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BIOGEOCHEMISTRY IN AN INTERTIDAL POCKET BEACH

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Abstract

Sandy beaches are places of active organic matter mineralization due to water renewal providing organic matter and electron acceptors in the porous and permeable sands. Recycled biogenic compounds are efficiently transferred to the coastal marine environment via wave and tidal-driven advective flows. The biogeochemical processes in beach aquifers were mainly studied in semi enclosed systems with low tidal amplitude, and with a connection to continental aquifers contributing to solute fluxes to the coast from terrestrial groundwater. We present here the study of a pocket beach isolated from terrestrial aquifers with a high tidal amplitude and a medium energy wave regime. In situ measurements, cross-shore profiles and vertical sampling were conducted during several tidal cycles in spring and autumn. Cross-shore transects, obtained at low tide from holes that represent a mixture of the upper 20 cm of the water saturated zone, showed concentration gradients of redox and recycled compounds. Increase in pCO₂, dissolved phosphate and ammonium concentrations downslope revealed that more products from organic matter mineralization accumulated in the lower beach. The related increase in total alkalinity downslope indicated that the part of anaerobic processes in organic matter oxidation was higher in the lower beach. Concentration and $\delta^{13}\text{C}$ of dissolved inorganic carbon in pore waters suggested that the carbon mineralized in pore waters came from marine plant debris that were mixed with the sand. Continuous probe records of dissolved oxygen saturation and vertical profiles revealed a tidally-driven dynamics of pore water in the first centimetres of the lower beach aquifer. Ventilation of pore waters corresponded to wave pumping and swash-induced infiltration of seawater in the upper 10–20 cm of sediment. Nutrients and reduced compounds produced through organic matter mineralization remained stored in pore water below the layer disturbed by wave. The flux of these components to seawater is possible when this interface is eroded, for example when wave energy increases after a less energetic period. The low extension of the studied aquifer, typical of pocket

beaches, limits the connection with continental groundwater. Both tidally-driven and wave-driven recirculation of seawater allows pocket beaches to be efficient bioreactors for marine organic matter mineralization. As such, they provide the coastal environment with recycled nutrients, and not new nutrients.

Keywords: Submarine groundwater discharge; beach aquifer; nutrients; carbon cycle; Yeu island

1. Introduction

Almost one third of the world's ice-free shoreline is sandy (Luijendijk et al., 2018). Sandy beach sediments are permeable and constitute unconfined aquifers where pore water can circulate. Both terrestrial and marine forces drive underground water flows within beach sedimentary bodies. Resulting submarine groundwater discharge (SGD) can be a significant source of dissolved ions, nutrients, or contaminants to the coastal ocean (Sawyer et al., 2016; Slomp and Van Cappellen, 2004). Marine forces such as tidal pumping and wave-induced pressure gradients may induce fluid flow into superficial permeable sediments (Ataie-ashtiani et al., 2001; Burnett et al., 2003; Moore and Wilson, 2005). Along tidal coasts, seawater circulating through intertidal sandy beach sediments can contribute to 90% or more of SGD and its associated geochemical signature (Li et al., 1999). Hence, intertidal sandy beaches are the place of a large recirculation of seawater.

Sandy beaches generally have low sedimentary organic matter content (Boudreau et al., 2001). However, numerous studies have shown that they are very active places of remineralization due to water renewal in the porous and permeable media providing organic matter and electron acceptors, mostly oxygen from seawater (Anschutz et al., 2009; Billerbeck et al., 2006; Charbonnier et al., 2013; Rocha et al., 2009; Santoro, 2010). When remineralization products are discharged into the coastal ocean via recirculation of seawater, they eventually stimulate primary production and may even cause harmful algae blooms and eutrophication on the inner shelf (Paerl, 1997).

Terrestrial hydraulic gradients result in a net fresh water input into the ocean when terrestrial groundwater mixes with recirculating seawater. The transition between saline pore water and fresh groundwater is called subterranean estuary (Li et al., 1999; Moore, 1999). The sharp interface between water masses of distinct salinities may represent a redox front, which has the potential to reduce nutrient and contaminant loading via groundwater through removal processes such as denitrification and phosphate sorption onto mineral surfaces (Charette and Sholkovitz, 2002; Kroeger and Charette, 2008; Slomp and Van Cappellen, 2004; Spiteri et al., 2006).

Beach hydrology and biogeochemistry response to groundwater, tidal, and wave forcing have been extensively studied. Most of the field studies were conducted on relatively protected beaches characterized by micro- (< 2m range) or meso-tidal regimes (2 to 4 m range), such as Waquoit Bay in the western Cape Cod (Abarca et al., 2013; Gonnee and Charette, 2014), Cape Henlopen at the outlet of the Delaware Bay (Hays and Ullman, 2007; Ullman et al., 2003), Magdalen Island (Couturier et al., 2017), Long Island sound (Tamborski et al., 2017), Tolo Harbour, Hong Kong (Liu et al., 2018) and Ria Formosa (Rocha et al., 2009). The low-energy conditions occurring along these beaches allowed scientists to deploy fragile sampling devices such as wells and in-situ monitoring probes. In agreement with hydrological models tested with similar low hydrodynamic conditions (e.g., Boudafel, 2000; Robinson et al., 2007), most of the aforesaid studies showed that a beach aquifer was often characterized by a sharp transition zone between saline water and fresh water. This interface has a steep slope and is generally located close to the low tide level. Infiltration of saline water into the fresh groundwater in the tidal zone creates a reactive intertidal recirculation cell (Austin and Masselink, 2006; Turner and Masselink, 1998; Bratton, 2010) that has been referred to as the upper saline plume (e.g., Brovelli et al., 2007; Robinson et al., 2006; Santos et al., 2009; Vandenbohede and Lebbe, 2006). The magnitude of the tidal circulation cell and the associated transport of solutes within the beach aquifer increase with higher tidal range, steeper beach slope and hydraulic conductivity as well as lower in-land hydraulic gradient (Li et al., 2009; Robinson et al., 2007). The highly transient nature of the intertidal circulation cell over neap tide-spring tide and seasonal time scales has been evidenced on a micro-tidal beach system (Heiss and Michael, 2014). Studies were also carried out on high-energy beaches shaped by both a high mean tidal range and a high wave exposure, such as the Truc Vert beach on the French Atlantic coast (Anschutz et al., 2009) or the Spiekeroog Island off the North-West German coastline (Beck et al., 2017). In such hydrodynamic context, seasonal erosion and accretion processes drastically reshape the beach profile (Castelle et al., 2014). The unconfined aquifer of the intertidal zone generally consists of saline waters, the intertidal salt water plume being deeper than in low-energy beaches. In the upper meters of the intertidal aquifer, water has a short residence time due to the combined effect of efficient tidally driven recirculation of saline pore water and high permeability of sands (e.g. Charbonnier et al., 2013). Circulation of saline pore water may drag parcels of brackish waters from meteoritic groundwater mixture that may seep out the aquifer at the lower beach level (Buquet et al., 2016), contributing to the flux of nutrients to the ocean (Anschutz et al., 2016).

Small pocket beaches are common worldwide along rocky shorelines (Short and Masselink, 1999). Undergoing limited sediment supply, pocket beaches present generally thin thickness of sand deposited in the intertidal zone above the underlying bedrocks. Thus, the fresh water aquifers connected to pocket beaches are generally restricted, making the offshore fresh groundwater

discharge very weak. Consequently, pore water in tidal pocket beaches mainly consists of recirculating seawater. Here, the purely marine dimension of the biogeochemical reactor that a beach represents can be studied, with associated features and implications such as benthic-pelagic coupling and organic carbon respiration. To our knowledge, documentation on pocket beach pore water circulation coupled to biogeochemical processes is scarce, even missing in case of pocket beach with high tidal range.

In this paper, we report observations on a high tidal range pocket beach located on the Yeu Island off the French Atlantic coast to investigate short-term (tidal scale) dynamics of biogeochemical compounds in pore waters at two contrasted seasons (spring and autumn) by way of detailed cross-shore, and vertical characterization of biogeochemical compounds in the sandy beach aquifer. Two field campaigns were then conducted during periods of contrasted beach morphology, in October 2012 and April 2013. In October the beach displays the highest volume of sand stored in the intertidal area during summer good weather accretion, whereas in April the beach is highly eroded after winter storms. During each campaign, cross-shore profiles allowed to describe biogeochemical processes from the upper to the lower beach. Vertical profiles obtained in the sandy beach aquifer from home-made vertical sampling collectors enabled to collect pore waters at several depths and at different time after the tidal emersion. Finally, *in-situ* probes were buried in sediments to describe the dynamics of dissolved O₂ during several consecutive tidal cycles.

2. Material and methods

2.1. Study area

Located in the Bay of Biscay, Yeu Island is 8 km long and 3 km wide, elongated in a NW-SE direction about 20 km off French Atlantic coast (Fig. 1). The orthogneiss substrate of the island is gently sloping from erosional cliffs along the southern coast that reaches 32 m at its highest point, to nearshore bedrock outcropping at 0 to -2 m along the north-eastern coast (Durand et al., 2014). This NE coast consists of a succession of sandy pocket beaches partitioned by hard rock headlands. Sandy sediments form discontinuous lenses of less than 3 m thick over the very irregular bedrock. Aeolian dunes of low elevation (< 3 m) formed on the backshore are dominated by beachgrass (*Ammophila arenaria*) and couch grass (*Agropyrum junceum*), and backed by marine pine and oak forests or by holiday cottages. Beach sediment is mostly composed of coarse to medium-sized siliciclastic particles derived from the erosion of adjacent rocky outcrops, mixed with numerous shell debris, and thinly overlays the orthogneiss substrate.

The studied area, so called Ker Chalon beach, is a 750 m long embayed coastline that features typical pocket beach characteristics: curved in plan between two headlands, low sediment supply. Ker Chalon beach, facing the mainland, is protected from the SW to NW Atlantic swell and

thus is subject to a moderate wave climate issue from the wave refraction around Yeu Island. Tides are semi diurnal and high-mesotidal to low-macrotidal with a mean range of 3.5 m (from 1.4 to 5.5 m). This hydrodynamical context allows Ker Chalon beach to display a large flat dissipative profile in the lower beach, comprised of fine sands developing under the swash at low tide, and a relatively narrow steep reflexive profile in the upper beach, with interlayers of fine sands, coarse sands and gravels (functioning under the breaking waves at high tide).

2.2. Sampling collection

Seawater and pore water chemistry of the Ker Chalon beach was monitored during spring tide periods between the 16th and the 18th of October 2012, and between the 8th and the 11th of April 2013. The weather was rainy during both campaigns. For both campaigns, topography of the beach was measured cross-shore using an optical level, from the foot of the dune to the water line at low tide (Fig. 1). Profile elevations are given in reference to the Chart datum (mean lower low water).

For both campaigns, we applied three different techniques to get pore water properties of three successive days. To obtain a beach water table cross-shore transect, pore waters were sampled at the top of the water-saturated zone along parallel transects during the midday low tide (Fig. 1). In addition, pore waters were collected every 10 cm depth using 1-m long home-made vertical sampling collectors deployed at four sites at increasing distances from the dune foot. Autonomous O₂ probes were buried in the water-saturated sand for the whole duration of the field campaigns.

Pore waters from the top of the water-saturated zone were sampled according to Anschutz et al. (2009), and Charbonnier et al. (2013). Briefly, holes were dug every 10 m along a cross-shore transect from the low-tide swash zone to the high-tide watermark. Seawater samples were collected at the beginning of each transect. The water-saturated zone of sediment was reached with one shovelful in the lower beach because sand was water-saturated up to the surface, whereas we had to dig holes more than 0.5 m deep in the upper beach to reach the saturated zone. The first water that filled the bottom of the hole was removed with a polypropylene beaker. Due to the permeability of sand, holes refilled slowly and the measurements were performed on the replenish water. Temperature, salinity, pH, and dissolved oxygen saturation were directly recorded in the waters of the excavated holes within one minute using WTW probes. The probes were calibrated before and after each field campaign using an aerated solution (100% saturation) for the oxygen saturation, an IAPSO seawater and deionized water for the salinity, and with NBS standard solutions (pH = 4.01 and pH = 7.00) for the pH. Oxygen and salinity probes were compensated automatically for *in-situ* temperature. Waters were sampled using a 50-mL syringe and filtered through a 0.45 µm cellulose acetate syringe-membrane. One subsample was acidified with a 1% equivalent volume of concentrated HCl for later analyses of dissolved inorganic phosphorus (DIP), dissolved iron and

manganese; another subsample was kept frozen until later analyses of other dissolved nutrients (ammonium, nitrite, nitrate and dissolved silica (DSi)). We stored samples for the total alkalinity (TA) in polypropylene bottles after filtration using a syringe equipped with glass fiber filter (0.7 μm). The stable isotope signature of the dissolved inorganic carbon ($\delta^{13}\text{C-DIC}$) was measured in waters of the first campaign. The $\delta^{13}\text{C-DIC}$ samples were collected using 120 mL glass serum bottles sealed with a rubber stopper and poisoned with 0.3 mL of HgCl_2 at 20 g L^{-1} to avoid any microbial respiration during storage. Vials were carefully sealed taking care that no air remained in contact with samples. Vials were also stored in the dark to prevent photo-oxidation. Several samples of the solid fraction from the holes of the cross-shore transect were sampled in October 2012 in order to measure particulate organic carbon (POC) and inorganic carbon (PIC), inorganic phosphorus and reactive particulate iron and manganese concentrations.

Vertical profiles of pore water composition were obtained along a cross-shore transect at two stations in October 2012 (stations 1 and 3) and at four stations in April 2013 (Fig. 1). Each station was equipped with a pore water collector. The device was made with 1.2 m long and 10 cm diameter PVC tube pierced every 10 cm on the height. Tygon® tubes (6 mm ID) were installed inside the PVC tubes to connect each orifice to the top of the collector. The maximum dead volume was 30 mL for the longest tubes. A 100 μm mesh was put to the end of each tube in contact with sand to prevent blocking during the sampling. Pore water collectors were positioned 5 m away from the cross-shore transects described above. The upper station (station 4) was located 22 m off the dune foot in April 2013. The other stations were aligned cross-shore in the seepage face, 30 m or 20 m away one from another (Fig. 1B). Station 3 was located in an area highly colonized by sandworms (*Arenicola marina*). 20 mL pore waters were collected with syringes through each tube after discarding the corresponding dead volume to rinse the sampling device. The sampling was always performed from the topmost surface level to deepest levels to avoid vertical mixing of pore waters. Collectors were installed the day before starting the sampling. Collection of pore waters started on the falling tide, just after the emersion of each station, and was repeated every 2-3 h until immersion of each station during flood tide. Oxygen saturation and temperature were measured immediately after pore water collection using a FireSting O_2 meter equipped with microOptode and temperature probe (*PyroScience GmbH*). Since these probes were available only during the second campaign (April 2013), oxygen saturation was not measured in samples of vertical pore water collector during the first campaign (Oct. 2012). The microOptode was calibrated daily for the oxygen saturation using an aerated seawater solution (100% saturation). Oxygen microOptode was compensated automatically for water temperature. Water samples were filtered through a 0.2 μm cellulose acetate syringe-membrane. One subsample was acidified with 10 μL of concentrated HNO_3 per 10 mL for later analyses of DIP, dissolved iron and

manganese, salinity and sulphate; another subsample was kept frozen until later analyses of dissolved inorganic nitrogen compounds (ammonium, nitrite and nitrate).

Autonomous probes equipped with a data logger were buried directly into the sediment during 2 or 3 days at each campaign according to Charbonnier et al. (2016). Four stations were examined simultaneously along the cross-shore profile (Fig. 1B). In October 2012, station 1 and 2 were located in the lower beach at 102 and 72 m from the dune foot, respectively. Station 3 located 42 m from the dune was in a beach portion highly colonized by *Arenicola marina*. Station 4 was located on the upper beach. In April 2013, the stations were located at the same distance from the dune foot than the pore water collectors. The depth at which the probes were buried was in top 20 cm of the water table at low tide, i.e., at 20 ± 5 cm depth in the lower beach and at 60 ± 5 cm depth in the upper beach. They measured continuously water pressure that is converted in water head (CeraDiver sensors, Schlumberger®, precision of ± 0.2 cm), salinity (STPS 100SI loggers, NKE instrumentation; precision of ± 0.1), water temperature and dissolved oxygen concentration (SDOT300 logger, NKE instrumentation, equipped with an Aanderaa optode model 3835; precision of ± 0.05 °C and ± 5 % respectively, without oxygen consumption or significant drift). Each probe recorded pore water parameters *in-situ* every 2 min. Note that optodes were placed in water during 24 h before burial to moisten their foil as recommended by Aanderaa.

2.3. Analyses

For pore waters sampled at the top of the water-saturated zone, the dissolved inorganic compounds were analysed colourimetrically according to standardized techniques. NO_2^- and NO_3^- were analysed according to Schnetger and Lehnert (2014). Precision was $\pm 5\%$ for NO_2^- and $\pm 10\%$ for NO_3^- . DSi and DIP were measured by colorimetric procedures (Mullin and Riley, 1955; Murphy and Riley, 1962). Ammonium and DIC (only for Oct. 2012 samples) were analysed with the FIA method described by Hall and Aller (1992). The frozen subsample was defrosted on the day of the nitrite, nitrate and ammonia analyses. The subsample was allowed to stand at room temperature for at least 24 h for depolymerization of silica before carrying out DSi analysis. The obtained precision of these procedures was $\pm 5\%$. DIP was analysed on the acidified sample to prevent dissolved Fe(II) oxidative precipitation and subsequent post-sampling phosphate sequestration on Fe(III) phases. Dissolved iron (Fe^{2+}) was analysed by colourimetry by adding a ferrozine solution in an aliquot (Stookey, 1970). Total dissolved manganese was determined colourimetrically using the Cd-TCPP complex according to Madison et al. (2011) and Charbonnier and Anschutz (2019). Dissolved Fe and Mn were determined with $\pm 5\%$ precision.

For samples collected with vertical pore water collectors, nitrate, nitrite, and ammonium were measured using standard methods on a QuAatro AutoAnalyzer (*Seal Analytical*). DIP was measured on the acidified sample by colourimetric method of Murphy and Riley (1962). Measurements of dissolved Fe, Mn, S and Na were performed by an ICP-OES ICAP 6300 ThermoFischer after a 50-fold dilution with a 1% ultrapure nitric acid. Analysed elemental sulphur was interpreted as sulphate since sulphide is volatile and was removed from the sample solution under acidic conditions. Salinity was calculated from sodium concentration.

TA of surface water table samples were analysed on filtered samples by automated electro-titration on 50 mL filtered samples with 0.1N HCl as titrant. The equivalence point was determined from pH between 4 and 3 with the Gran method (Gran, 1952). Precision based on replicate analyses was better than $\pm 5 \mu\text{mol L}^{-1}$. We calculated the partial pressure of CO_2 ($p\text{CO}_2$) from TA, pH, and water temperature measurements using the carbonic acid dissociation constants of Millero (1979) and the CO_2 solubility from Weiss (1974) from the CO2SYS software (Lewis and Wallace, 1998). The $\delta^{13}\text{C-DIC}$ measurements were performed with the headspace technique using a isotope ratio mass spectrometer coupled to an elemental analyser (EA-IRMS, Micromass IsoPrime) as described in Gillikin and Bouillon (2007).

The grain-size distribution was measured using a Malvern laser diffraction particle size analyser. The freeze-dried solid fraction was homogenised for chemical analyses. POC was measured by infrared spectroscopy (LECO 200 C-S analyser) after removal of carbonates with 2M HCl from 50 mg powdered sample (Etcheber et al., 1999). Total particulate carbon was measured with the same technique, on aliquots that were not acid-leached. PIC converted into CaCO_3 was deduced from the difference between total carbon and POC. Detection limit was 0.2%. The precision of measurements was better than 5%. One gram crushed bulk sediment was leached with a 1N HCl solution to extract reactive P, Fe, and Mn adsorbed on particles and precipitated as oxide, carbonate, and phosphate minerals (Anschutz et al., 2005; Kostka and Luther, 1994). Extracted iron, manganese, and phosphorus contents were analysed colourimetrically according to standardized techniques with $\pm 5\%$ precision (Anschutz and Deborde, 2016; Charbonnier and Anschutz, 2019).

3. Results

3.1 Beach morphology and sediment characteristics

Ker Chalon beach was 140 m wide at low tide in October 2012 and 130 m in April 2013 (Fig. 1). The difference was due to the spring tide amplitude, which was higher in October (4.8 m) than in April (4.4 m). The beach profile changed between autumn 2012 and spring 2013. In autumn, the beach showed a slightly convex to almost linear profile with a relatively steep gradient (7%) normally

resulting from summer aggradation. At 20 m from the dune foot, an erosional step of about 0.5 m high was dug at the upper limit of wave run-up at high tide. This kind of feature in the upper foreshore indicated the beginning of the winter erosion processes. The upper part of the beach was eroded in April relative to the measurements carried out 6 months earlier. In spring, the beach showed a concave profile with a slope break at 30 m from the dune foot, dividing the beach into two distinct parts. The upper foreshore was 2 m below the topographic profile measured in October with a slope of 17%, whereas the very flat lower foreshore displayed a dissipative morphodynamic beach state with a slope of 2%. A source of seeping waters appeared at the slope break at low tide. The lower beach remained always wet during tidal emersion. A continuous longshore zone of 5 to 10 m width located at the upper part of the seepage face was highly inhabited by *Arenicola marina* both in October and April (Fig.1), as evidenced by the presence of 2 to 10 fecal mounds per m². The bedrock, located at several decimetres below the topographic surface was not reached with the sampling devices.

The particle size showed a large degree of heterogeneity between the upper and the lower beach. The upper beach was composed of very permeable coarse sands and gravels, whereas the surface sediment of the seepage zone, which constituted the main part of the studied beach, consisted of well sorted particles composed of fine sand with a median grain size of $166 \pm 10 \mu\text{m}$ (Table 1) and a POC concentration of $0.20 \pm 0.08\%$ ($166 \pm 70 \mu\text{mol g}^{-1}$), and a CaCO_3 content of $18.2 \pm 4.6\%$ ($15167 \pm 3833 \mu\text{mol/g}$). Reactive inorganic Fe, Mn, and P concentrations were $48.6 \pm 3.5 \mu\text{mol g}^{-1}$, $1.21 \pm 0.20 \mu\text{mol g}^{-1}$, $12.1 \pm 2.8 \mu\text{mol g}^{-1}$, respectively (Table 1). Sediment colour was light olive grey (5Y 7/2) over the first 10 cm and grey (5Y 5/1) below. Hydraulic conductivity was not measured in sands of the Ker Chalon beach but several empirical formulae allow to estimate this parameter based on grain size distribution (Vienken and Dietrich, 2011). The USBR formula (Vuković and Soro, 1992) gave 2.4 m d^{-1} whereas the Seelheim (1880) formula gave 8 m d^{-1} . Ker Chalon beach is thus an original example of an aquifer with a high tidal range, medium energy, gentle slope and medium permeability.

3.2. Water head measurements

As expected, water head levels increased during flood and decreased during ebb (Fig. 2). In the seepage zone of the lower beach (stations 1 to 3), water level was almost constant at low tide during emersion of the stations (Fig. 2), because the whole sediment column remained saturated with water. For these three stations, the water head was the sediment surface. In the upper beach (station 4), the water head slowly decreased as sediment emerged.. The drop in water level occurred in the porous medium due to the exfiltration of water downslope. It decreased continuously until the

flooding of the next tide. The upper beach sediments were unsaturated in water down to 80 cm at low tide (Fig. 2).

3.3. Temperature and salinity

Seawater and pore water temperature was relatively constant, around 16 °C in October 2012 and between 9 and 11 °C in April 2013. Pore water salinity remained close to 34, which was similar to the value measured in seawater (Fig. 3). Some surface pore waters showed lower values around 30. Vertical profiles from collectors indicated that salinity remained around 33-34 in the middle part of the beach, at stations 2 and 3 (Fig. 4). Nevertheless, brackish waters were detected in April 2013 in the upper beach below 1 m depth (station 4 with salinity around 30) and below 40 cm depth in the lower beach (station 1 with salinity down to 23). Profiles were steady during time series of sampling with pore water collectors (Fig. 4). Continuous measurements with probes located in the upper part of the water-saturated zone also showed constant (± 1) salinities, except at station 4 in April 2013, where slightly lower salinities were recorded after emersion period (not shown on Fig. 2).

3.4. Dissolved oxygen records

Waters collected along cross-shore profiles in the superficial part of the water table showed a decrease in oxygen concentration from the upper beach to the lower beach (Fig. 3). A similar trend was observed for both investigated seasons. In the upper beach section near the dune foot, oxygen saturation of sediment pore water varied between 80% and 100% whereas most of the values were between 20% and 40% in the lower beach section (Fig. 3). These values from holes dug in the sand represented pore water samples that integrated the upper decimetre of the water table. In April 2013, the vertical profiles determined from pore water collectors showed that oxygen concentration decreased with depth and reached a value lower than 20% below 40 cm depth at stations 1 and 2 (Fig. 4). This clearly showed that the sediment was divided into two parts. An upper part was enriched in dissolved O₂, and a lower part, corresponding to the grey sand, was depleted in dissolved O₂ and almost anoxic. Time series during emersion showed that the oxic layer displayed redox oscillations, with values close to 100% saturation just after emersion, and a rapid decrease at a rate of about 10–15% saturation per hour (Fig. 4). Vertical profiles were more scattered at station 3 inhabited by the sandworms *Arenicola marina*. They also showed a decrease in O₂ saturation during emersion, but waters remained oxic down to 1 m depth (Fig. 4). At station 4, the water table deepened during emersion and pore waters remained oxic. *In-situ* probes recorded oxygen changes during whole tide cycles (Fig. 2). They were placed in the transition zone between oxic and anoxic sediments. Probes were mainly in the anoxic part of the sediment at station 1 and at station 2 in October 2012 (Fig. 2). In April 2013 at station 2, the probe buried at 20 cm depth recorded redox

oscillations, which showed an oxygen saturation decrease after emersion reaching values below 5%, in agreement with pore water collector sample data. The probe also showed that oxygen level increased abruptly about 1 or 2 hours after the start of sediment immersion in the swash zone. The maximum value remained below 25%, and this value was reached approximately 2 to 3 hours after high tide (Fig. 2): redox oscillations followed the tidal cycle with a delay of a few hours at depth. In addition, we observed a second peak of dissolved O_2 that occurred at low tide slack water on the decreasing shoulder of the main peak. Such double-peaks were also observed at stations 3 and 4 (Fig. 2). At these stations, oxygen saturation also changed with tide and presented oscillation of 10% amplitude.

3.5. Nutrients

The cross-shore profiles of dissolved nutrients (NO_2^- , NO_3^- , NH_4^+ , DIP, DSi) of October 2012 and April 2013 had similar shapes and showed relatively low temporal variability (Fig. 3). Pore waters at the top of the water table showed opposite cross-shore profiles for nitrate and ammonium (Fig. 3). Pore water nitrate concentrations in the upper beach and in the zone inhabited by *Arenicola marina* were higher than in seawater. Most of the values were between 20 and 40 μM (Fig. 3). NO_3^- remained below 10 μM in the lower beach. Vertical pore water profiles of NO_3^- showed values similar to those measured in the holes of the cross-shore profiles (Fig 3 and 4). Time series of depth profiles during emersion showed that nitrate concentration in deep pore waters did not change. However, we did observe some changes in NO_3^- concentration in the top 20 cm of the sediment (Fig. 4). Immediately after the station was exposed, the surface pore waters had a nitrate concentration close to that of seawater. Afterwards, the concentrations returned to the values found deeper, so that vertical profiles became homogeneous a few hours after emersion until the next immersion (Fig. 4). Nitrite concentrations were often close to the detection limit, except at the bioturbated station 3, where concentrations up to 5 μM were measured in surface pore waters (Fig. 3 and 4). Waters of surface table along the cross-shore transects were characterized by increasing gradients of ammonium, DIP, and DSi, from the upper beach to the low tide swash zone (Fig. 3). Ammonium reached 100 μM in the lowest part of the beach, downstream from station 1 (Fig. 3). Vertical pore water profiles of NH_4^+ showed mean concentrations similar to those measured in holes (Fig. 3 and 4). As with nitrate, NH_4^+ profile time-series between emersion and flood showed changes in the surface sediment with concentrations going from the value of seawater to the value measured deeper after a few hours (Fig. 4). DIP concentrations were around 10 μM in holes of the lower part of the beach, whereas they were close to seawater concentration in upper beach pore waters (Fig. 3). Here again, depth values from vertical profiles were similar to those from the holes, with concentrations close to 2 μM at the investigated stations (Fig 3 and 4). Dissolved Si cross-shore profiles had the same shape

as DIP profiles (Fig. 3). DSi concentration was below 10 μM in seawater and in holes of the upper beach, whereas a concentration of 50 μM was reached in holes of the lower beach.

3.6. Other redox species

In the cross-shore profiles, concentrations of dissolved Fe and Mn higher than seawater values were found in pore waters depleted in dissolved oxygen, in the lower part of the beach (Fig. 3). Presence of dissolved Fe above 10 μM occurred more particularly in samples depleted in dissolved nitrate (Fig. 3). Hence, dissolved Fe concentration was close to seawater values in the bioturbated zone of the beach and in the upper beach, and Fe and Mn concentrations reached up to 75 μM and 8 μM , respectively, in holes of the lower beach. Vertical pore water profiles of the stations 1 and 2 showed that dissolved Mn and Fe were absent in the top 20 cm sediments, and up to 3 and 6 μM below (Fig. 4). Sulphate was measured in samples from pore water collectors. Sulphate concentrations were close to the value expected from the salinity measurement, with values between 27 and 28 mM at salinity close to 33, but some low sulphate deficit relative to salinity occurred in pore waters of stations 1, 2, and 3 in April 2013 samples (not shown).

3.7. Carbon

Parameters linked to carbonate chemistry were only measured in waters collected in holes of cross-shore transects because the required water volume for analyses was too high to do it on samples collected with pore water collectors. These parameters were similar for the three replicate cross-shore transects realized during each campaign (Fig. 5). They also showed comparable trends in October 2012 and in April 2013. TA of seawater was 2.3 mM (Fig. 5). Values measured in cross-shore transects were close to seawater value in the permeable upper beach. TA increased up to 4 mM in the seaward direction. The pH of seawater was close to 8.0 (Fig. 5). Minimum values were 0.2 pH unit below seawater value in pore waters of the lower beach, except near the low tide mark in the profile of April 11th 2013, in which the minimum value was 7.4 (Fig. 5). pCO_2 calculated from pH and TA was in equilibrium with the atmosphere in seawater samples (close to 400 ppm) (Fig. 5). Values were higher in pore waters with a pCO_2 value varying roughly between 800 and 2000 ppm. Interestingly the lower beach pore water of April 11th reached a pCO_2 of 5000 ppm (Fig. 5). $\delta^{13}\text{C}$ -DIC was only measured in samples of October 2012 (Fig. 5). Seawater $\delta^{13}\text{C}$ -DIC value was 0.15 ‰. It was lighter (more negative) in pore waters than in seawater. These pore waters with low $\delta^{13}\text{C}$ -DIC values were associated with high TA, with a minimum value of -7.92 ‰ measured in the sample with the highest TA value (Fig. 5).

4. Discussion

4.1. Physical context of the beach

The Ker Chalon beach aquifer was mainly composed of saline water (Fig. 3), and did not have an underlying terrestrial groundwater body, as in most of the beaches studied. The main reason is that the Ker Chalon beach aquifer has a limited vertical extent. It is a sand lens (< 3 m thick) lying on the crustal bedrock, which prevents the intrusion of terrestrial groundwater in the tidal zone. The beach aquifer is mostly recharged by tidally driven recirculation of seawater. In addition to tidal forcing, waves increase the penetration of saline waters in the beach aquifer (Robinson et al., 2014). Some brackish waters were evidenced in the lower beach, more particularly in April 2013 (Fig. 4), suggesting that some fresh water may be occasionally carried with the recirculating salt water. The freshwater mixed with recirculating seawater could be provided by meteoric waters infiltrating the low backshore dune bordering the beach.

4.2. Vertical and horizontal gradients of biogeochemical compounds

There was a significant difference in temperature between October 2012 (15-16°C) and April 2013 (9-12°C). However, the profiles shown in Figures 3, 4 and 5 did not indicate sufficient differences to interpret the results in terms of seasonal changes. The shape of vertical profiles looked like well-known diagenetic profiles of marine sediments, where oxidized dissolved compounds were present in pore waters close to the sediment surface (Fig. 4). Their concentration decreased with depth, and reduced and recycled compounds accumulated at depth (Fig. 4). This was particularly true at stations 1 and 2 located in the lower part of the beach. In these stations, ammonium, dissolved Fe, and Mn concentrations increased below 20 cm depth, when dissolved O₂ and nitrate dropped to low values (Fig. 4). These results suggest that sediment organic matter was oxidized through reactions in which the available terminal electron acceptor that yielded the maximum energy was used, as encountered in diffusion-dominated marine sediment (Froelich et al., 1979). At stations 4 and 3, located in the upper part of the beach, sediment remained oxic, and reduced compounds such as dissolved Fe and Mn kept concentrations close to those of seawater, below 2 µM, down to a depth of 1 m (Fig. 4). Nitrate concentration in these stations was higher than the value of seawater. As the pore water salinity was similar to that of seawater, higher nitrate concentrations were most likely the result of oxic mineralization of sedimentary organic matter (Charbonnier et al., 2013).

Cross-shore transects also showed horizontal concentration gradients of redox and recycled compounds across the beach (Fig. 3). Waters of the upper beach were well oxygenated. In the lower beach, the presence of reduced compounds and high concentrations of recycled nutrients indicated that organic matter was mostly mineralized through anaerobic processes such as denitrification, reduction of Mn and Fe oxides, and sulphate reduction. N:P ratio deduced from pore water NH₄⁺:DIP ratio varied between 10 and 15 in holes of the lower part of the beach and in the vertical profiles of

station 1, suggesting that the mineralized organic matter was of marine origin. Increase in $p\text{CO}_2$ (as well as DIP and NH_4^+) downslope the cross-shore transect (Fig. 5) clearly indicated that more products from organic matter mineralization accumulated in the lower part of the beach. The related increase in TA downslope suggests that the part of anaerobic processes in organic matter oxidation was higher in the lower beach, as anaerobic processes create alkalinity (Kempe, 1990; Soetaert et al., 2007; Thomas et al., 2009). Denitrification and ammonification, when ammonium is not subsequently nitrified, produce alkalinity (Abril and Frankignoulle, 2001). Both processes generate one mole of TA per mole of nitrogen involved in the reaction. Sulphate reduction was suggested by a sulphate deficit in the pore water of some samples and the grey colouration of sediments below a few cm in depth, indicative of the presence of authentic iron sulphides. This process produces two moles of TA per mole of sulphate reduced to S(-II) species, either escaping to the atmosphere when sediments are exposed at low tide (Kristensen et al., 2000), or buried as Fe-sulphide. Fe(III) and Mn(III,IV) oxide reduction is a net source of alkalinity if reduced Fe(II) and Mn(II) do not precipitate again in the sediment. Carbonate mineral dissolution is also a source of alkalinity. However, this process is not in agreement with the observed increase in $p\text{CO}_2$ and the decrease in $\delta^{13}\text{C-DIC}$ (Fig. 5). DIC added in pore water shifted the isotopic signature to negative values, suggesting that this additional DIC came from a lighter source of carbon. Marine carbonates have a relatively heavy signature close to 0‰, whereas organic matter has a negative signature. In addition, a graph showing the evolution of alkalinity as a function of DIC shows a slope of 0.9 (Fig. 6A). This slope reflects the stoichiometry of the predominant reactions controlling dissolved carbonate chemistry and thus can be used to infer which reactions control dissolved carbonate chemistry (Cai et al. 2003; Liu et al. 2017). This slope would be 2 if the increase in alkalinity was due to the dissolution of carbonates. A slope close to 1 is compatible with denitrification and sulphate reduction processes. All of these results strongly suggest that the carbon added to the interstitial waters came from anaerobic mineralization of sedimentary organic matter. This result means that the studied beach is a source of alkalinity, as are larger systems such as the Wadden Sea intertidal zones (Beck et al., 2008, Santos et al. 2015). Therefore pocket beaches may account for the alkalinity budget of coastal ocean, and as a net total alkalinity generation setting, they have the potential to increase the capacity of the coastal ocean buffer and the uptake of atmospheric carbon dioxide (Hu and Cai, 2011; Brenner et al., 2016; Voynova et al., 2018).

Respiration processes in the beach aquifer occur when labile organic matter is present. A Keeling-like diagram, where $\delta^{13}\text{C-DIC}$ is plotted vs. $1/\text{DIC}$ concentration (Pataki et al., 2003), was used to determine the isotopic composition of the organic carbon source respired in beach sediments. Concentration and $\delta^{13}\text{C-DIC}$ in pore waters could be explained by a mixture of seawater DIC with a lighter source of additional DIC with a mean isotopic signature of $-12.2\text{‰} \pm 0.6\text{‰}$ (Fig. 6B). This value

is less negative than marine phytoplankton isotopic signature ($-22.7\text{‰} \pm 0.8$; Dubois et al., 2012) and is close to the isotopic value of marine and saltmarsh angiosperms ($-11.9\text{‰} \pm 1.4$; Dubois et al., 2012), as well as macroalgae such as seaweeds ($-18.8\text{‰} \pm 2.6$; Dubois et al., 2012) from neighbouring rocky shores and *Ulva sp.* ($-11.1\text{‰} \pm 1.8$; Riera et al., 1996), suggesting that the carbon mineralized in pore waters came from the sea or the seashore, and not from land sources, such as terrestrial higher plants and river SPOM ($-28.0\text{‰} \pm 1.3$; Dubois et al., 2012). It may have resulted from marine plant debris left on the beach at high tide (also known as wracks) mixed with the sand during seasonal erosion-accretion episodes or sand remobilization due to waves. Wracks, of which small piles have been observed on the beach, may have represented here a source of labile organic matter as observed in other beach aquifers (Dugan et al., 2011; Lastra et al., 2018).

4.3. Pore water residence time

The role of water residence time on biogeochemical processes was pointed out in several studies focussing on seawater recirculation (e.g., Billerbeck et al., 2006; Waska et al., 2019). The studied transects covered oxic to anoxic redox-conditions as it has been described for other beaches (e.g. Reckhardt et al., 2015). The existence of the well-developed horizontal redox zonation from oxic to anoxic conditions towards the sea in the beach aquifer implied either that organic matter fuelling biogeochemical reactions was not the same, or that physical controls differed between the upper and the lower beach. We have measured similar concentrations of organic matter along the transect and we did not observe a gradient of $\delta^{13}\text{C}$ -DIC end-member, suggesting that the origin of organic matter was not the variable that explained the horizontal redox gradient. Field observation suggests that water residence time was the main parameter that explained differences in biogeochemical processes. The water table of the upper beach dropped at each low tide (Fig. 2), indicating that the porous environment of the upper aquifer transiently filled with air was replenished with seawater at each tide. This zone was infiltrated by well-oxygenated, low-nutrient seawater at each flood. During ebb tide, a part of the pore water was flushed towards the sea, leaving the sand water unsaturated until depth around 80 cm at station 4 (Fig. 4). Deep layers of the upper beach aquifer that remained water-saturated, sampled at low tide in holes and at the bottom of pore water collectors, had a residence time sufficiently long to become enriched in dissolved nitrate, slightly enriched in DIP and DSi, but not long enough to become anoxic.

The upper beach acted as the recharge zone of the beach aquifer. Seawater infiltrated the beach face during flood faster than it discharged during ebb. Hence, the time-averaged beach water table was higher than the mean sea level (Turner et al., 1997), creating a seaward hydraulic gradient at low tide. Pore water circulated into the beach aquifer and seeped out in the lower intertidal zone (Horn, 2006; Robinson et al., 2007; Xin et al., 2015). In the Ker Chalon beach, the seepage face

represented almost all the beach, from 40 m off the dune foot to the low tide mark located 100 m seaward farther. The fact that the water table height was almost equal to the lower beach surface is characteristic of fine sand beaches (Horn, 2006). The upper part of the seepage face, at 40 m from the dune foot corresponded to the portion that was highly inhabited by *Arenicola marina*. It was oxic down to 1 m depth (Fig. 4), suggesting that this section of the beach received pore waters from the recharge zone as a result of the seaward hydraulic gradient that occurred at low tide. In this portion of the beach, oxygen concentrations showed a reverse profile after 5 hours of exposure with concentrations at depth between 40 and 90 cm that were higher than those between 10 and 40 cm (Fig. 4). The ventilation of the deep layer was not due to *Arenicola marina* bioirrigation because worm activity does not reach the sediment deeper than 40 cm depth (Meysman et al., 2005). Deep ventilated waters most likely came from the recharge zone and flowed towards the lower beach aquifer during low tide.

Oxic waters were only present in the first decimetres of the lower beach aquifer (Fig. 4). Below 30 cm depth, concentrations of reduced compounds (dissolved Fe and Mn) and metabolic products (ammonium, DIP, DSI) increased seaward. This suggests that waters flowing seawards in the beach aquifer became depleted in dissolved oxygen and nitrate because these waters were old. Oxygen was consumed and denitrification occurred before waters were flushed through tidally-driven recirculation. Considering the value of hydraulic conductivity estimated from grain size of fine sand (between 2.4 and 8 m d⁻¹), and the slope of the beach, the Darcy velocity was between 1 and 3 cm h⁻¹. This meant that pore water covered only a few centimetres seawards during each low tide. This is in agreement with the fact that profiles from pore water collectors sampled three consecutive days at each campaign remained relatively constant below 30 cm depth. The estimated Darcy velocity implied that anoxic pore waters found in the lower beach aquifer originated from seawater that entered the upper beach sand several months before sampling. Such a low renewal of pore water in the lower beach explains why waters became anoxic and enriched in metabolic products. It also suggests that rain waters that infiltrated the sand in the upper beach and in the dune aquifer were not totally diluted in saline pore water because of its low renewal. This may explain why we found brackish pore waters at some places of the lower beach. The fresh water end-member probably originates from rain events that occurred several months before seeping.

4.4. Tidal-scale dynamics

Continuous probe records of dissolved oxygen saturation and vertical profiles of some chemical components revealed a tidal dynamics of pore water in the first centimetres of the lower beach aquifer. Results showed that dissolved oxygen concentration increased abruptly when flood tide reached the studied stations (Fig. 2). Thus, sudden ventilation of pore waters corresponded

probably to wave swash-induced infiltration of seawater. Several studies have determined the dynamics of the swash zone moisture. Studies focusing on the estimates of swash infiltration into unsaturated sand (e.g. Heiss et al., 2014; Masselink and Turner, 2012; Steenhauer et al., 2011) showed that a water table mound thus formed was a consequence of a rapid rise in water content during rising tide. Seawater inflow was the highest near high tide as observed at station 4 located close to the high-tide water mark (Fig. 2). In most of the beach aquifer, from station 3 to the lower part of the beach, the sediment remained water-saturated up to the sediment surface at low tide. In this situation, the beach could behave as an impermeable surface (Packwood, 1983). Nevertheless, our observations suggest that swash-induced dynamics consisted of a replacement of pore waters by seawater in the upper 10–20 cm of sediment. It has been shown that waves caused cyclic infiltration-exfiltration corresponding to vertical flow across the sediment surface in saturated sediments (Turner and Masselink, 1998). More generally, pressure gradients due to wave generate fluxes across the sediment-water interface as observed in many different contexts (Huettel et al., 2014; Riedl et al., 1972; Santos et al., 2012; Shum and Sundby, 1996). Therefore, ventilation of the upper 10–20 cm sediment surface was the result of pore water-seawater mixing in the swash zone during flood and ebb tides, and also during the period of immersion, due to wave-induced pressure gradients (wave pumping). The increase in the intensity of this pump resulted in better irrigation and ventilation of the submerged sands and a deepening of the redox front (McLachlan and Turner, 1994; Riedl et al., 1972). Probes deployed at stations 1 and 2 showed very low concentrations of dissolved oxygen (Fig. 2), probably because they were placed at a depth at the limit of the influence of wave-induced water mixing. However, the periodic ventilation due to swash and immersion at each high tide could be seen clearly (Fig. 2).

The evolution of concentration profiles during emersion was observed from the hourly sampling of pore waters (Fig. 4). Pore waters collected at the beginning of beach exposure period were similar to seawater in the upper 10–20 cm. In this layer, oxygen concentration decreased rapidly except at the very surface of the sediment, which remained in contact with atmosphere. This observation has been made for the three periods of low tide monitored the 9th, 10th and 11th April 2013, confirming that ventilation of pore waters occurred at each high tide. The rapid shift in oxygen concentration at low tide corresponded to a loss of more than 100 μM in less than 5 hours at 10 cm depth at stations 1, 2, and 3. This change could not alone be explained by *in-situ* aerobic respiration process confined in the upper part of the aquifer, because such an efficient respiration rate would also have occurred at high tide, which was not the case. Oxygen loss due to aerobic respiration would have been consistent with an increase in nitrate concentration, which was also not the case at stations 1 and 2 (Fig. 4). In fact, oxygen and nitrate profile monitoring showed that surface pore waters evolved within a few hours from concentrations close to those of seawater just after

exposure to values close to those found at depth, suggesting that during emersion, concentration changes were mostly due to a transport process, and not to reactions. Solute transport on a distance of several centimetres in the sandy porous media may take about 1 month for transport by molecular diffusion (Huettel et al., 2014, 1996), which is much longer than observation. Therefore, the fluxes we observed were most likely explained by vertical advection of pore water. This flow can be interpreted as the vertical component of the seepage flow due to tidally-driven circulation of pore water at low tide (Chassagne et al., 2012; Robinson et al., 2007; Rocha et al., 2009; Xin et al., 2015). Dissolved Fe, Mn, ammonium, and DIP showed constant profiles during emersion at stations 1, 2, and 3 (Fig. 4). These dissolved species had a concentration close to that of seawater in the upper 20 cm, and higher concentrations below. The gradient did not move upward during emersion (within the 10-cm vertical resolution of our sampling), unlike dissolved oxygen and nitrate. This suggests that the reduced compounds did not cross the redox boundary imposed by the depth of oxygen penetration determined by the intensity of the waves and swash.

To summarize, the coupling between wave and tidally driven circulation on the wide seepage face corresponded first to a mixture with seawater of the 10–20 cm upper pore water during flood and immersion due to swash and wave pumping, and second to a 10–20 cm advective upward flux of waters at low tide due to pore water circulation in the intertidal circulation cell (Fig. 7). Seeping waters were mostly seawater that had filled the pores in the first decimetre of the beach aquifer during the previous high tide due to waves. Nutrients and reduced compounds produced during organic matter mineralization accumulated in pore water below the interface between the layer disturbed by wave mixing and the undisturbed aquifer. The flux of these components to the seawater may have occurred each time this interface was eroded, for example when wave energy increased after a period of lower energy.

5. Conclusions

The Ker Chalon beach is a pocket beach with fine sand that forms a small unconfined permeable aquifer. Pore water chemistry monitoring shows that biogeochemical processes are coupled to tidally-driven recirculation. Pore water composition is progressively modified along the intertidal cross-shore transect because of the production and consumption of chemical components due to marine organic matter mineralization. Aerobic processes occur in the upper part of the beach, where pore water are young. Seawards, in the seepage area, dissolved oxygen becomes depleted and anaerobic processes continue the mineralization of organic matter in older pore waters. Therefore, the porous media located a few decimetres below the beach face shifts from oxic in the upper beach to anoxic in the lower beach. In this beach, mineralization of organic matter creates a

horizontal redox stratification, not a vertical one as commonly observed in benthic environments, because of tidal pumping.

Wave-driven advection of pore water ventilates the upper part of the aquifer along the whole cross-shore transect. A vertical redox gradient occurs in the lower beach mostly because of this advection. The redox boundary that coincides with the depth of wave influence becomes shallower during emersion due to tidal pumping, inducing a flux of pore water toward the beach face. Then, the water that seeps out the aquifer at low tide is mostly composed of oxic pore water that recently entered the upper aquifer through wave pumping.

This mechanism implies that fluxes of nutrients and of reduced compounds occur mostly during immersion when redox boundary is eroded by wave-pumping after it has been shifted upwards during the previous low tide. Therefore, an increase in wave amplitude may deepens the mixed layer and may enhance the flux toward seawater of biogeochemical solutes issuing from organic matter recycling. The low hydraulic conductivity of Ker Chalon beach does not permit a rapid renewal of pore water in the aquifer. However, both tidally-driven and wave-driven recirculation of seawater allow the beach aquifer to be a bioreactor for marine organic matter mineralization that prevents accumulation of organic matter and metabolic products. The low extension of the studied aquifer, typical of pocket beaches, limits the connection with continental groundwater, unlike many other tidal environments, even if recirculating seawater comprises a high percentage of SGD (Slomp and Van Cappellen, 2004; Knee and Paytan, 2011; Anschutz et al., 2016; Tamborski et al., 2017). As such, the beach provides the coastal environment only with recycled nutrients, and not with new nutrients. This is a distinguishing feature of pocket beaches. This general characteristic of pocket beaches, however, should not blind us to the fact that the temporal and spatial properties of SGD are site-specific (e.g., Urish and McKenna, 2004; Pain et al., 2019). As a result, it is difficult to extrapolate across all pocket beaches from a single example. Other pocket beaches now deserve to be studied and compared to the example of Yeu island described here.

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Figure captions

Figure 1: Map of the Yeu Island and location of the Ker Chalon beach (A). Graphical representation of the sampling site on the tidal beach (B). Cross-shore topographic profiles obtained in October 2012 and April 2013 with the position of pore water collectors (C)

Figure 2: Dissolved oxygen saturation and water head recorded in October 2012 and April 2013 in sediment of the Ker Chalon beach at the station along the cross-shore transect (Fig. 1). Data were obtained from autonomous probes buried in the upper part of the water-saturated aquifer at low tide.

Figure 3: Salinity and concentrations of dissolved O_2 , nitrate, nitrite, ammonium, dissolved manganese, dissolved iron, inorganic phosphorus (DIP), and dissolved silica (DSi) in pore waters from cross-shore transects of the Ker Chalon beach in October 2012 (left column) and April 2013 (right column). Transects were replicated three times at low tide during each campaign. The position 0 m corresponds to the base of the dune. Seawater values are represented by a star at the 150 m position.

Figure 4: Vertical profiles of salinity and concentrations of dissolved O_2 , nitrate, nitrite, ammonium, dissolved manganese, dissolved iron, and inorganic phosphorus (DIP) in pore waters from pore water collectors deployed on the Ker Chalon beach in October 2012 (left) and April 2013 (right). Sampling was replicated several times during one emersion period to monitor the evolution of measured parameters in an hourly time scale. Such sampling was repeated at two or three low tides during each campaign; the evolution was the same each time. Profiles presented here are those from samples collected in October 17th 2012, and April 11th 2013. The positions of pore water collectors indicated on Fig. 1 are represented by vertical bars.

Figure 5: Total alkalinity, pH, and pCO_2 values in pore waters from cross-shore transects of the Ker Chalon beach in October 2012 (left) and April 2013 (right), and $\delta^{13}\text{-DIC}$ in pore waters from October 2012. Transects were replicated three times at low tide during each campaign. The position 0 m corresponds to the base of the dune. Seawater values are represented by a star at the 150 m position.

Figure 6. (A) TA concentrations versus DIC, and (B) Keeling plot of Ker Chalon beach pore waters collected in October 2012. The intercept at the origin of the Keeling plot gives the isotopic signature of the source of the mineralized organic carbon that produces additional DIC relative to seawater DIC concentration.

Figure 7: Schematic representation of hydrologic circulation during flow (left) and low tide (right) showing potential source contributing to ventilation of pore waters. 1: Parcels of meteoritic waters mixed with saline water; 2: Seeping of pore water at the slope break; 3: Vertical component of tidally-driven circulation pushing the redox gradient upwards. The dotted line indicates the boundary between the aquifer dominated by aerobic and anaerobic organic matter (OM) mineralization processes..

FIGURE 1

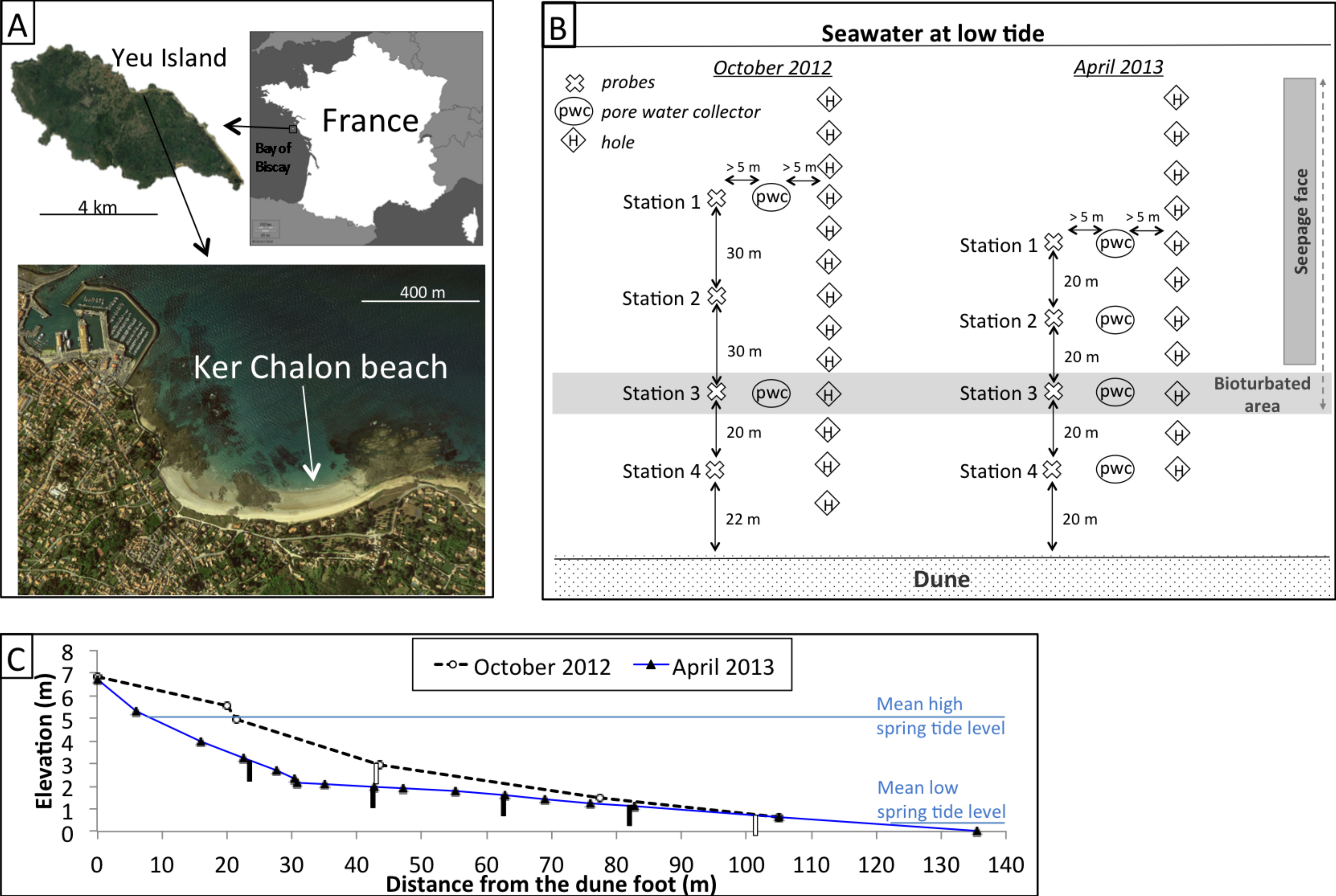


FIGURE 2

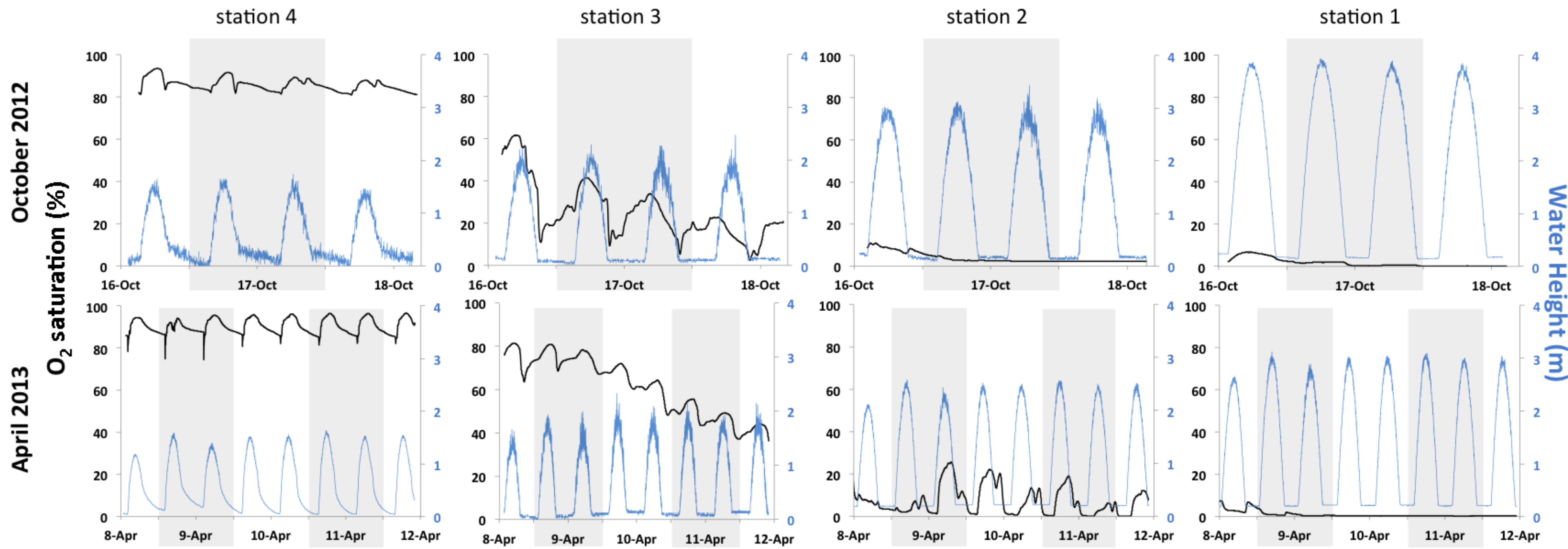


FIGURE 3

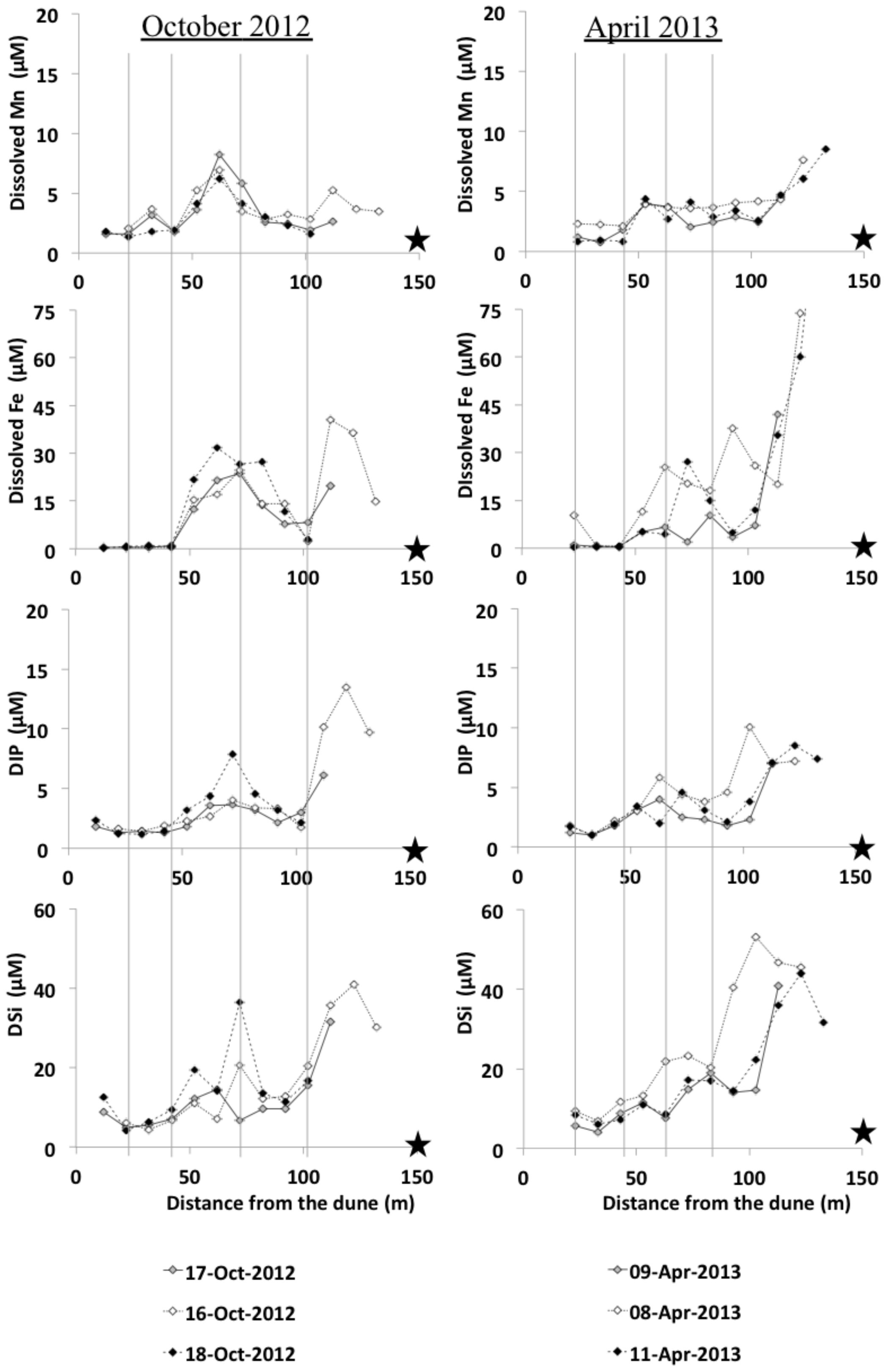
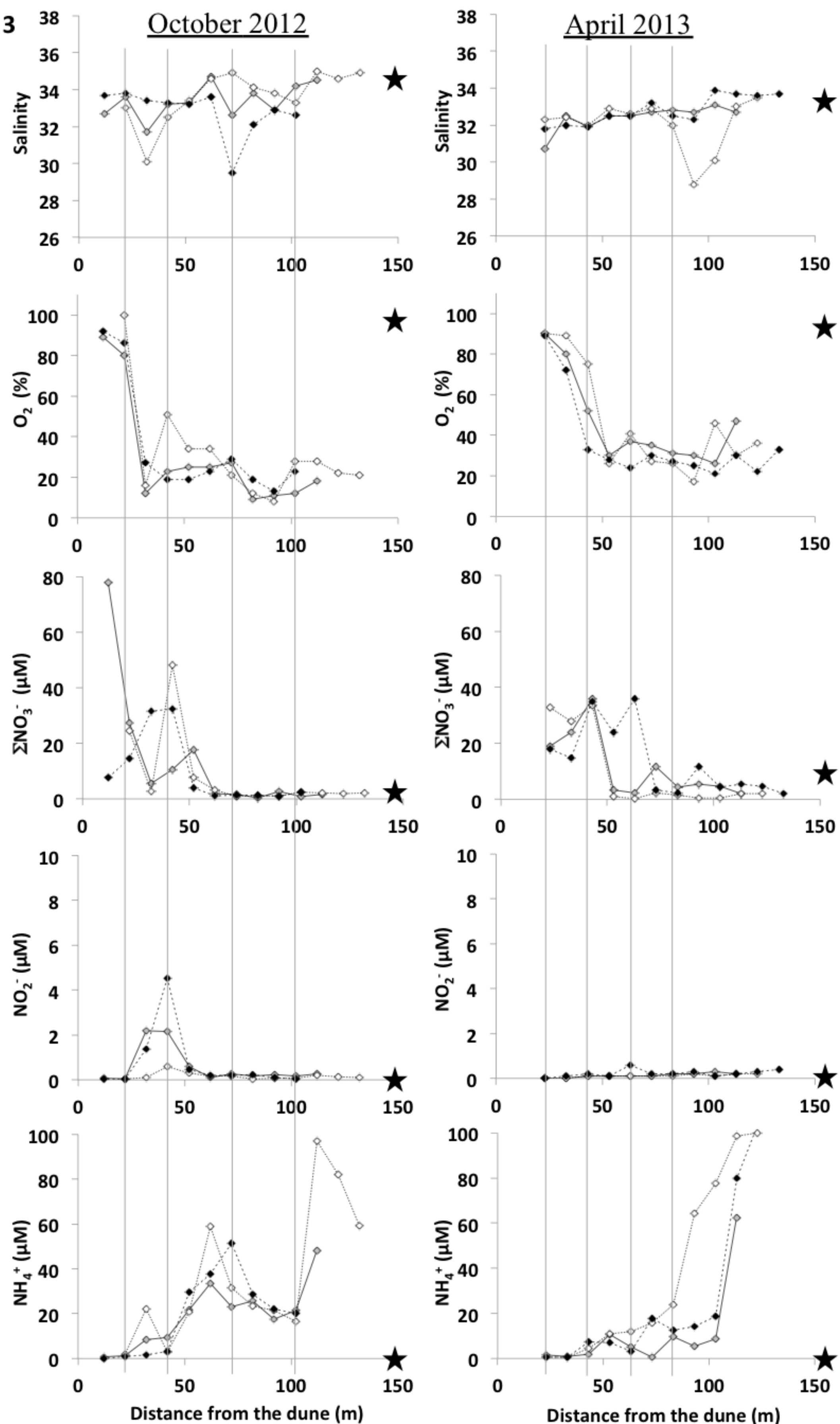


FIGURE 4 suite

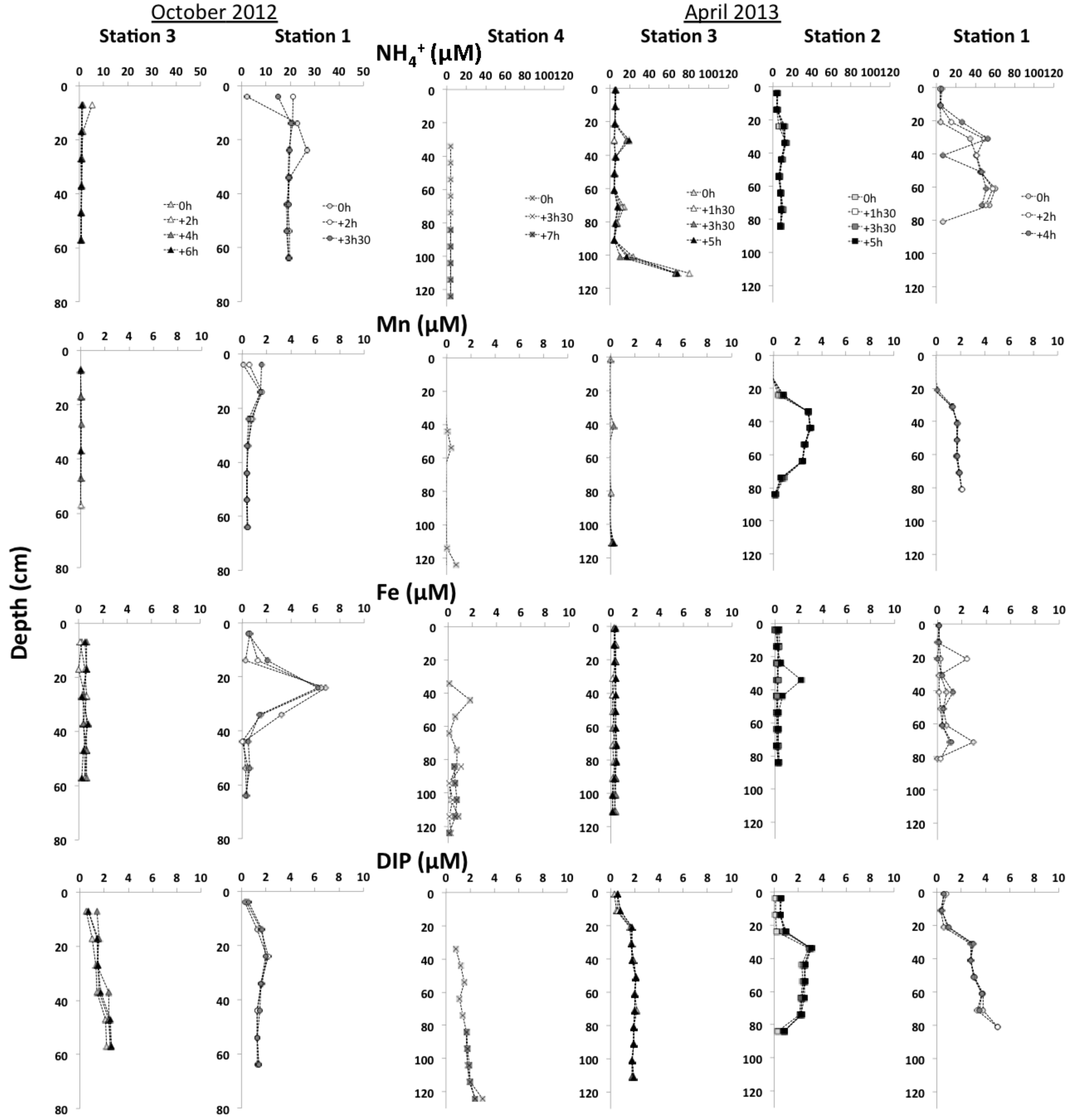


FIGURE 4

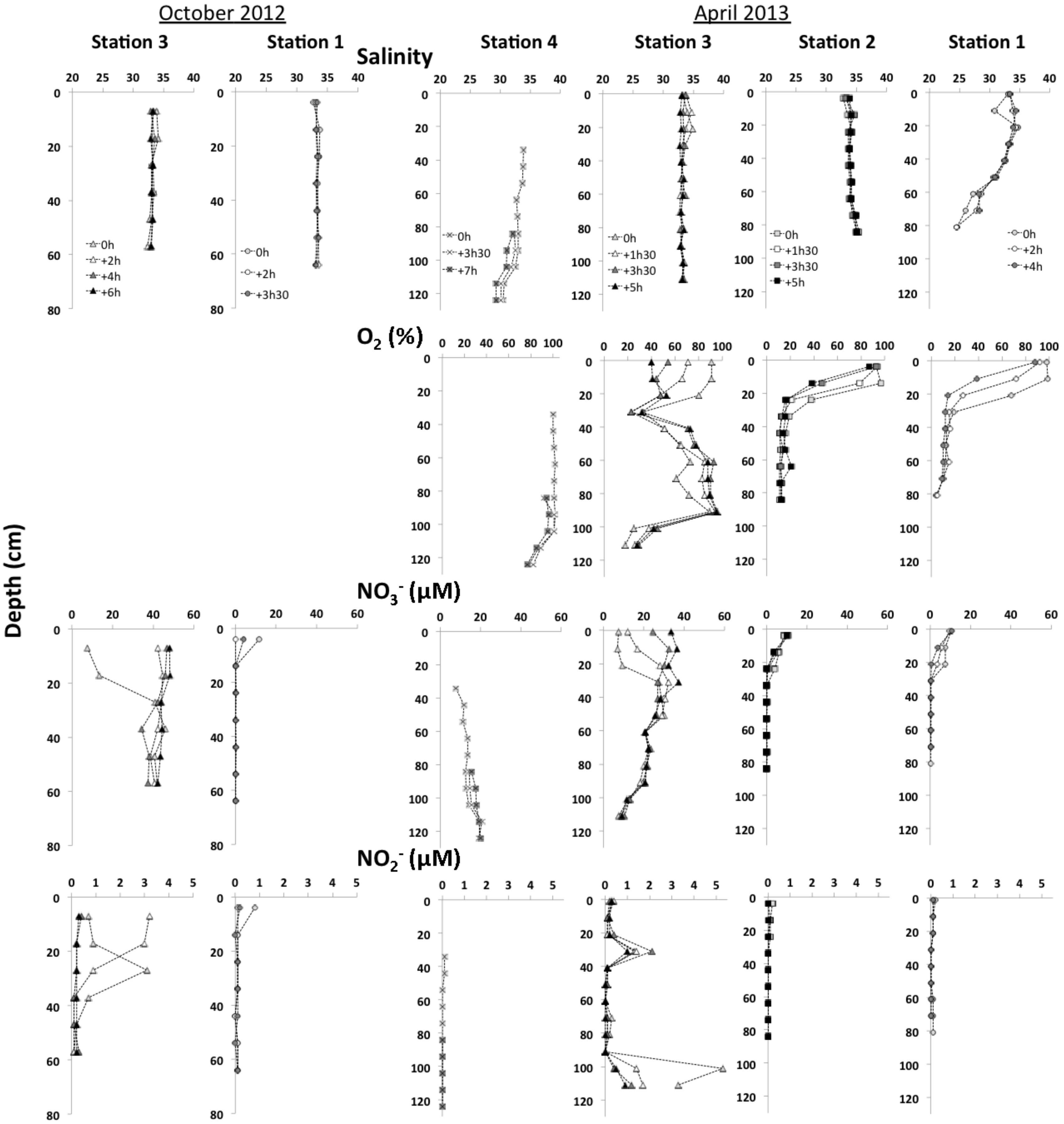


FIGURE 5

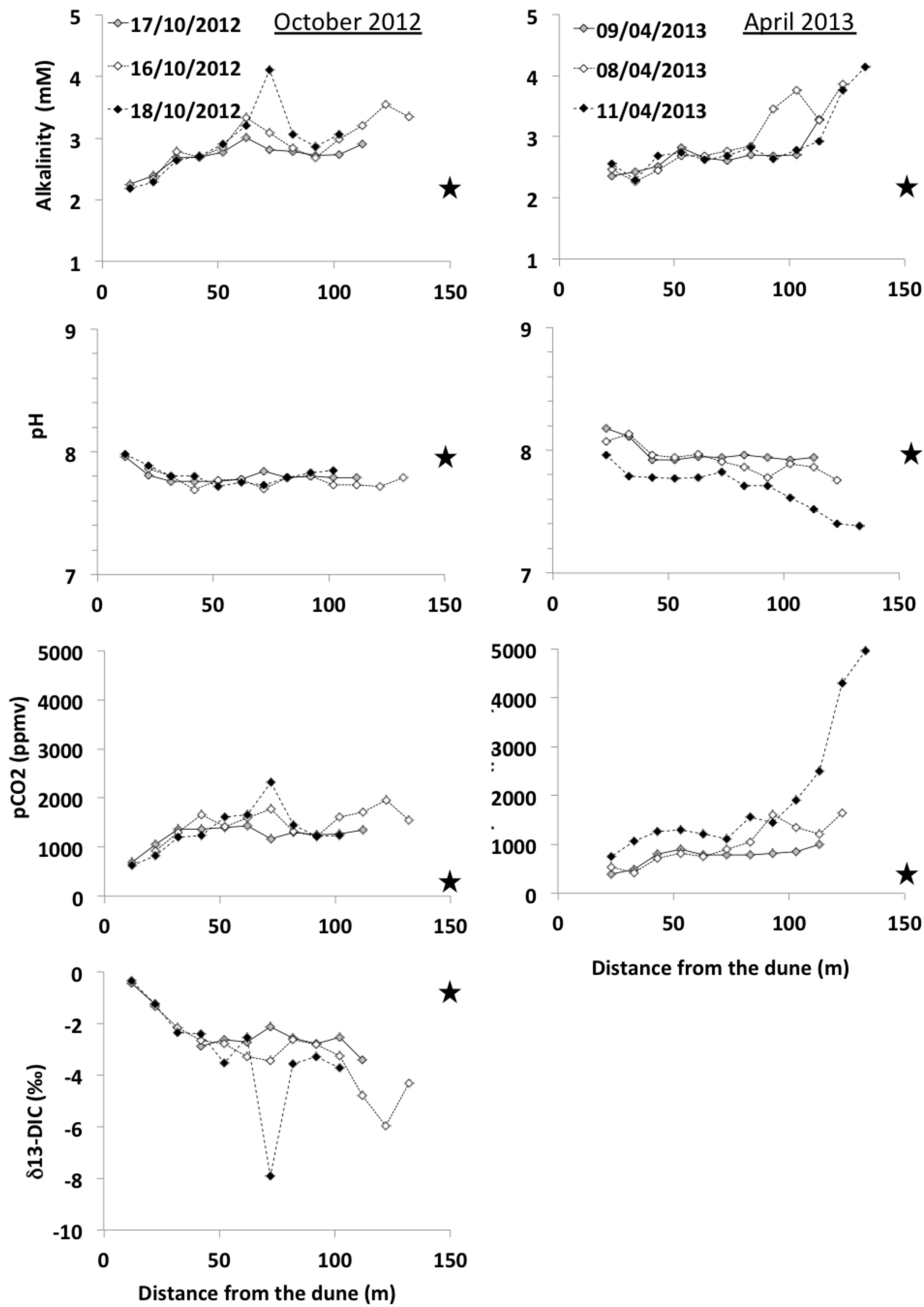


FIGURE 6

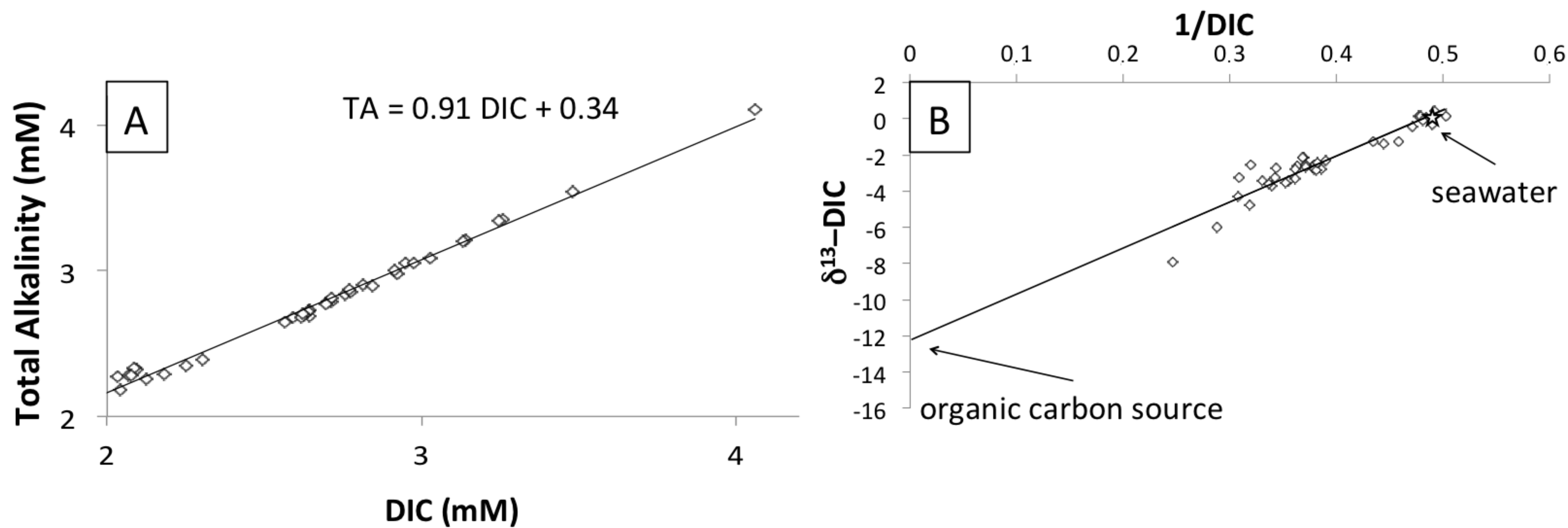


FIGURE 7

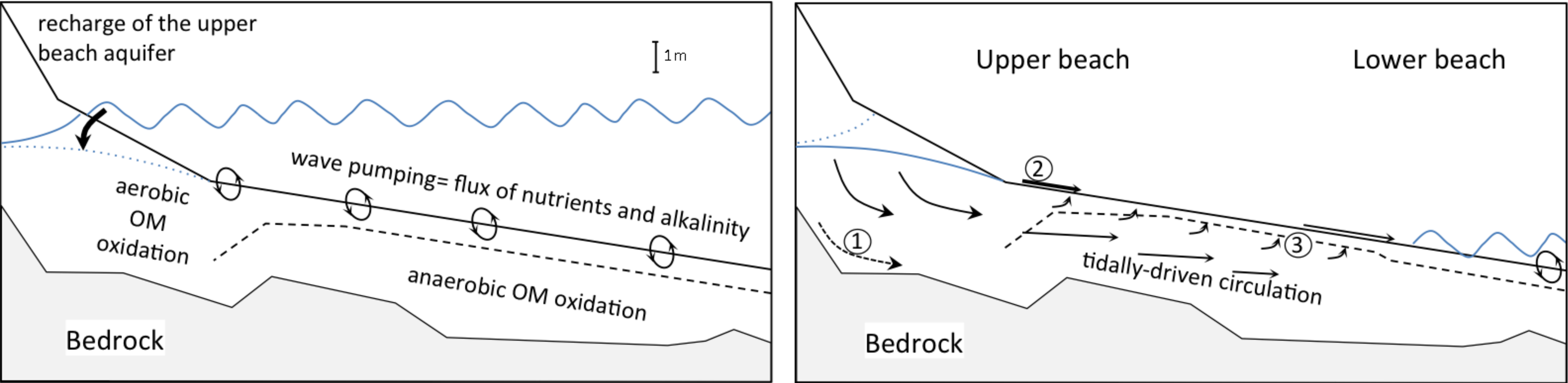


Table 1. Characteristics of sediments in the seepage zone of the Ker Chalon beach (mean $\pm$ standard deviation, N=6 for grain size and N=11 for other parameters)

Median grain size (μm)	166 $\pm$ 10
Particulate organic carbon (% dry sediment)	0.20 $\pm$ 0.08
Particulate CaCO_3 (% dry sediment)	18.2 $\pm$ 4.6
HCl-extracted Fe ($\mu\text{mol g}^{-1}$)	48.6 $\pm$ 3.5
HCl-extracted Mn ($\mu\text{mol g}^{-1}$)	1.21 $\pm$ 0.2
HCl-extracted P ($\mu\text{mol g}^{-1}$)	12.1 $\pm$ 2.8