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Advances in Distinguishing Groundwater Influenced by Oil Sands Process-Affected Water (OSPW) from Natural Bitumen-Influenced Groundwaters

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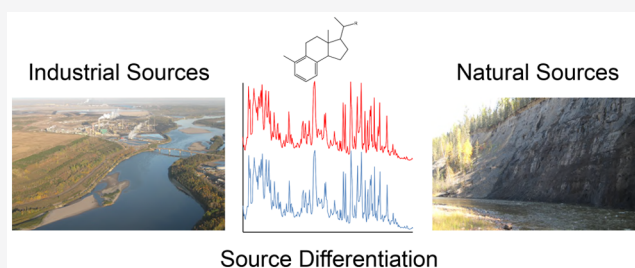


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Supporting Information

ABSTRACT: The objective of this study was to advance analytical methods for detecting oil sands process-affected water (OSPW) seepage from mining containments and discriminating any such seepage from the natural bitumen background in groundwaters influenced by the Alberta McMurray formation. Improved sampling methods and quantitative analyses of two groups of monoaromatic acids were employed to analyze OSPW and bitumen-affected natural background groundwaters for source discrimination. Both groups of monoaromatic acids showed significant enrichment in OSPW, while ratios of O₂/O₄ containing heteroatomic ion classes of acid extractable organics (AEOs) did not exhibit diagnostic differences. Evaluating the monoaromatic acids to track a known plume of OSPW-affected groundwater confirmed their diagnostic abilities. A secondary objective was to assess anthropogenically derived artificial sweeteners and per- and polyfluoroalkyl substances (PFAS) as potential tracers for OSPW. Despite the discovery of acesulfame and PFAS in most OSPW samples, trace levels in groundwaters influenced by general anthropogenic activities preclude them as individual robust tracers. However, their inclusion with the other metrics employed in this study served to augment the tiered, weight of evidence methodology developed. This methodology was then used to confirm earlier findings of OSPW migrations into groundwater reaching the Athabasca River system adjacent to the reclaimed pond at Tar Island Dyke.



INTRODUCTION

The Alberta oil sands deposit in Canada is the third-largest proven oil reserve in the world, comprising 142000 km^{2,2}. Mined oil sands undergo a caustic hot water extraction process to separate bitumen from the sand. The byproduct of this extraction, known as oil sands process-affected water (OSPW), is a complex mixture of dissolved inorganic and organic compounds. The organics fraction includes naphthenic acids (NAs) for which the aquatic toxicity has been well documented,^{3–9} although the chemical classes causing toxicity within the greater group of acid extractable organics (AEOs) have yet to be established.

To date, the oil sands industry has not treated and released OSPW and has operated under a zero-discharge practice with accumulated OSPW stored in large containments (tailings ponds), comprising a total area of >130 km².¹⁰ Despite containment infrastructures (interceptor wells, ditches, and relief wells), designed to capture and return seepage to the ponds,¹¹ the question remains as to whether OSPW-affected groundwaters migrate beyond such containments,^{12–14} reach surface waters,^{15–21} and cause adverse effects. Seepage into

groundwater from containments adjacent to the Athabasca River or its tributaries are of concern because if seepage is pronounced, there is potential for aquatic life exposures to OSPW-derived toxic substances.

In the case of the Mildred Lake Settling Basin (MLSB) containment, an OSPW-affected groundwater plume has been distinguishable from ambient local groundwater by elevated concentrations of routinely measured parameters, including salts or NAs.¹³ However, the detection of potential seepage at other ponds is hindered by several factors. First, inorganic and organic OSPW compositions closely resemble those in natural bitumen-affected groundwater within the McMurray geological formation.^{16,20,21} Further, as the caustic aqueous extraction of bitumen does not require the addition of industrial additives, potential tracers of OSPW migration are not readily available.

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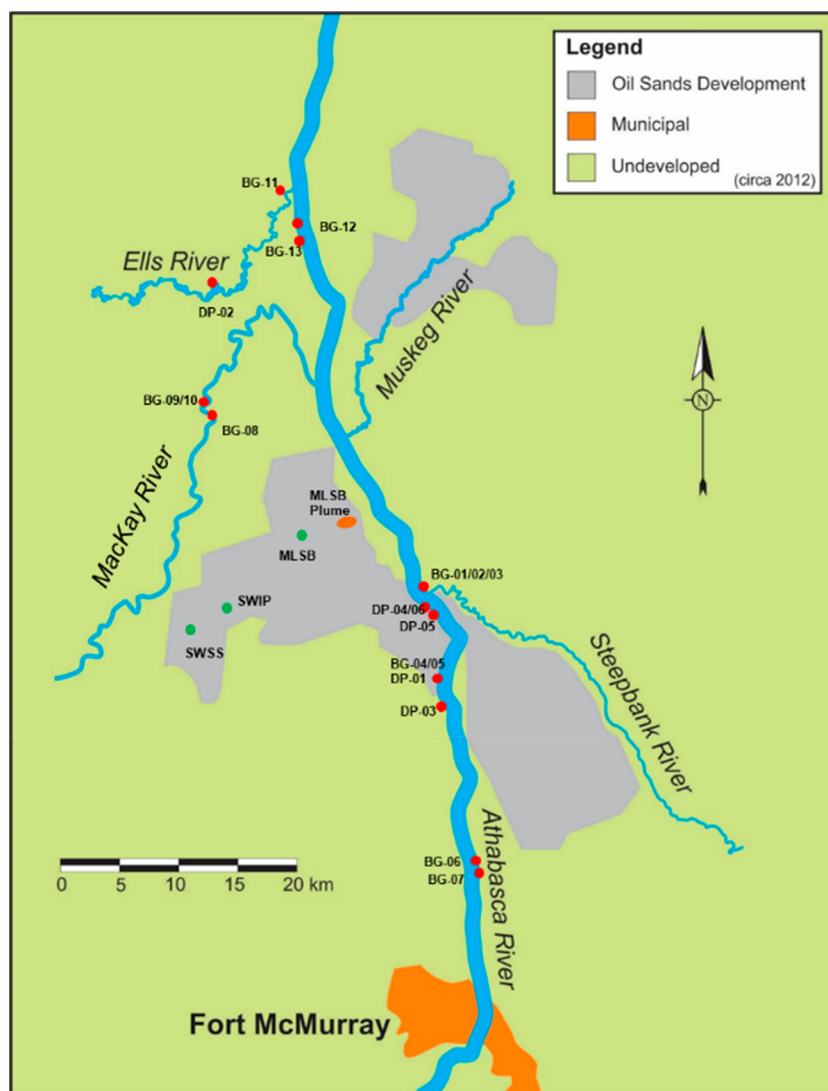


Figure 1. Overview map of the study area of the oil sands region north of Fort McMurray, AB, Canada, showing sampling sites for OSPW (green symbols), the MLSB plume (orange oval; individual well locations not shown) and riparian groundwater (red dots) comprising Background (BG-1 to 13; DP-1 to 3) and Unknowns (DP-4 to 6) by Tar Island Dyke (TID). Sample details are provided in Table 1.

Finally, despite recent discoveries of adamantane,^{22,23} bicyclic NAs,^{24,25} and alicyclic, aromatic, and sulfur-containing acids^{23,26} in OSPW, there are few commercially available authentic standards of OSPW constituents, hampering efforts to develop and standardize analytical methodologies for NAs and seepage detection.

In previous attempts to identify OSPW-affected groundwater, a wide range of chemical metrics has been applied. These have included: measurement of total NAs, arsenic, boron, fluoride, molybdenum, selenium, vanadium, nickel, rhenium, ammonium, sodium, and chloride and their ratios, characterization of water types (Piper plots), use of synchronous fluorescence spectroscopy (SFS) to determine aromatic species, monitoring of a variety of isotopic tracers from ²H and ¹⁸O of water to dissolved ions and to measurement of intramolecular $\delta^{13}\text{C}$ signatures, determination of the ratios of O₂/O₄ heteroatom classes, and the presence of SO_n containing species in OSPW.^{13,17,18,20,27–38} Since 2005, with advances in available analytical instrumentation and an improved understanding of the organic compositions of OSPW,^{22,24–26,39,40} studies on NAs as tracers of OSPW

migration have expanded to include the broader class of AEOs and have progressed from standard gas chromatography–mass spectrometry (GC-MS) methodologies¹⁸ to deployment of state of the art instrumentation, including high performance liquid chromatography-time-of-flight mass spectrometry (HPLC-ToF-MS¹⁹), HPLC-high resolution mass spectrometry (HPLC-HRMS²¹), comprehensive multidimensional GC-MS (GC × GC-MS^{16,41}), and Fourier transform-ion-cyclotron-resonance mass spectrometry (FT-ICR-MS^{29,34,35,42}). It is important to note that although the ability to characterize bitumen influence has evolved through these studies, the differentiation of OSPW-affected groundwaters from natural bitumen sources has not been achieved. The analytical challenge in source discrimination of these inherently similar, so-called “supercomplex”⁴³ mixtures, is considerable.

The only study to assert some success in this endeavor was that of Frank et al.¹⁶ that applied a tiered, weight of evidence approach that included routine analytical methods (Level-1) and two advanced methods for profiling AEOs (Level-2) to directly compare each source. One Level-2 method used GC × GC-TOF-MS to identify two groups of monoaromatic

naphthenic acids (denoted “Family A and B”) shown to be elevated in OSPW and OSPW-influenced groundwater. The other Level-2 method used high resolution mass spectrometry (HRMS) profiling of AEOs to associate OSPW influence with a greater proportion of O₂ heteroatom ion classes relative to O₄ species, although subsequent studies have shown conflicting evidence regarding such ratios to differentiate source.^{21,34,35} Both Level-2 methods demonstrated source differences for two samples of shallow groundwater collected adjacent to Tar Island Dyke (TID) at the edge of the Athabasca River, which the authors concluded were likely influenced by OSPW.¹⁶ While this study showed promise in differentiating groundwaters affected by OSPW from those influenced by natural bitumen sources, limitations included the extent of sampling within each source type and the qualitative capability of the Level-2 analyses.

The primary objective of the present study was to explore further the utility of the Family A and B groups of monoaromatic acids, and ratios of O₂/O₄ classes, to distinguish OSPW-affected groundwater from ambient groundwater influenced by the natural bitumen deposits of the McMurray Formation. This was accomplished through sampling and analysis improvements that included the (i) use of larger sample volumes (>1000 fold) to facilitate the greatest possible level of detection; (ii) quantitation of the Family A and B acids; (iii) use of a larger set of OSPW (2 different mining operations per parameter); (iv) use of a larger set of natural “background” groundwaters from the area where the McMurray formation outcrops, including several that were fairly saline; and (iv) validation of the combined methodology against a known plume of OSPW-affected groundwater.¹³

A secondary objective was to assess whether anthropogenic artificial sweeteners and per-and polyfluoroalkyl substances (PFAS) have diagnostic capabilities as tracers of OSPW seepage. Both groups are hypothesized to have some diagnostic capabilities due to their association with anthropogenic activities, persistence and mobility in groundwater, and utility in contaminant source apportionment.^{44–48} The final objective was to reassess the presence of OSPW-affected groundwater at select locations at the edge of the Athabasca River adjacent to TID, as indicated by Frank et al.,¹⁶ using the improved methodology.

METHODS

Sample Collections. Sample locations are presented in Figure 1 and sampling details are summarized in Table 1. OSPW samples were collected with stainless dippers into precleaned stainless steel containers fitted with Viton seals or glass bottles, with assistance and access provided by the respective oil sands operators’ personnel. Shallow riparian groundwater samples were collected using a stainless steel drive-point system⁴⁹ at depths of 30–120 cm below the streambed of the Athabasca River and associated tributaries (Figures S1–S3). The locations (e.g., edge of river, outside of meanders) and timing (e.g., low river flow periods) of sampling were all chosen to best ensure collection from groundwater discharge zones. Furthermore, the drive point was pushed deeper prior to sampling if the water quality parameters indicated a similarity to surface water, which might indicate hyporheic exchange. Sampling included a repeat sampling of 3 background sites (>1 km from any OSPW influence) and 3 sites adjacent to TID (“unknowns”) initially sampled by Frank et al.¹⁶ and was followed by collections from 13 additional

Table 1. Sets of samples collected for this study (locations shown in Figure 1)^a

Sample Groupings	Number of samples	Notes
OSPW	5 ponds sampled from 2 operators	1. Syncrude Mildred Lake Settling Basin (MLSB); September 2010 & December 2013 2. Canadian Natural Resources Limited (CNRL); September 2011 3. Syncrude West In-Pit (WIP); September 2011 4. Syncrude Southwest In-Pit (SWIP), August 2015 5. Syncrude Southwest Sand Storage (SWSS), September 2015
Background Groundwater	16	13 new locations (BG, within McMurray formation, near surface) (August 2013) Resampled Drive-points (DP) 1–3 from Frank et al. ¹⁶ (September 2012)
MLSB plume (Figure 3)	12	In Plume: 01A, 01B, 02B, 02C, 03B, 03C, 04B Nonplume: 03A, 04A, 05B, 05C, 06A All sampled December 2013
Unknowns by TID	3	Resampled DP 4–6 from Frank et al. ¹⁶ (September 2012)

^aIncluding oil sands process water (OSPW) from several tailings ponds, shallow riparian groundwater (Background and Unknowns by Tar Island Dyke), and groundwater from wells near Mildred Lake Settling Basin (indicated as inside/outside the OSPW plume¹³).

background sites. Finally, MLSB OSPW and groundwater from 6 sets of nested wells (each with wells at 1–3 depths) along the MLSB plume¹³ were collected (Figures 1 and 3). All groundwater sampling commenced following purging of the well or mini-profiler system and equilibration of field-parameters (temperature, electrical conductivity, pH, dissolved oxygen) measured with hand-held meters. No surface water sampling was conducted.

Samples for standard geochemistry (i.e., ammonium, major ions, metals, alkalinity) and artificial sweeteners were collected in separate bottles and preserved as described (Table S1). In addition, a large volume sample for Level-2 profiling, total NAs, SFS, and PFAS analysis was collected in an 18 L stainless steel collection vessel fitted with a Viton seal. All samples were refrigerated on the day of collection, shipped at 4 °C to Burlington, ON within 1 week of collection, and stored at 4 °C. Subsampling and extraction were conducted within 1 week of arrival.

Screening (Level-1) Analyses. Level-1 analyses included geochemical parameters comprising determination of anions (including chloride, sulfate, and nitrate) analyzed by ion chromatography, major cations (including sodium and calcium) analyzed by direct aspiration using an inductively coupled argon plasma mass spectrometry, and ammonium analyzed by spectrophotometry (absorbance of phenol-hypochlorite at 640 nm¹⁶). Alkalinity was determined by titration method 2320.⁵⁰ Samples were analyzed for a suite of trace metals at Environment and Climate Change Canada’s National Laboratory for Environmental Testing (NLET) (Burlington, ON) using inductively coupled plasma-sector field mass spectrometry (SOP 2003). Level-1 analyses also included determination of total AEO concentrations (referred to subsequently as NAs). Low resolution ESI-MS analyses for total NAs were conducted with a Quattro Ultima (Waters

Corp., Milford, MA) triple quadrupole mass spectrometer equipped with an ESI interface operating in negative-ion mode, as described by Frank et al.¹⁶ Additionally, expected maxima in an SFS profile associated with previously identified mono- and diaromatic acids³⁹ were analyzed as described previously.⁵¹ In this investigation, samples that exhibited the characteristic bitumen profile with three maxima with a signal intensity of 100 at 272 nm were identified as positive for this profile.

Advanced Separation and High-Resolution (Level-2)

Analyses. Samples were extracted for detailed profiling of acidic and neutral bitumen organics by ESI-MS and ESI-HRMS. Quantitative determinations of monoaromatic acid Family A (expanded to 8 isomers, using m/z 145) and B acids (2 isomers, using m/z 237, 310), as their methyl esters, were conducted by comprehensive two-dimensional gas chromatography with time-of-flight mass spectrometry (GC \times GC-TOF-MS). For all OSPW samples, 2.0 L volumes were utilized due to the elevated concentrations expected, whereas 15–18 L were used for all groundwaters. Prior to extraction, samples were acidified to pH 2 with 12.0 M HCl, and extractions were conducted using two 900 mL dichloromethane aliquots, each with 1 h thorough mixing within sample collection vessels. Each extract was subsequently evaporated to dryness under a stream of N₂ and reconstituted to an initial volume of either 1.0, 5.0, or 10.0 mL in dichloromethane, depending on source. Aliquots (0.5–1.0 mL) were withdrawn and methylated with freshly prepared diazomethane for GC \times GC-TOF-MS analysis of monoaromatic acids. The remaining extracts were then adjusted to either 5.0 or 1.0 mL; a 500 μ L aliquot was removed, solvent exchanged into methanol, and split evenly for (i) ESI-MS analyses of total NAs with profiling by ESI-HRMS and (ii) untargeted profiling by LC-QToF.⁵²

Level-2 AEO profiling of sample extracts using ESI-HRMS was performed on a LTQ Orbitrap Elite mass spectrometer (Thermo Fisher Scientific, San Jose, CA) using electrospray ionization in negative ion mode.¹⁶ Class distributions were determined using acquired accurate mass data and Composer version 1.0.2 (Sierra Analytics, Inc. Modesto, CA) with an O₂ mass error of 0.065 ppm.

GC \times GC-TOF-MS Analysis of Monoaromatic Acids.

Quantitative analysis of the Family A and B monoaromatic acids¹⁶ were conducted on methylated extracts using an Agilent 7890A GC (Agilent Technologies, Wilmington, DE) fitted with a Zoex ZX2 GC \times GC cryogenic modulator (Zoex Corporation, Houston, TX) interfaced with an Almsco BenchTOFdx TOF-MS (Almsco International, Llantrisant, Wales, UK). The primary column was a 60 m \times 0.25 mm \times 0.25 μ m Rxi-1 ms (Restek, Bellefonte, U.S.A.), followed by a 1 m \times 0.1 mm deactivated fused silica modulation loop. The secondary column was a 2.5 m \times 0.1 mm \times 0.1 μ m BPX50 (SGE, Melbourne, Australia). Samples (1 μ L) were injected at 300 °C splitless using a helium carrier at 1.0 mL/min. Oven programming was from 120 °C (5 min hold) to 250 °C at 15 °C/min, then to 340 °C at 2.5 °C/min, and held for 10 min. The hot jet was programmed to track the primary oven at 100 °C above. The modulation period was 3 s. The MS transfer line temperature was 300 °C and ion source 275 °C. Data processing was conducted using GC Image v2.3 (Zoex, Houston, TX, U.S.A.) with the CLIC (Computer Language for Identifying Chemicals) expression tool.^{26,53}

For this study, quantitation of the Family A and B acids was calibrated on one isomer isolated from an OSPW sample (WIP, sampled in 2009) using preparative GC. Following a

protocol used by Ahad et al.,²⁷ the OSPW extract was repeatedly injected into an Agilent 7890A GC equipped with two 30 m \times 0.5 mm i.d. DB-5MS columns (0.5 μ m film thickness) and a flame ionization detector (FID) coupled to a Gerstel (Mülheim an der Ruhr, Germany) preparative fraction collection (PFC) system. Approximately 5% of the eluent was directed to the FID, with the remainder sent to the PFC. The isolated subfractions were recombined and reinjected into the PFC several times to improve purity. Subfractions were purified by silica gel and analyzed by GC-MS to verify purities.

Anthropogenic Chemicals Analyses. Artificial sweeteners (acesulfame, saccharin) were analyzed with a Dionex 2500 ICS ion liquid chromatography system combined with an Applied Biosystems AB Sciex Q Trap 5500 triple quad mass spectrometer using an electrospray ionization source in negative mode.⁴⁴

Per- and polyfluoroalkyl substances were concentrated from aqueous samples using weak anion exchange solid phase extraction (SPE) following previously published protocols,⁵⁴ and concentrations were adjusted for surrogate recoveries (Tables S2–S4). While the full suite of perfluoroalkylcarboxylates (four to 14 carbons) and perfluoroalkylsulfonates (C₄, C₆, C₈, C₁₀) were determined, we focused on perfluorobutane sulfonate (PFBS), perfluorohexane sulfonate (PFHxS), and perfluorooctane sulfonate (PFOS) to highlight sample trends.

Statistical Analysis. Mood's median test was used to test equality of the background and OSPW group medians, as a nonparametric alternative to a one-way ANOVA or *t* test that is more robust against outliers than a Kruskal–Wallis test.⁵⁵ Differences between background and OSPW group means were tested for significance using *t* tests assuming unequal group variances. To improve normality of the data and equality of group variances, data were transformed by $\log(x+x_{\min}) - \log(x_{\min})$, which is appropriate for data with zeros whose smallest positive value is not close to 1.⁵⁵

To assess joint variation among 32 samples from the 4 sample groups (Table 1), Principal Components Analysis (PCA) was conducted on a correlation matrix computed for 31 chemical variables (Level-2 parameters and a select group of Level-1 parameters). Data for all variables except O₂/O₄ ratio were log-transformed to improve normality of distributions. Significance of eigenvalues was determined by the broken-stick test.⁵⁵

RESULTS AND DISCUSSION

Evaluation of Previous Level-2 Metrics with OSPW vs Background Groundwater.

The first step in advancing our previous methodology was to evaluate the source discrimination ability of our previously proposed Level-2 metrics¹⁶ against an expanded set of samples from OSPW and natural groundwater sources. A comparison of background sites and OSPW ponds, analyzed for the Family A and B acids and ratios of heteroatomic ion classes (O₂/O₄), is presented in Figure 2.

Consistent with the qualitative data from Frank et al.,¹⁶ the quantitative analyses of Family A and B acids showed both are present naturally. Here, both groups showed clear enrichment in OSPW (Family A, 148-fold; Family B, 38-fold based on means), with no overlap in the ranges. Further, the results of the Mood Median tests on untransformed data and *t* tests on transformed data (Table S5) both indicated significant differences between the OSPW and background samples for the sum of Family A isomers, sum of Family B isomers, and all

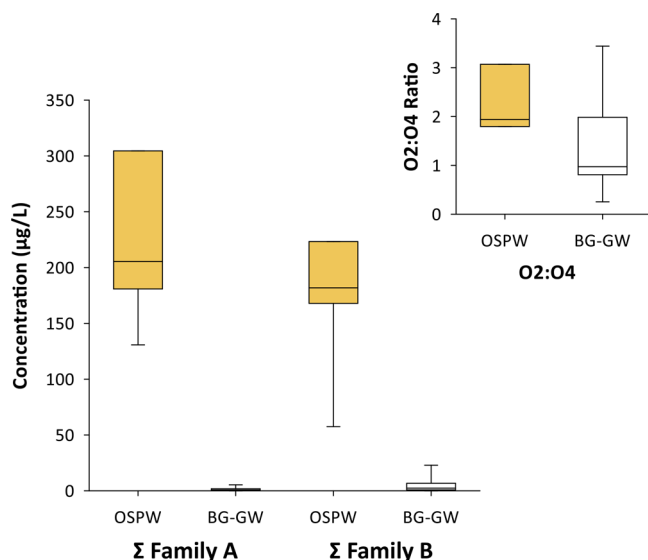


Figure 2. Box plots showing mean and middle quartile (box) and range (whiskers) for total Family A and B monoaromatic acid concentrations and HRMS speciation ratios of O₂- and O₄-containing species compared between OSPW ($n = 4$ Family A-B, $n = 3$ O₂/O₄) and background groundwater (BG-GW; $n = 16$) samples.

individual A and B isomers (8 and 2 isomers, respectively). There was clearly less difference for Family B, as the lowest value for OSPW was only 2.5-fold larger than the maximum of the background groundwater samples; likewise the t test T -value was of lower magnitude for Family B (8.3, compared to 15.4 for Family A). When the individual components were considered, those showing the greatest difference between OSPW and background groundwater (based on the T -value; Table S5) were, in order, isomers A5, A3, A1, A4, with some separation then to A2, A7, A6, A8, and finally B1 and B2. Note also that isomers A5 and A3 were only found above detection limits in two background samples (BG-3, BG-13), whereas all other components were found in four or more background samples (Table S6). The higher concentrations of the Family A and B acids in OSPW are likely from the caustic bitumen extraction and recycling of water (including evaporative enrichment³¹) that leads to a general enrichment of AEOs as a class.

In contrast, ratios of heteroatomic ion classes containing O₂/O₄ species from OSPW and background groundwater overlapped, though the OSPW data set was shown to be significantly higher than that of the background groundwaters with the Mood Median Test and t test on transformed data. Both sets had one or more values considered high (>1). While this may be related to the differing bitumen contents within the set of background samples (Figure S4), based on the extensive sampling conducted in this study, the ratios of O₂/O₄ species do not clearly differentiate OSPW from background bitumen-affected groundwater. This is consistent with Sun et al.²¹ who reported high O₂/O₄ ratios for OSPW (1.2 to 1.8) but also for far-field natural groundwater (A5w-GW; ratio of 1.3). Similarly, Yi et al.³⁴ also reported a wide range of O₂/O₄ ratios that overlapped among sample types.

Given these results, we next sought to evaluate the ability of the Family A and B compounds to identify known OSPW-affected groundwater. For validation, we sampled the monitoring well network of the known OSPW-affected groundwater plume emanating from MLSB (tailings pond;

Figure 3. Locations deemed “in plume” (i.e., 01A, 01B, 02B, 02C, 03B, 03C, 04B) and outside the plume (i.e., 03A, 04A,

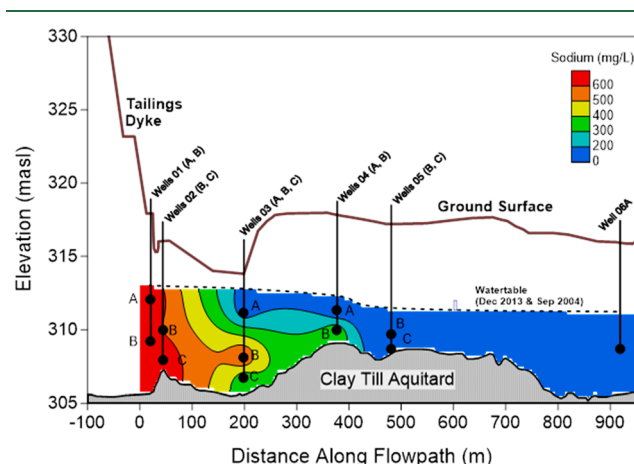


Figure 3. Schematic cross section of the Mildred Lake Settling Basin (MLSB) plume showing: base of constructed tailings dyke; ground surface elevation; well and associated sample intake locations; water table location measured in December 2013 and September 2004;¹⁵ and contour of December 2013 data as defined by sodium concentration (c.f.¹⁵). Note that plume contours and inferred extent differed somewhat for the different chemical tracers considered. Elevations in meters above sea level (masl). Modified after.¹⁵

05B, 05C, 06A; “non-plume”) have not changed from what was first reported,¹³ based on total NAs, sodium (Figure 4) and other Level-1 metrics (i.e., SFS, boron, fluoride; Figure S5). While this plume can be distinguished from the surrounding and relatively pristine groundwater by routine analyses (i.e., major ions, ammonium, and total NAs), it is nevertheless important that potential OSPW-tracers, such as the Family A and B compounds, be assessed in their ability to identify OSPW-affected groundwater throughout the length of the plume. A robust OSPW tracer should be able to match OSPW of present and historical conditions. Here, changes across the extent of the plume could reflect changes in pond composition over the past few decades (i.e., locations closer to the source MLSB representing more recent OSPW conditions; further locations representing more historical conditions). Second, an ideal OSPW tracer would not be slowed substantially by sorption or removed by degradation processes⁵⁶ during transport through the aquifer.

The monoaromatic Family A and B acids showed good utility for identification of the plume (Figure 4A) paralleling the concentrations of total NAs and sodium (and chloride, Figure 4B), where the elevated concentrations at well 4A, may reflect faster transport of this conservative ion. The leading edge of the plume is indicated at well site 4 where the deeper 4B sample is the last sample to show OSPW-affected groundwater by these metrics. The absence of detectable Family A and B acids in samples from wells 5 and 6 combined with the lower concentrations present in samples from shallow wells 3A and 4A also reflect the nature of the background at this site, which is affected minimally by natural bitumen (Figure 4A¹³). Interestingly, when expressed relative to the totals measured in the OSPW sample from MLSB, the Family A acids were generally reduced throughout the plume, whereas the Family B acids were consistent or elevated. Further differences were apparent between the individual components (Figures S3, S4). For instance, isomers A1–A4 and especially

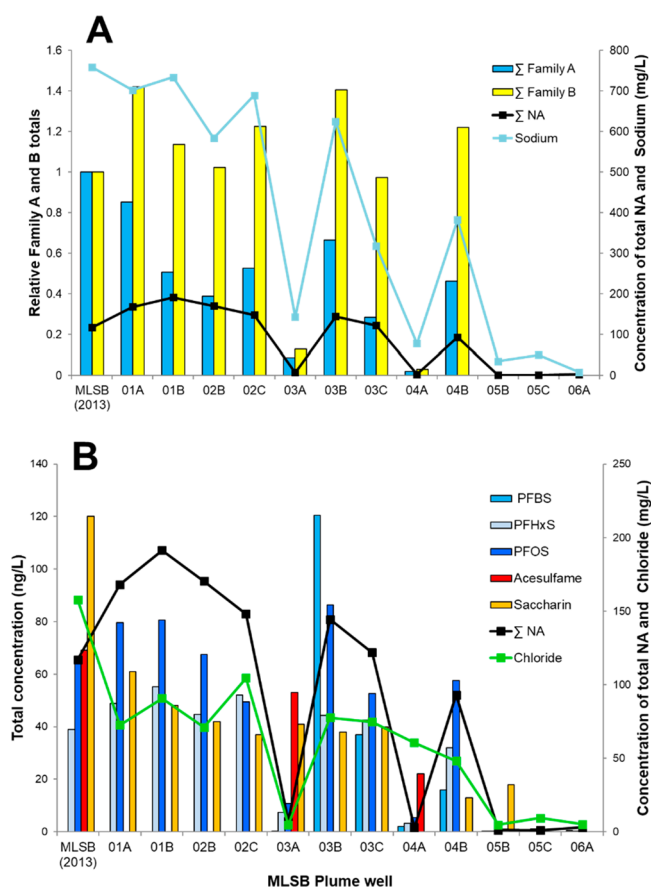


Figure 4. Total Family A and B monoaromatic acids (A), artificial sweeteners (acesulfame and saccharin) and PFAS surfactants (PFBS, PFHxS, PFOS) (B) present in well samples of the MLSB plume monitoring network and the OSPW-source MLSB tailings pond and are arranged from left in direction of flow path. Panel A depicts total monoaromatic acids expressed relative to the source OSPW. Results are plotted against Total NAs, (Σ NA, black line), chloride- and sodium+ ions (green and blue lines, respectively), which have been previously shown to delineate the plume at this site.¹³

A6 were found throughout the plume at levels only slightly lower than the source OSPW (Figure S5), attributable to dilution, biodegradation/sorption, or enrichment in the source OSPW over time. In contrast, isomers A5, A7, and A8 were at much lower levels or absent from wells at the front of the plume (e.g., 3C, 4B; Figure S6), possibly due to sorption or biodegradation within the aquifer.⁵⁶ Alternatively, this may reflect a historical change in OSPW composition, making them less diagnostic. Of the Family B components, B2 showed a general enrichment relative to the source OSPW, while B1 was consistent (Figure S7). It is possible that B2 is a degradation product or its concentration in OPSW has declined recently. In conclusion, this evaluation supports the utility of using the Family A and B combination as tracers of OSPW as they appear capable of application to historical conditions and generally behave conservatively when grouped, with some individual isomer differences noted during transport in this groundwater flow system. We note that this conclusion is based on the MLSB groundwater plume and results may differ at other ponds under other geological conditions.

Evaluation of Anthropogenic Chemicals. Artificial sweeteners and PFAS are two groups of anthropogenic contaminants known to be highly mobile and persistent in

groundwater with source apportionment capabilities.^{44–48} Despite the fact that they may not be unique to OSPW, their presence in OSPW may augment a weight of evidence approach. Artificial sweeteners were analyzed in 10 OSPW samples from three tailings pond sites, with concentrations of acesulfame and saccharin ranging from nondetect (<8 ng/L) to 530 ng/L and ~30 to 420 ng/L, respectively, significantly enriched over background groundwater (Figure S8).

The elevated amounts of sweeteners in many of these OSPW samples may result from tailings ponds historically receiving treated or untreated on-lease wastewater. They may also be derived via recycling of Athabasca River water containing these substances from upstream sources, during bitumen extraction. These compounds would typically be below detection limits for pristine waters. However, groundwater may be affected by other sources associated with oil sands developments, including wastewater treatment plants (WWTPs) and septic systems. For example, in a study of urban groundwater,⁴⁴ four sites (three with known wastewater influences) had maximum acesulfame concentrations >2500 ng/L and two sites (one wastewater, one landfill influenced) had maximum saccharin concentrations >2000 ng/L. Similarly, in a study of 12 septic system sites, Robertson et al.⁵⁷ found that acesulfame concentrations in groundwater plumes were typically >2000 ng/L. Finally, one background groundwater sample of Roy et al.,²⁰ collected along the Athabasca River in the oil sands area and near a WWTP had (unreported) acesulfame and saccharin concentrations of 290 and 50 ng/L, respectively. Thus, these other sources, which may occur near the tailings ponds in association with anthropogenic activity, combined with the low concentrations found in some tailings ponds may confound the identification of OSPW-affected groundwater using artificial sweeteners alone.

In this study, OSPW concentrations of PFBS, PFOS, and PFHxS ranged from nondetection (0.03 ng/L) to 60 ng/L, 0.5 to 75 ng/L, and 0.2 to 97 ng/L, respectively (Figure S9). The origin(s) of the PFAS in OSPW are not presently known. The predominant historical usage of PFOS is in aqueous film forming foams (AFFF) for fighting fuel-based fires.⁵⁸ Due to the industrial manufacturing process, PFOS-based AFFF also contains PFBS and PFHxS as minor impurities.⁵⁹ Given that the profile of PFAS congeners in OSPW did not resemble that of the Athabasca River and its tributaries, particularly considering the relatively low levels of perfluorooctanoate (PFOA) and perfluorononanoate (PFNA), recycling of Athabasca River water containing these substances from upstream sources during bitumen extraction can be eliminated as a source (Figure S10). For example, in OSPW samples, the concentrations of PFOA ranged from <LOD to 9.5 ng/L and those for PFNA ranged from <LOD to 0.20 ng/L, and similarly, the concentrations in the Athabasca River itself were 0.93 ± 0.33 ng/L PFOA, 0.097 ± 0.006 ng/L PFNA, <LOD PFHxS, 0.40 ± 0.05 ng/L PFOS, and 7.9 ng ± 2.1 ng/L PFBS. Conversely, the elevated presence of PFOS, PFBS, and PFHxS in OSPW relative to background groundwaters (Figure S9), and their relative similarities to AFFF, suggest the mining operation as a source. These findings are consistent with Dauchy et al.⁶⁰ that showed high levels of PFOS (9.5–19 μ g/L), PFBS (0.2–0.6 μ g/L), and PFHxS (1.3–2.7 μ g/L) in a wastewater lagoon sampled from an oil refinery that conducted on-site AFFF training. Similarly, Milley et al.⁶¹ indicated that commercial and military airports are sources of PFAS when AFFF is released during training, emergency responses,

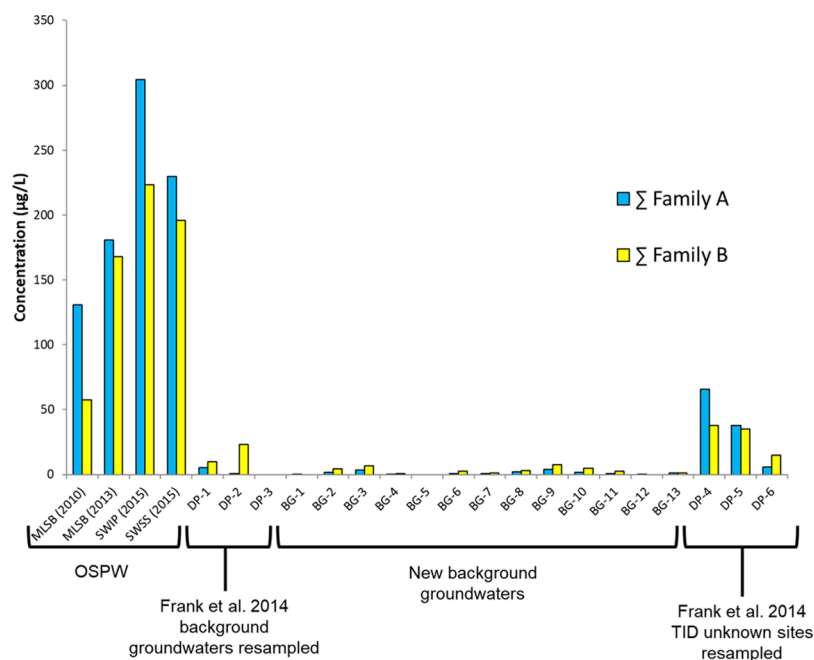


Figure 5. Total Family A and B concentrations in samples of OSPW, Background Groundwater, and Unknowns by Tar Island Dyke (TID, Table 1; Figure 1).

maintenance, and testing. Concentrations in groundwater affected by AFFF use may reach several orders of magnitude higher than those noted here for OSPW.^{45–47,62} For example, Anderson et al.⁶² reported mean groundwater concentrations corresponding to 309 $\mu\text{g/L}$ PFOS, 20 $\mu\text{g/L}$ PFHxS, and 7 $\mu\text{g/L}$ PFBS at U.S. air force operations. Similar to artificial sweeteners, PFAS should not be considered unique markers of OSPW. One pond with the lowest PFAS had just become operational when the pond was sampled.⁵¹

Generally, sweeteners and PFAS were found within the “in-plume” samples (Figure 4B). Saccharin and the PFAS generally tracked the plume well in relation to the metrics previously used to track OSPW migration at this site (e.g., NAs, sodium, chloride)¹³ and determined again here (Figure 4) with other Level-1 parameters (Figure 4; Figure S5). Plume concentrations of PFAS were elevated relative to the combined set of background groundwater in this study (Figure S9), which were comparable to groundwater samples from the United States: 1.26–11 ng/L PFBS, 1.88–45 ng/L PFHxS, and 3.25–3.41 ng/L PFOS.⁴⁸ Acesulfame was only present in well samples 3A and 4A, which are above the main OSPW-affected groundwater plume. Saccharin was also elevated in well 3A. These 2 wells are the shallowest within the well nests at locations 3 and 4, suggesting an alternate source of wastewater or landfill contamination at or near ground surface in that area. The saccharin found at well 5B (clearly in front of the plume) may have a similar source.

This study is the first to report elevated concentrations of artificial sweeteners and PFAS in OSPW. However, concentrations were not elevated in all of the tailings ponds and both have other sources that may exist as part of mining operations. Thus, their use as OSPW-indicators should serve as part of a weight of evidence approach with other parameters and must be applied with proper consideration of site conditions.

Evaluation of Unknown Samples Adjacent to Tar Island Dyke. Frank et al.¹⁶ collected shallow groundwater from 3 locations adjacent to TID along the shoreline of the

Athabasca River (DP-4, DP-5, DP-6); 2 of these (DP-4, DP-5) were determined to be likely affected by OSPW. These same 3 locations were resampled in 2013 (Unknowns, Table 1) and then reassessed for OSPW-indicators, utilizing the advancements made in this study. Total Family A and B concentrations for these locations are presented in Figure 5, along with those of 4 OSPW samples collected from three ponds from two mining operations and the 16 background groundwater samples. All unknown samples had quantifiable concentrations, but only DP-4 and DP-5 had values (especially for Family A) notably above the maximum of the background groundwaters. Both had relatively low concentrations of components AS, A7, and A8 (Table S6), consistent with samples at the front of the Mildred Lake plume (Figure S6). These two samples also had elevated SFS signatures and concentrations of F and NA, while those of DP-6 were low (Figure S4). Note that it is possible that DP-6 has some minor OSPW-influence but, based on the data generated here, not enough to clearly distinguish it from background. Unfortunately, there are no data on artificial sweeteners and PFAS from the source at TID (Pond 1, now reclaimed), further complicating interpretation of these data. Regardless, the most noteworthy finding for the anthropogenic tracers was that the concentrations of PFHxS and PFOS at DP-4 were greater than those of any OSPW sample (Figure S9). This may reflect an OSPW influence at DP-4, though it may also result from past PFAS uses around TID. The sweetener data for these 3 sites (Figure S8) was suggestive of an OSPW influence at DP4 and DP 6 but again may reflect other sources around TID.

A final assessment was conducted using a PCA with 31 chemistry parameters, including Family A and B (sums and individual components), O_2/O_4 , PFAS, artificial sweeteners, and a select group of inorganic compounds (typical screening parameters) and a Pearson Correlation matrix (Table S7). The biplot of this PCA (Figure 6) shows clear distinction along Component 1 (x -axis; strongly aligned with Family A and B vectors) between OSPW-related samples (OSPW of MLSB

(induction of hepatic ethoxyresorufin-O-deethylase activity) relative to sites upstream of mining activity.⁶⁴ At present, there is no linkage to any of these response patterns to OSPW seepage specifically; however, application of the methodology developed will now be able to confirm if seepage is entering surface waters.

This study has shown that the Family A and B monoaromatic acids with unknown chemical structures, and possibly acesulfame and perfluorinated surfactants, are components of the migratory OSPW mixtures beside TID. Other OSPW-derived organics, or in situ degradation products⁵⁶ with similar properties (e.g., water solubility), may also be present and pose undefined risks to aquatic biota. Determination of the drivers of toxicity within OSPW and impacted groundwaters, as well as the natural bitumen-affected groundwaters, is necessary to address the ecological relevance of OSPW-affected groundwater reaching surface waters. Given the complexities and inherent similarities in the chemical compositions of these sources, such studies need to be conducted carefully. Work in this area has been initiated,^{65,66} which will guide and inform the effects-based monitoring of fish populations and benthic communities of the Athabasca River watershed under the Canada-Alberta Oil Sands Monitoring Program.

■ ASSOCIATED CONTENT

SI Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.est.9b05040>.

Photos of field work, sample collection details, PFAS Analysis, additional methods, information on sample preservation (PDF)

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Notes

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