yamijala, R. Shinde and B. M. Wong, *Phys. Chem. Chem. Phys.*, 2020, **22**, 6804–6808.
- 52 L. M. Dorfman, F. Jou and R. Wageman, *Ber. Bunsen-Ges.*, 1971, **75**, 681–685.
- 53 E. J. Hart and J. W. Boag, *J. Am. Chem. Soc.*, 1962, **84**, 4090–4095.
- 54 H. Ghosh, D. Palit, A. Sapre, K. RamaRao and J. Mittal, *Chem. Phys. Lett.*, 1993, **203**, 5–11.
- 55 L. Ambrosone and R. Ragone, *J. Colloid Interface Sci.*, 1998, **205**, 454–458.
- 56 P. Palladino and R. Ragone, *Langmuir*, 2011, **27**, 14065–14070.
- 57 K. Lunkenheimer, K. Geggel and D. Prescher, *Langmuir*, 2017, **33**, 10216–10224.
- 58 K. Lunkenheimer, D. Prescher and K. Geggel, *Langmuir*, 2022, **38**, 891–902.
- 59 X. Cui, S. Mao, M. Liu, H. Yuan and Y. Du, *Langmuir*, 2008, **24**, 10771–10775.
- 60 H. Kunieda and K. Shinoda, *J. Phys. Chem.*, 1976, **80**, 2468–2470.
- 61 N. Muller and H. Simsohn, *J. Phys. Chem.*, 1971, **75**, 942–945.
- 62 H. Nakayama, *Bull. Chem. Soc. Jpn.*, 1967, **40**, 1592–1595.
- 63 K. Shinoda, M. Hato and T. Hayashi, *J. Phys. Chem.*, 1972, **76**, 909–914.
- 64 K. Shinoda and K. Katsura, *J. Phys. Chem.*, 1964, **68**, 1568–1569.
- 65 J. L. Lopez-Fontan, F. Sarmiento and P. C. Schulz, *Colloid Polym. Sci.*, 2005, **283**, 862–871.
- 66 K. Shinoda, M. Hato and T. Hayashi, *J. Phys. Chem.*, 1972, **76**, 909–914.

