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Sea spray aerosols can be a source of PFAS pollution in coastal aquifers

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Pollution with PFASs is found in several types of environmental matrices across the globe. In groundwater, the occurrence is usually attributed to point sources like firefighting training areas, landfills or direct industrial use. During the last 15 years it has become clear that some PFASs are highly preconcentrated in sea spray aerosols and recently this was proposed to be a significant source of PFASs on land. To see if such a source is strong enough to affect groundwater, we analysed a nationwide dataset for PFASs in shallow wells. By focusing on wells located in forests or other nature areas, it became clear that groundwater within 5 km of the 400 km long Danish west facing North Sea coast is clearly affected by a diffuse PFAS source, most likely sea spray aerosols. PFOA dominated, but PFHxS was detected almost as frequently and the concentration ratio between these two PFASs was relatively constant. Four very shallow monitoring wells with 2–21-year-old groundwater were repeatedly sampled over 3–5 years, and all showed an almost constant concentration of PFOA, PFHxS, PFOS and PFBS. A screening of 60 PFASs showed that legacy PFAAs dominated wells affected by sea spray aerosols. The observed diffuse PFAS pollution in groundwater is most likely a world-wide coastal phenomenon, but additional studies are needed to achieve sufficient understanding of the drivers.

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Environmental significance

In Denmark, like in most other countries, many groundwater wells are polluted with PFASs. Often, known sources can explain this; however sometimes PFASs are detected in groundwater without any likely source. We demonstrate a very high detection frequency for certain PFASs in shallow groundwater wells located in Danish nature areas close to the North Sea coast with concentrations often exceeding local drinking water guideline values. Based on several observations, it is concluded that the only plausible explanation is transport from sea to land *via* sea spray aerosols. This process therefore needs to be considered as an important source of diffuse PFAS pollution in coastal areas.

Introduction

Per-and polyfluorinated alkyl substances (PFASs) are a diverse group of compounds, that may be divided into some major subgroups with specific chemical characteristics. The compounds most commonly detected in aqueous environmental matrices all belong to the group of perfluorinated alkyl acids (PFAAs) which can be further divided into the subgroups of perfluorinated carboxylic acids (PFCAs) and perfluorinated sulfonic acids (PFSAs), with their most "famous" members being PFOA and PFOS, respectively. Many PFCAs and PFSAs have surfactant-like molecular characteristics that make them adsorb to air/water interfaces.¹ While these molecular characteristics are well studied, their environmental implications are only beginning to be recognized.

A potentially critical aspect of the air/water adsorption of PFAA compounds is their tendency to accumulate in aerosols may travel hundreds of kilometres before the PFAAs present in the aerosols at some point will be scavenged from the atmosphere by wet and dry deposition providing a transport pathway from sea to land. In a subsequent study, it was estimated that in coastal areas more than 100 ng PFAA per m² per year may be emitted from sea water and that 15 to 30% of the PFAAs emitted from the global oceans is transported and deposited on land, with more than 80% of this deposition occurring in what is defined in the study as coastal grid cells. Theoretically, the relatively low background pollution of the oceans may hence be an important regional diffuse PFAS source on land. The actual size of such a source is largely unknown, but it is well known

derived from sea spray. In laboratory experiments, it has been

shown that compounds within the PFCA and PFSA subgroups may be enriched more than 1000 times during the aerosol

emission process.²⁻⁵ Aerosol sampling at coastal sites in Norway have provided further evidence for PFAA enrichment in sea

spray aerosols using sodium ions as a tracer for the sea-derived

aerosols.6 The same authors underpin that sea spray aerosols

that those PFAS compounds that are present in the world's

oceans and that possess the ability to adsorb to air/water

Department of Geochemistry, Geological Survey of Denmark and Greenland (GEUS), Øster Voldgade 10, DK-1350 Copenhagen, Denmark. E-mail: cal@geus.dk interfaces are found ubiquitously in rain.8-10 For example, concentrations of PFOA are typically in the range of 0.2-2 ng L^{-1} at inland locations, which theoretically would be higher in coastal areas, if sea spray aerosols are major contributors. The toxicologically derived guideline value for PFOA in drinking water is 2 ng L^{-1} in Denmark and 4 or 4.4 ng L^{-1} in several other EU countries, and recently the US-EPA has set an enforceable maximum contaminant level of 4 ng L⁻¹ for PFOA. Diffuse atmospheric PFOA pollution, which ultimately will make it to groundwater by percolation through the soil, is therefore potentially important with regards to maintaining a safe groundwater-based water supply. Recently, it was concluded that the most likely reason for a groundwater-fed waterworks close to the Danish North Sea coast having PFASs above the drinking water standard is PFASs in sea foam and sea spray aerosols.11 In order to explore the hypothesis that the transport of PFASs with sea spray aerosols is large enough to affect groundwater resources, we analysed publicly available data for PFASs in Danish groundwater. In addition, we studied four selected PFAS-contaminated shallow coastal monitoring wells with regards to time trends of PFAS concentrations, groundwater age, potential sources of PFASs and occurrence of a broad range of PFAS compounds.

Methods

Two types of groundwater wells were included in the study; wells belonging to the national groundwater monitoring program (GRUMO) and waterworks wells used for drinking water abstraction. The GRUMO wells typically have short screens (1-3 m) and are not directly affected by groundwater abstraction. The waterworks wells typically have 3-10 m long screens, but longer screens exist. To focus the data analysis on diffuse PFAS sources, we did not include groundwater wells which were sampled for the purpose of investigating PFAS point sources, but such sources may still influence some of the monitoring and waterworks wells. In Denmark, PFASs are only rarely detected in groundwater wells below 40 m depth, and we therefore exclusively focused on shallow wells with 40 m or less to top of the screen. This also secured a relatively homogeneous distribution of wells across the country. Before 2021, when new drinking water standards were introduced, reporting limits for PFASs in groundwater were often relatively high in Denmark. To minimize the risk of including non-detects solely due to high reporting limits, we therefore included only wells that were analysed between 2021 and 2024. A total of 1538 groundwater wells fulfilled these criteria, of which 1136 were drinking water abstraction wells and 402 belonged to the national monitoring program.

The groundwater wells were analysed for 22 PFAS compounds; PFCAs with 3–12 fluorinated carbons, PFSAs with 4–13 fluorinated carbons and two PFAA precursors (6:2-FTS and PFOSA). Names and typical reporting limits are shown in Table S1. The 22 PFASs were selected due to a national guideline value of 0.1 $\mu g \ L^{-1}$ for the sum of these. The analyses were executed at different commercial accredited laboratories in Denmark. We sampled four selected monitoring wells near the

Danish North Sea Coast in November 2024 and had them analysed for 60 PFASs at Eurofins Sweden AB. Compound names and details on analytical methods are in Table S2. The age of the water was also determined in these four wells using the tritiumhelium-method, with analyses performed at Bremen University as previously described. 12,13 The concentration of the ultrashortchain PFAS trifluoroacetate (TFA), which also provides information on water age, was analysed for four monitoring wells using an in-house LC-MS/MS method with a reporting limit of 0.03 $\mu g \ L^{-1}$, as described previously. 13

The data on 22 PFASs in 1538 groundwater wells were extracted from the open database Jupiter14 (https:// www.geus.dk/produkter-ydelser-og-faciliteter/data-og-kort/ national-boringsdatabase-jupiter) on 30th August 2024 and quality assured. Some data below Reporting Limits (<RL) had relatively high RL and were therefore excluded from the analysis: for the four PFASs with a guideline value of 2 ng L^{-1} (PFOA, PFNA, PFHxS, and PFOS), the values <RL were excluded in cases where RL was higher than 1 ng L-1 were excluded. For the rest of the PFASs, values <RL were excluded in cases where RL was higher than 50 ng L⁻¹ were excluded. For all analyses passing these criteria, the values <RL were substituted with 0, and were aggregated (summarized), so that each well-screen is represented by a single value per parameter for the study period. The aggregation was done in two steps: first at a sample level, and then at the well-screen level. Median was used at both steps. The sample-level aggregation was done to ensure that each sample would have equal weight in the aggregation at the well-screen level over the period. This was important, because in some cases replicates were taken, and it is also possible that the same laboratory analysis is reported twice, using both the current and obsolete database code for the specific parameter. Median was used for aggregation at the wellscreen level to avoid bias due to potential outliers.

For the purposes of this study, we calculated percentage of specific land-use within a 200 m buffer zone around each well. The 200 m buffers were calculated in QGIS v. 3.38 (http:// qgis.org). The zonal statistics calculation (percent area per land use type within buffer) was done in R v. 4.4.0 (ref. 15) with the function "exact extract" from the R package "exactextractr" v. 0.10.0 (https://CRAN.R-project.org/ package=exactextractr). This function provides a precise estimation because it considers raster pixels partially covered by polygons. The land-use/land-cover (LULC) raster used for this calculation was Basemap03 (ref. 16) with 10m resolution. For the purposes of this work the following LULC categories were aggregated into two types of LULC:

- (1) "Nature and fresh waters": including the categories 411 000 lake, 412 000 stream, 321 000 nature, dry (habitat types on dry ground, which are not categorised as forest), 322 000 nature, and wet (habitat types on wet ground, which are not categorised as forest); In total this aggregated category has area 3345.4 km² (7.7% of the terrestrial area in Denmark).¹⁶
- (2) "Forest" including the categories: 311 000 forest, 312 000 forest, and wet (forested land on wet ground). In total this aggregated category covers 5634.7 km² (13.1%). 16

Nature and forest

5 km distance

The distance from the wells with PFAS data to the Danish coastline was calculated in OGIS, as the distance to the "nearest hub", where the hubs are the vertices of the coastline shape-file (line type). This could introduce some minor uncertainty in the distance calculation, especially for long straight stretches of the coastline. The distance to the coast for the well-screens included in this study is from 0.1-44.0 km.

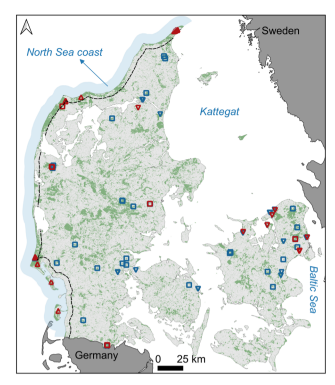
To determine which well-screens are relevant to the North Sea coast, all well-screens within a manually outlined area along the west coast of Denmark were flagged. The flagged wells, which were with <5 km distance from the coastline, were included in the sub-set "<5 km from North Sea coast". Additional supporting data for Cl⁻, Mg²⁺ and Na⁺ concentrations in the wells were also extracted from the Jupiter database on 28 August 2025. The data were quality assured and aggregated in a similar manner to the PFAS data, except there were no values <RL and there was no period restriction. The periods with data were well-screen specific, but overall, the data spanned the period 1966-2025.

The Kruskal-Wallis chi-squared test¹⁷ and post-hoc Dunn test for multiple comparisons18 with Bonferoni adjustment of the pvalue (as implemented in the R package 'FSA' v. 0.10.0, https:// CRAN.R-project.org/package=FSA) were used for testing for significant differences between subsets. Three different tests were performed for testing for significant differences in the two subsets "<5 km from North Sea coast" and the other wellscreens in forest and nature ("other (F&N)": (1) difference in depth to top of well-screens, (2) difference in common PFASs, and (3) difference in the PFOA/PFHxS ratio. The post-hoc Dunn test was used when the Kruskal-Wallis test was positive (significant difference) when testing for differences in more than two subsets, e.g. "<5 km to North Sea coast (F&N)", "<5 km to other coast (F&N)", and ">5 km to coast (F&N)" or "<5 km to North Sea coast (F&N)", "Nationwide dataset", "Nationwide w/o <5 km to North Sea coast (F&N)", and "North Sea marine samples". For all statistical tests, the significance level was set to $\alpha = 0.05$, but we also report exact (adjusted) *p*-values.

Results and discussion

PFASs in Danish groundwater

A total of 1538 shallow groundwater wells fulfilled the criteria to be included in the nationwide dataset. The wells were distributed across the country (Fig. S1). At least one PFAS was detected in 300 of the 1538 wells, with PFOA as the most frequently detected (206 out of 1538 wells corresponding to 13%), followed by PFHxS and PFOS, which were detected in 11 and 10% of the wells. None of the wells in the dataset are deliberately placed near point sources of PFASs, but the wells are placed in all landscape types including urban and industrial areas, and the PFAS-affected wells in the complete dataset are most likely polluted by a variety of diffuse and point sources. To see any effect of a diffuse sea spray aerosol source, we therefore focused the subsequent analyses on 137 wells in the dataset that are located in forests or nature areas where the chance of having PFAS point sources affecting the groundwater is very low (Fig. 1).



Detected PFAS

- <5 km from North Sea coast
- <5 km from other coast
- >5 km from coast

Not detected PFAS

- <5 km from North Sea coast
- <5 km from other coast
- >5 km from coast

Fig. 1 Map with location of the 137 Danish well-screens located in forest and nature (F&N) areas. Not detected (blue) denotes a median value below the detection limit for all 22 PFASs; detected (red) denotes the median value above the detection limit for at least one of the 22 PFASs. Well screens with detection are shown on top and may therefore cover well screens without detection. For more information about "F&N" see the main text. The coast line defined as the North Sea in this study is shown with blue shading, with the dotted line depicting the 5 km distance to the North Sea coast.

For these 137 wells a clear pattern appeared when dividing the wells into coastal and inland wells located less or more than 5 km from a sea shoreline, respectively, and furthermore, dividing the coastal wells into wells located along the North Sea coast or close to other Danish coasts like Kattegat and the Baltic Sea: 52 wells were located in forest/nature more than 5 km from a seacoast, and for these, the detection frequencies were similar or lower as for the whole dataset. 40 wells were located less than five kilometres from seacoasts other than the North Sea and for these, the pattern was similar. However, for the 45 wells located within 5 km from the North Sea coast, almost all wells had detections of at least one PFAA. PFOA, PFHxS and PFOS showed the highest detection frequency (78 to 89%) but PFBA, PFPeA, PFHxA, PFHpA and PFBS were also detected very frequently (in

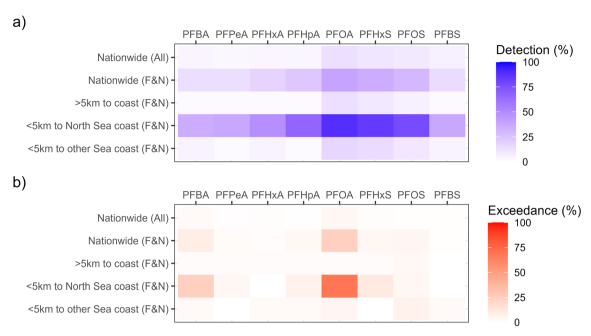


Fig. 2 Heatmap with frequency of detections (a) and exceedances of the 2 ng L $^{-1}$ national threshold limit (b) of most common PFASs in different subsets, where row 1 "Nationwide (All)" is the complete nationwide dataset of shallow wells (n = 1521 to 1538 depending on the compound); row 2 "Nationwide (F&N)" is the subset with the shallow wells located in areas dominated by forest or other nature types (n = 137); row 3 is the subset of "Nationwide (F&N)" containing only the wells located more than 5 km from the coast (n = 52); row 4 is the subset of "Nationwide (F&N)" with less than 5 km to the North Sea coast (n = 45) and row 5 is the subset of "Nationwide (F&N)" with less than 5 km to any other sea coast than the North Sea (n = 40). Below is a similar heatmap of wells where concentrations of the individual compounds exceed the threshold limit for the sum of PFOA, PFNA, PFHxS and PFOS in Denmark (>2 ng L $^{-1}$).

36–67%). For all the compounds shown in Fig. 2, there is a statistically significant difference between the wells close to the North Sea and all other wells in forest/nature areas (Table S7) and they all have significantly higher detection frequency in the wells close to the North Sea using a Kruskal–Wallis chisquared test (Table S7). In Denmark, the threshold value in drinking water is 2 ng L⁻¹ for the sum of PFOA, PFNA, PFHxS and PFOS. For the individual compounds, PFOA very frequently solely exceeded this limit in wells close to the North Sea coast.

The 45 wells located within 5 km from the North Sea coast in general have very little industry or build areas nearby (on average 0.16% of the 200 m buffer, while for the remaining 92 wells located in forest/nature in the rest of the country it is 1.5%). The source of PFASs for the North Sea wells is therefore very unlikely to be local point sources, leaving a diffuse atmospheric source as the most plausible explanation. The question is then if the much higher detection frequency in wells close to the North Sea is simply caused by a faster transport of PFASs? Well depth is often used as a proxy for the transport time from the surface to well screen, but the depth of the screens in the subsets are not different to a degree that would be likely to cause such a big difference in PFAS occurrence (average 17 m for wells <5 km from the North Sea coast and 24 m for the remaining wells); moreover, there is no statistically significant difference (p = 0.08) (SI Table S5). The estimated yearly recharge rate of the wells (Fig. S2a) is also very similar at 324 mm on average for the wells close to the North Sea and 286 mm for the remaining wells.

In addition, for many of the PFAS-polluted wells close to the North Sea, the PFAS concentrations are much higher than one would expect from the general atmospheric deposition. For example, the median concentration of PFOA is 4 ng L⁻¹ and 25% of the wells have 5 ng L^{-1} or higher (maximum 19 ng L^{-1}). For comparison, the concentration of PFOA in precipitation in Denmark has recently been measured to have an average of 0.25 ng L⁻¹, while slightly higher concentrations, typically between 1 and 2 ng L-1, have been measured in the neighbouring country Sweden.10 The most plausible explanation for the observed geographical pattern of PFASs in groundwater is therefore transport from sea water to land with sea spray aerosols, as no other known PFAS source could cause such a pattern. PFASs in sea spray aerosols has not been measured at sea or coastal areas around Denmark, but it is well known that PFASs are present in sea water all over the world, although the measured concentrations vary geographically, in time and from study to study.20 For example, within the sea areas most relevant for aerosol deposition in Denmark, the median seawater concentrations of PFOA between 2010 and 2014 were around 0.1, 0.5, and 1.1 ng L^{-1} for samples from the North Atlantic Ocean, the Baltic Sea, and the North Sea, respectively. A study of sea spray aerosols 20 km from the coast at a site in Southern Norway located only a few hundred kilometres from the Danish North Sea coast, showed some influence of sea spray aerosols on PFAS deposition.6 For a parallel study site in Northern Norway, located just 1 km from the coast, the relationship between PFASs and aerosol tracers (magnesium and sodium) was much

stronger indicating that distance to the coast is important for the strength of sea spray aerosols as a PFAS source on land.6 While aerosol tracers such as Cl⁻, Mg²⁺ and Na⁺ could be less useful for groundwater, since some fractionation may occur during infiltration, the concentration of all three ions are much higher in the 45 wells close to the North Sea coast compared to the rest of the dataset (Fig. S3). Also, the ratio between Mg²⁺ and Na⁺ is on average 0.14 in wells close to the North Sea with little variation (std. dev. 0.03), which is equal to the ratio of 0.12-0.14 found in sea spray aerosols in Norway and the Netherlands. 6,21 The ratio between Mg²⁺ and Na⁺ in the remaining 92 wells was much more varying and having a very different average of 0.47. While the direct relationship between aerosol tracers and PFAS concentration was not very strong (Fig. S4), this is a clear indication that the wells close to the North Sea Coast, of which almost all had detectable PFASs, are affected by sea spray

In addition to the small number of analyses of sea spray aerosols, there are several measurements of PFASs in sea foam, which can be seen as a closely related environmental medium. In the North Sea, high concentrations of a range of PFASs have been found in collapsed/condensed sea foam with PFOA and PFOS as the dominating compounds (1800-73000 ng L⁻¹) and also with PFHxS occurring in high concentration (60-16600 ng L⁻¹).²²⁻²⁴ PFNA and PFDA were also found in sea foam at relatively high concentrations (300–17400 ng L^{-1}) and are probably also deposited inland by sea spray aerosols as was also concluded by Sha et al.6 The reason we do not (yet?) detect PFNA and PFDA in groundwater could be due to higher retention in the soil caused by sorption to organic matter and air/water interfaces. Such retention is probably also causing the relatively lower concentration and detection frequency of PFOS in the groundwater despite an expected historical input with sea spray aerosols of at least the same magnitude as PFOA. On the other hand, PFPeA, PFHxA and PFBS are retained only to a very minor degree in soil, which could explain their frequent detection in wells near the North Sea coast, despite smaller preconcentration factors in sea spray aerosols.3,5

PFOA and PFHxS as indicators of a sea spray aerosol source?

PFOA and PFHxS are both ubiquitously present in sea water across the world, with PFOA typically at about 10 times higher concentration, though this varies with location.20 These two compounds have quite similar enrichment factors in sea spray aerosols3-5 and will be retained in soil also to quite similar degrees both with regards to sorption to organic matter²⁵ and air/water interfaces in soil.26,27 One could therefore hypothesize that the presence of both compounds at concentration ratios not too different from that of the sea water source would be found in groundwater polluted by a diffuse sea spray source. To test this hypothesis, we calculated the ratio between PFOA and PFHxS in all groundwater samples. Both compounds were present in almost all the 45 shallow wells close to the North Sea coast and within these, PFHxS was never present without PFOA, while the opposite was the case for two wells where the PFOA concentration was close to the reporting limit. For the whole

dataset (nationwide), PFHxS was not detected in 35% of the 206 wells where PFOA was detected and PFOA was not detected in 20% of the 164 wells where PFHxS was detected. While even in the whole dataset there is a high overlap between the two compounds, this clearly shows that the overlap is exceptionally high in wells near the North Sea coast where the sea is the most likely source of PFASs. In addition, the ratio between the two compounds is also within a much narrower interval in wells near the North Sea coast compared to the whole dataset (Fig. 3). The vast majority of these wells have a ratio between 2 and 6 while the complete range goes from 0.7 to 13. For the nationwide dataset, most wells are within a ratio from 0.4 to 7 but the complete range goes from 0.05 to 109. The ratios for the wellscreens within 5 km from the North Sea (n = 38 with both compounds detected) are statistically different from the ratios for the rest of the well-screens (n = 94 with both compounds detected) (p = 0.003, Table S8). For comparison, the two compounds have recently been detected in 23 seawater samples taken from surface waters of the North Sea near the Danish Coast. In these samples the ratio was typically between 2 and 6 with a whole range of 2 to 9 (Fig. 3). There is no statistical difference (p = 1.00, Table S8) between the ratios in the surface water marine samples and the coastal (North Sea) groundwater wells. This supports the hypothesis that the presence of PFOA and PFHxS at concentration ratios not too different from that of the sea water would be found in groundwater polluted by

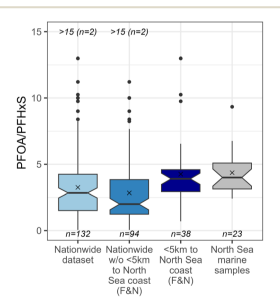


Fig. 3 Ratio between PFOA and PFHxS (ng L^{-1}) in groundwater wells where both compounds were detected. The ratio in 23 marine surface samples from the coastal North Sea taken in 2022 and 2023 is shown for comparison (https://miljoedata.miljoeportal.dk/, data extracted on January 13. 2025, Table S4). F&N - forest and nature; boxplot elements: 25th and 75th percentiles for the hinges (box); whiskers extend from the hinges up/down to the largest value \leq / \geq 1.5 \times IQR (where IQR is the inter-quartile range, or distance between the first and third quartiles); data beyond the whiskers is outlier (black dot); median – horizontal thick line; mean – x symbol; the notches extend 1.58 * IQR/sqrt(n), which is approximately a 95% confidence interval for the medians.

 Table 1
 Selected characteristics and PFAS detections for four monitoring wells located in remote nature areas close to the Danish North Sea coast

Well no.	Distance to the Depth to the top Well no. North Sea (km) of screen (m)	Distance to the Depth to the top Screen North Sea (km) of screen (m) length (t	n)	Screen Depth to GW table Lithology above length (m) last 5 years (m) screen	Lithology above screen	Surrounding vegetation Tritium-helium TFA (in order of dominance) GW age (y) $(\mu g L^{-1})$	Tritium-helium GW age (y)	$TFA \\ (\mu g \ L^{-1})$	TFA PFAS detected in $(\mu g L^{-1})$ latest analysis ^a
23.487-2 1.8	1.8	9.0	1	0.5-1.3	7 m aeolian sand then old seabed sand	Coniferous forest	2	0.25	PFBA, PFOA, PFBS, PFHxS, PFOS, PFPrA, and HFPO-DA
120.236-2 2.7	2.7	13.0	8	1.2–2.5	7 m aeolian sand then old seabed sand	Coniferous forest	16	0.29	PFBA, PFPeA, PFHpA, PFOA, PFBS, PFPeS, PFHxS, PFOS, PFPFA, and PFECHS
120.237-2 4.8	4.8	15.8	2	1.4-2.4	8 m aeolian sand Coniferous then old seabed sand heathland	Coniferous forest, heathland	21	0.22	PFBA, PFPeA, PFHpA, PFOA, PFPeS, PFHxS, PFHpS, PFOS, and PFECHS
148.62-1 3.4	3.4	16.5	17	1.3 to 2.5	5 m aeolian sand Coniferous then old seabed sand heathland	Coniferous forest, heathland	9	0.47	PFBA, PFPeA, PFHYA, PFHPA, PFOA, PFBS, PFPeS, PFHXS, PFOS, PFPA, and PFECHS
Concentra	tion of most domi	Concentration of most dominating PFASs in latest analysis (ng ${\bf L}^{-1}$	est analysis (1	ng L^{-1}), sum of branched and linear	thed and linear				

^a The full name of the compounds and analytical reporting limits can be found in SI Table S2.

2.3 0.53 3.6 4.2

0.96 2.6 2.5 5.8

<0.3 0.64 2.5 0.81

0.61 0.49 <0.3

3.5 8.7 7.1 11.0

<0.3 1.1 0.45 1.9

<0.3 0.43 0.78

4.3 2.2 0.94 4.6

> 120.236-2 120.237-2 148.62-1

23.487-2

PFHxS

PFBS

PFOA

PFHpA

PFBA

a diffuse sea spray source. In sea foam sampled close to Denmark, the PFOA/PFHxS ratio has been found to be between 3 and 11,22-24 which also fits with the ratios we observe in Danish groundwater between these two compounds. In conclusion, PFOA seems to always be detected in wells polluted by sea spray aerosols and unless the concentration of PFOA is very low, PFHxS will also be detected, typically at 2-6 times lower concentration. It should be emphasized, though, that although much more common near the North Sea coast, this ratio is also seen in wells further inland with other expected sources of PFASs, and a PFOA/PFHxS ratio in this interval therefore should not in itself be used as a strong indication of the source.

Shallow coastal monitoring wells

33 of the 40 PFAS polluted wells near the Danish North Sea coast are groundwater abstraction wells used to provide drinking water. Due to long well screens combined with pumping of large volumes that may vary over time, abstraction wells often represent a mixture of water with different origins and ages, which complicates interpretation of origin and age of pollution with organic micropollutants.28 In the Danish groundwater monitoring system, we have four PFAS-polluted wells near the North Sea coast that are in nature areas with no other suspected source of PFASs nearby, except for atmospheric sources including sea spray aerosols. For all four wells there is more than 1 km to the nearest area with registered activity that could potentially lead to any kind of soil pollution, including PFASs and there are more than 500 m to the nearest house (more than 1 km for three of the wells). In all four wells, the well screens are short (1 to 3 m) and

shallow (9-17 m below surface) and there is no pumping of water, except when the wells are sampled. All four wells are situated in quite similar geological and geographical settings, in very sandy sediments, with shallow unconfined groundwater tables and with coniferous trees (spruce and pine) as the dominant vegetation (Table 1 and Fig. S5). These wells are therefore expected to represent a constant and relatively small recharge area over time and the groundwater age (recharge time) would be expected to have a small interval and to be constant. Using the tritium-helium tracer method we estimated the groundwater recharge time, to be from a few years in the well with the youngest water to around 20 years in the well with the oldest water. Concentrations between 0.22 and 0.47 $\mu g L^{-1}$ of the ultra-short chain PFCA trifluoroacetate (TFA) that functions as a tracer for young groundwater¹³ confirm that the groundwater has recharged recently (Table 1).

The four compounds PFOA, PFBS, PFHxS and PFOS were the most frequently detected short- and long-chained PFASs and were repeatedly detected in all four wells (Fig. 4). Over time, there is no clear trend in concentration in the four wells, though PFOS was in all cases the highest at the last sampling point. PFOS will expectedly have the highest retardation through the soil layers above the groundwater wells, since it has the highest soil: water and air: water partitioning coefficients among the four compounds.29 If an increase in PFOS is currently occurring, this does not necessarily imply an increase in the source compared to the other PFASs but could be caused by a delayed arrival in the groundwater. The fact that PFAS concentrations are not yet decreasing even in the very young groundwater of the

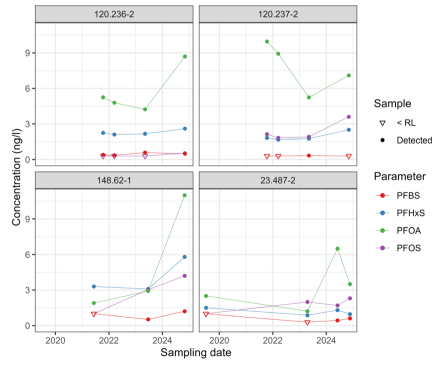


Fig. 4 Time series for repeatedly detected PFAS compounds in four wells with the sea spray aerosols from the North Sea as the suspected source. Further details of the individual wells are shown in Table 1. Open triangles denote that the compound was not detected at that sampling date with the concentration depicting the Reporting Limit (RL) of that analysis.

four monitoring wells, suggests that PFASs from sea spray aerosols may be found in years to come in coastal groundwaters with longer travel times than in these four wells. This seems to be the case even if the PFASs are banned or phased out like PFOA and PFOS, a ban that has resulted in a significant decrease in these compounds from 2007 to 2017 in sea water in the southern part of the North Sea.³⁰ It is more likely that the concentrations in coastal groundwater in general can be expected to increase in the years to come. At the same time, longer-chained PFAAs with higher retardation factors in soil such as PFNA and PFDA, which have recently been found at relatively high concentration in water and foam from the North Sea^{22–24,30} and in sea spray aerosols in Norway,⁶ may appear.

To see if other PFASs than the usual legacy PFAAs can be found in groundwater wells with sea spray aerosols as the most likely source of PFASs, the four wells were sampled and analysed for 60 PFAS compounds (Table S2). Three additional compounds were found: The ultrashort-chain PFCA PFPrA (perfluoropropanoic acid) was found in three of the wells in concentrations from 7-14 ng L⁻¹, PFECHS (perfluoro-4ethylcyclohexane-sulfonic acid) was found in the same three wells at lower concentration (0.4-0.9 ng L⁻¹) and HFPO-DA (GenX) was found in one well at a concentration of 0.2 ng L^{-1} . The concentration of these three compounds in sea water is little investigated but HFPO-DA and PFECHS have been detected in the seas around Denmark.30 HFPO-DA was found at higher concentration than PFECHS in sea water, but at air/water interfaces, a relatively larger enrichment factor can be expected for the long-chained PFECHS, compared to HFPO-DA and PFPrA. Indeed, PFECHS has been detected in sea foam at much higher concentration than in corresponding bulk water,²² supporting that the PFECHS detected in the three groundwater wells could be derived from sea spray aerosols. HFPO-DA has also been found in sea foam, with a highly varying enrichment factor of 1 to 410 compared to bulk sea water.31 It is unknown if PFPrA will be preconcentrated to a level that sea spray aerosols may be the main cause of the detected PFPrA, but there is evidence that PFPrA can be present in precipitation at relatively high concentration, possibly as a degradation product of HFCand HCFC-gases and other volatile PFCA precursors. 10,32

Concluding discussion

All in all, the most plausible explanation for the high concentrations of certain PFAA compounds (PFOA, PFHxS, PFOS, etc.) in shallow groundwater near the Danish North Sea coastline is transport from sea to land via sea spray aerosols. While this transport process has been previously demonstrated, the fact that it may pollute groundwater to a level above drinking water guideline values has not been demonstrated previously in the scientific literature. On the other hand, there is no clear sign of sea-derived PFAS pollution in groundwater wells in other parts of the country, despite many groundwater wells close to other coastlines. Likely reasons for this observation could be (a) less PFASs in other sea bodies, (b) less wave energy along other coastlines leading to less preconcentration in sea spray aerosols, (c) different geological settings leading to less or slower

transport of PFASs to groundwater and (d) different land use in the groundwater recharge areas such as agricultural land or deciduous forests instead of coniferous plantations that dominated most affected wells along the North Sea coast. We find explanation (a) less likely, as the PFAS concentration in the Baltic Sea is known to be of the same magnitude as that in the North Sea^{20,30} and recently similar levels of PFASs were found in sea foam from the Baltic Sea just southeast of Denmark²⁴ and from the North Sea. The other three explanations seem more likely and may work in combination as the reason that we do not (yet) see PFAS contamination in groundwater caused by sea spray aerosols in other parts of Denmark than along the North Sea coast. Concerning explanation (b), the prevailing wind direction in Denmark is between Southwest and Northwest and most strong wind fields come from those directions. Strong wind fields also arrive from easterly directions, but these are less frequent and on a yearly basis, and the wave energy is much higher on the North Sea coast.33 The North Sea coast may hence experience both more long-range transport of PFASs in sea spray aerosols, as well as a larger contribution from the surf zone, which can be a major contributor of sea spray aerosols tens of km inland.34,35

The combination of soil type (c) and land use (d) could also influence, as the North Sea coast is dominated by aeolian sand combined with shallow unsaturated zones and coniferous plantations. Although sandy soils are common in many other areas of Denmark (Fig. S2b) and coniferous plantations and shallow unsaturated zones are also common, a combination of these features is not common along other coasts in Denmark. This combination may promote transport of PFAAs like PFOA to the groundwater compared to other areas due to three reasons: (1) water saturated coarse sand shows little retention of PFOA, 36,37 (2) sorption to air/water interfaces in the unsaturated zone will be of less importance compared to areas with deeper unsaturated zones and (3) conifers may effectively filter out the aerosols carrying the PFASs.38,39 Future research should address these possible explanations for the clear PFAS signal along the Danish North Sea coast and the lack of signal along other coastlines. The observed diffuse PFAA pollution in coastal groundwater is most likely a world-wide phenomenon, as PFAAs are present in sea water across the world.20 However, additional studies are needed to achieve a sufficient understanding of the drivers, so that current and future concentration levels of different PFAS compounds in coastal groundwater wells may be predicted.

Author contributions

C. N. A.: conceptualization, investigation, methodology, formal analysis, visualization, writing – original draft. D. V.: data curation, formal Analysis, visualization, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Data availability

This study was carried out using publicly available data from the Danish National well database (Jupiter) at https://www.geus.dk/produkter-ydelser-og-faciliteter/data-og-kort/national-

boringsdatabase-jupiter (accessed on 30th August 2024). Additional data for 60 PFASs in 4 monitoring wells have been included as part of the SI. Results for noble gas analyses for estimation of groundwater age have been included as part of the SI. Results of in-house-analysis of trifluoroacetic acid (TFA) are in Table 1. Publicly available data for PFOA and PFHxS in sea water have been included as part of the SI. Supplementary information is available. See DOI: https://doi.org/10.1039/d5va00181a.

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