Solute transfer across the sediment surface of a eutrophic lake: I. Porewater profiles from dialysis samplers

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ABSTRACT

Porewater profiles often are used to identify and quantify important biogeochemical processes occurring in lake sediments. In this study, multiple porewater profiles were obtained from two eutrophic Swiss lakes using porewater equilibrators (peepers) in order to examine spatial and seasonal trends in biogeochemical processes. Variability in profile shapes and concentrations was small on spatial scales of a few meters, but the uncertainty in calculated diffusive fluxes across the sediment surface was, on average, 35%. Focusing of Fe and Mn oxides toward the lake center resulted in systematic increases in porewater concentrations and diffusive fluxes of Fe²⁺ and Mn²⁺ with increasing water depth; these fluxes are postulated to be regulated by the pH-dependent dissolution of reduced-metal phases. Despite higher concentrations of inorganic carbon, NH₄, Si and P in pelagic compared to littoral sites, diffusive fluxes of these substances across the sediment surface increased only slightly or not at all with increasing water depth. Porewater profiles did reveal temporal changes in Fe2+, Mn2+, Ca2+ and Mg2+ that were an indirect result of the large, seasonal changes in seston deposition, but no clear seasonal variations were found in diffusive fluxes of nutrients across the sediment surface. The intense mineralization occurring at the sediment surface was not reflected in the porewater profiles nor in the calculated diffusive fluxes. Calculated diffusive fluxes across the sediment surface resulted from decomposition occurring primarily in the top 5-7 cm of sediment. Diffusive fluxes from this subsurface mineralization were equal to the solute release from mineralization occurring at the sediment-water interface. Buried organic matter acts as a memory of previous lake conditions; it will require at least a decade before reductions in nutrient inputs to lakes fully reduce the diffusive fluxes into the lake from the buried reservoir of organic matter.

Introduction

In both the oceanographic and limnological studies, porewater profiles frequently are used to identify biogeochemical processes or to calculate process rates. Processes amenable to such study include mineral dissolution, authigenic mineral formation,

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organic matter decomposition, and microbial dynamics (e.g., Lerman, 1979; Froehlich et al., 1979; Berner, 1980). Solute concentrations may indicate the existence of mineral phases (e.g., Emerson, 1976; Cook, 1984) or of microbial processes (e.g., Lovely and Goodwin, 1988). Diffusive fluxes calculated from concentration gradients in porewater profiles have been equated to rates of microbial processes (e.g., Rudd et al., 1986; Lovely and Klug, 1986; Canfield, 1991), rates of mineral dissolution or precipitation (e.g., Lerman and Liezke, 1977; Wersin et al., 1991), and rates of solute transfer from the sediments to the water column (e.g., Rudd et al., 1986, 1990; Carignan and Lean, 1991). Problems inherent in the interpretation of porewater profiles include reproducibility, the effects of depth resolution, the assumption of steady state, internal element cycles, and spatial and temporal variability.

The objective of this paper is to demonstrate the spatial and temporal variability in porewater profiles. This serves not only to point out appropriate methodologies and interpretations of data, but also to illustrate the dynamics of the biogeochemical processes involved in early diagenesis of lake sediments. Specifically, comparison of a large number of solute profiles from pore waters of two lakes allows us to address four questions. First, what is the uncertainty in measured concentrations and calculated fluxes? Reproducibility depends on both the precision of the technique and the spatial heterogeneity of the sediments. Artifacts of the dialysis sampling technique (Hesslein, 1976) have been examined previously (e.g., Carignan, 1984; Adams, 1991; Brandl and Hanselmann, 1991; Carignan et al., 1994). However, although previous papers have presented replicate porewater profiles (e.g., Carignan, 1984; Carignan et al. 1994), the reproducibility has seldom been quantitatively examined. Comparison of profiles among lakes (e.g., Rudd et al., 1986; Cook et al., 1987) or use of calculated diffusive fluxes in lake mass balances (e.g., Carignan and Lean, 1991; Rudd et al., 1990) requires knowledge of the uncertainty about individual measurements. The sampling of this study allows a preliminary estimate of the uncertainty due to small-scale spatial variability.

The second question addressed in this study concerns the sediment depth at which diagenetic reactions occur. Intense mineralization of organic matter occurs in the first 1–3 mm sediments (e.g., Lovely and Klug, 1986; Sweerts et al., 1986, 1989). On the other hand, Carignan and Lean (1991) reported that significant release of nutrients continued down to sediment depths of 50 cm in a mesotrophic lake. Fineresolution sampling would be required to characterize sharp concentration gradients resulting from surface mineralization. Development of microelectrodes (e.g., Revsbech et al., 1980, 1988) and gel porewater collectors (Davison et al., 1991, 1994; Davison and Zhang, 1994) has begun to allow us to measure profiles for some species with a resolution of < 250 µm. Unfortunately, microelectrodes are not available for all species, and gel collectors can be employed only by differs in shallow lakes (Davison et al., 1994). The effects of coarse-resolution sampling on major ions generally is ignored. In this study we compared fluxes calculated from profiles with 0.5 and 1.5 cm resolution and found few systematic differences. By comparison of diffusive fluxes calculated from these coarse-resolution profiles with fluxes measured in benthic chambers we demonstrate the relative importance of surface and deep mineralization.

Third, systematic spatial variations in solute concentrations in pore waters and in diffusive fluxes across the sediment surface were investigated. Focusing of

organic matter (Davis, 1968; Lehman, 1975) could result in higher rates of organic matter decomposition and nutrient release in the deepest areas of lakes. Such a pattern was reported for one mesotrophic lake (Carignan and Lean, 1991) but was not observed in an oligotrophic lake (Sherman et al., 1994). Systematic spatial variation in rates of decomposition might offset systematic variations in organic matter deposition.

Finally, this study examined the temporal variations in pore water profiles. Temporal variability could be problematic with regards to pore water measurements. Profiles are assumed to be at steady state for the calculation of diffusive fluxes (e.g., Lerman, 1979), and the long equilibration period for porewater dialysis samplers (2–3 weeks; Brandl and Hanselmann, 1991; Carignan et al., 1994) requires that no significant changes occur over this timeframe. However, large seasonal and annual variations in nutrient inputs to lake sediments occur. Sedimentation rates in summer may be more than 10-fold higher than in winter, and interannual variations in sedimentation rates may approach 25% (Gächter et al., 1989; Urban et al., 1994). Such variability in inputs to sediments should result in a corresponding variability in element release to porewaters unless the diagenetic processes are much slower than the variations in inputs. Results from this study clarify the four questions above and point to serious limitations of the dialysis sampling technique.

Methods

Site description

The major focus of this study was Lake Sempach. This lake ocupies a deep, glacially-scoured river valley in central Switzerland. It has an area of 14.4 km², a maximum depth of 86 m, and an average depth of 46 m (Gächter et al., 1983). Because the catchment: lake area ratio (4.3) is small, the water retention time is relatively long (17 yr). Situated among rich farmlands, the lake receives considerable nutrient inputs from agricultural runoff (Baumann, 1988) and is, consequently, eutrophic. Varved sediments, an indication of anoxia in bottom waters, have been deposited in the center of the lake since about 1930 to the present (Züllig, 1956, 1982; Gächter and Meyer, 1990). Installation of sewage treatment plants around the lake decreased total phosphorus concentrations from above 150 to around 100 µg/L (Stadelmann, 1988). For the past 10 years artificial aeration and forced circulation have been utilized in an attempt to reduce internal recycling of P (Gächter et al., 1989; Stadelmann, 1988). These restoration measures have succeeded in maintaining oxygen concentrations above 4 mg/L in the hypolimnion, but internal recycling of P has not been reduced significantly (Gächter and Wüest, 1993).

To augment the discussion of profile reproducibility and spatial variability, additional porewater profiles were obtained from Lake Baldegg. This nearby lake has a similar morphometry (max. depth 65 m; area 5.2 km²; Stadelmann, 1984), and is similarly eutrophic. Laminated sediments in the center of L. Baldegg have been deposited since about 1880 (Niessen and Sturm, 1987; Züllig, 1982). Despite a reduction in P inputs and 10 years of artificial aeration, the lake remains highly eutrophic (Wüest et al., 1991).

Porewater collection and analysis

Porewater profiles were measured on a total of 10 occasions in L. Sempach between September 1989 and June 1993. On seven occasions (Sept. 1989, June 1990, July 1990, October 1990, April 1991, April 1992, June 1993) profiles were obtained from the deepest area (86 m) of the lake. Profiles from littoral areas (water depth < 20 m) were collected twice (March 1991, October 1992), and in August 1992, profiles were obtained from an intermediate water depth (50 m). In addition, porewater profiles from L. Baldegg were obtained in November 1992, January 1994, and March 1994 from pelagic regions (56–65 m water depth), and in November 1992 also from 25-m water depth.

Porewater equilibrators (peepers; Hesslein, 1976) were used on all occasions to obtain the profiles. Peepers consisted of a 50 × 15 cm solid, plexiglass frame into which were milled cells (1 or 0.5×10 cm) at a spacing of either 0.5 or 1.5 cm. The cells were filled with distilled, deionized water, and a 0.2 µm filter (HT 200, Tuffryn polysulfone membrane; Gelman Inc.) was laid over the surface of the frame and held in place by an outer frame of plexiglass. To remove oxygen from the water in the cells, the peeper was immersed in a tank of distilled water that was then purged for 24 hours with nitrogen (cf. Carignan et al., 1994). When the peeper is placed in the sediments, solutes from porewater in contact with the filter diffuse into the cells such that, after equilibrium is reached, concentrations inside each cell equal those at the corresponding depth in the porewater. A two week period was allowed for equilibration (Hesslein, 1976; Brandl and Hanselmann, 1991). Peepers were mounted on a steel tripod that was lowered to the sediment surface. On several occasions, the orientation of the peepers was inspected by means of a video camera mounted on the same tripod. Exposure of the peeper to lake water following retrieval from the sediments lasted no more than five minutes.

The cells of the peeper were sampled onboard ship immediately upon retrieval from the lake. Water from within the cells was withdrawn into pipets and immediately injected into containers containing appropriate fixative agents. Samples for analysis of Fe, Mn and major cations were acidified with Suprapure $^{\rm TM}$ HNO3. Samples for CH4 analysis received NaOH to raise the pH above 10, and zinc acetate was added to samples for sulfide analysis. Samples for measurement of pH and alkalinity were stored in syringes until analyzed. The sampling of peepers was done as quickly as possible to prevent evasion of gaseous species and invasion of oxygen; the entire sampling generally required 15–30 min.

Analysis for all species generally was completed within 48 hours after retrieval from the lake. Major cations (NH $_4^+$, Ca $^{2+}$, Mg $^{2+}$, Na $^+$, K $^+$) and anions (Cl $^-$, NO $_3^-$, SO $_4^{2-}$) were analyzed by ion chromatography(Metrohm). Silica and PO $_4^{3-}$ were analyzed photometrically with molybdenum blue (APHA); total P was measured identically following digestion with K $_2$ S $_2$ O $_8$. Alkalinity was determined by acid titration to pH 4.3, CH $_4$ was measured by gas chromatography with flame ionization detection, and Fe and Mn were measured by atomic absorption spectrophotometry.

Calculation of the diffusive fluxes of solutes across the sediment-water interface generally is based on application of Fick's Law to the concentration profile.

At steady state, the flux (F) is equal to the product of the concentration gradient (dC/dz) and the diffusion coefficient (D):

$$F = D \cdot dC/dz \tag{1}$$

The validity of the assumption of steady state will be discussed below in light of our results. Only in the diffusive boundary layer (DBL) can the molecular diffusion coefficient in water (D) be used. Because of the coarse resolution of the profiles, it was not possible to determine the concentration gradient in the DBL. Instead, concentration gradients within the sediments are used together with an effective diffusion coefficient $(D_{\rm eff})$, defined as:

$$D_{\rm eff} = \phi^{\rm n} \, \mathbf{D} \tag{2}$$

where ϕ is the porosity, and n is a value between 1 and 3 (Lerman, 1979). Porosity, measured as weightloss upon drying of sediments at 60 °C, was 0.92 for pelagic and 0.80 for littoral surface sediments. In this paper, we use a value for n of 1. Values of D were taken from Li and Gregory (1974) and Lerman (1979) corrected to a temperature of 5 °C. In this paper, only diffusion of major ions was considered; complexes (e.g., Furrer and Wehrli, 1993) and electrostatic interactions among ions (e.g., Lasaga, 1979) were ignored. Numerous authors select an arbitrary number of points from the concentration profiles and apply a linear regression to calculate the concentration gradient at the sediment surface (e.g., Rudd et al., 1986; Brandl and Hanselmann, 1991; Sherman et al., 1994). We followed this procedure only for solutes whose concentrations change sharply across the sediment surface (viz., NO₃, SO₄-, PO₄-, Fe²⁺, Mn²⁺). For many substances that are released during decomposition (e.g., HCO₃, NH₄, Si, CH₄), concentration profiles follow a smooth curve that can be fit to a second- or third-order polynomial (cf. Berner, 1980). The concentration gradient at the interface is equal to the first derivative of the polynomial at z = 0. The polynomial curve fitting smooths the profiles and minimizes the effects of measurement errors on the calculated fluxes.

One major uncertainty in calculating diffusive fluxes is the location of the sediment surface within the porewater profile. We used the appearance of a brown layer of iron oxides on the peeper surface to indicate the depth of the sediment-water interface. The formation of iron oxyhydroxides does not necessarily occur at the sediment surface, but at the oxycline. The presence of a mat of Beggiotoa at the sediment surface in pelagic sites of both lakes Sempach and Baldegg strongly suggets that the oxycline lies at the sediment surface; observations with a video camera confirmed that the layer of iron oxyhydroxides occurred at the sediment surface in pelagic sites. Equating of the sediment surface and the oxycline may not be valid in littoral sediments. However, the depth of iron oxyhydroxide precipitation coincided with sharp changes in slopes of concentration profiles even of species (e.g., Si) that should not be sensitive to oxygen.

Results

Identical peepers spaced three meters apart yielded solute profiles that were very similar in general shape and concentration (Fig. 1). To conserve space, Fig. 1 presents only one set of replicate profiles for sulfate and one for phosphate; the remaining eleven sets of replicate profiles also showed very close correspondence between the concentration profiles. Such close correspondence was observed in both littoral and pelagic sediments of both L. Sempach and L. Baldegg, and in all months (April, June, August, October, November) for which such paired profiles were collected.

The variability among profiles also may be assessed by comparing concentrations below about 20 cm in the profiles. At this depth below the sediment surface, concentrations of many solutes (e.g., CH_4 , NH_4^+ , Fe^{2+} , HCO_3^- , Ca^{2+} , SO_4^{2-}) approach constant values. Only for SO_4^{2-} did t-tests consistently (i.e., 5 of 5 pairs for SO_4^{2-} , ≤ 1 of 5 for all other solutes) indicate that these "constant" concentrations were significantly different among pairs of replicate profiles. The range of variation $(100 \times S.D./mean)$ in the concentrations varied from 3% for Ca^{2+} and Mg^{2+} to 70% for SO_4^{2-} ; relative standard deviations for all solutes except SO_4^{2-} were below 20%. The large uncertainty in porewater SO_4^{2-} concentrations results from the low concentrations that are near the detection limit of the analytical technique; analytical errors are probably as large as the actual variability within the pore waters. It even has been suggested that the low concentrations of SO_4^{2-} measured at depth in lake pore waters are artifacts that result from oxidation of reduced S by oxygen diffusing

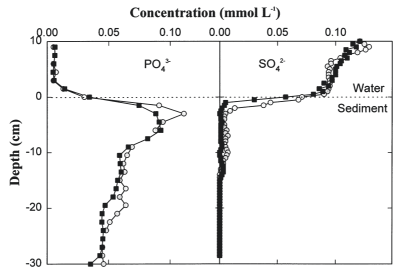


Figure 1. Replicate porewater profiles. Replicability of profile shapes and concentrations generally is good as shown here for o-PO $_4^{3-}$ and SO $_4^{2-}$. The PO $_4^{3-}$ profiles were from 86-m water depth in L. Sempach in October, 1990; SO $_4^{2-}$ profiles are from 50-m water depth in L. Sempach in August, 1992

Table 1. Variability among diffusive fluxes calculated from identical replicate peepers

Solute	NO ₃	SO ₄ ²⁻	Cl-	Ca ²⁺	Mg ²⁺
Number of replicates	2	2	2	1	1 3
Relative error (%) ^a	18, 28	51, 76	4–18	17	

a Relative error = $100 \cdot (x_1 - x_2)/x_1 + x_2$

out of the peeper frame (Carignan et al., 1994). The similarity in solute concentrations and profile shapes suggests that spatial variability is small over the scale of a few meters in both littoral and pelagic sediments during all seasons in these eutrophic lakes.

Despite the appearance of low spatial variability, there is a large uncertainty in the calculated diffusive fluxes across the sediment surface. On four occasions, one or more solute was measured in replicate porewater profiles obtained within 3 m of each other with identical peepers; for any single solute only one or two pairs of profiles were obtained. Relative errors among the diffusive fluxes calculated from these replicate profiles ranged from 3% to 76% (Table 1). These estimates of the uncertainty in fluxes are based on a limited number of replicate measurements. If the variance in the relative errors were the same for all solutes (additional data are needed to test this assumption), the relative standard deviation for all of the paired fluxes would be 37%.

Systematic changes with water depth were observed in ion concentrations. Littoral porewater profiles were characterized by lower concentrations of Mn, Fe, P, Si, NH $_4^+$ (Fig. 2) and alkalinity when compared to pelagic profiles. These differences in concentrations were observed in both lakes Sempach and Baldegg on all sampling dates. Diffusive fluxes also differed significantly for many solutes between littoral and pelagic regions (Table 2; as discussed below, the absence of seasonal variations allows a comparison of littoral and pelagic sites across seasons). Diffusive fluxes of NH $_4^+$, Fe $^{2+}$, and Mn $^{2+}$ were significantly greater in pelagic than in littoral regions. Only the fluxes of SO $_4^{2-}$ and P were significantly larger in littoral than in pelagic areas of one or both lakes (Table 2).

There was little evidence of seasonal changes in diffusive fluxes of most solutes across the sediment-water interface. Only for Mn²⁺, NO₃ and Ca²⁺ were diffusive fluxes higher during the period of lake stratification than during the period of mixis. For all other solutes neither grouping of fluxes into stratified and non-stratified lake conditions nor into seasons of spring, summer, fall and winter revealed any significant trends. For solutes for which no systematic seasonal changes in fluxes occurred, all of the profiles gathered at the deepest site may be pooled to estimate the uncertainty in the average annual flux. The relative standard deviations among 7–11 fluxes calculated for the deepest site of L. Sempach ranged from 30 to 80% for individual solutes.

While it is easy to assess the variability in diffusive fluxes, it is more difficult to assess the accuracy of the calculated fluxes. One factor that might contribute to inaccuracy in the calculations is the depth resolution with which the concentration gradient is assessed. In this study, profiles were collected with depth resolutions of

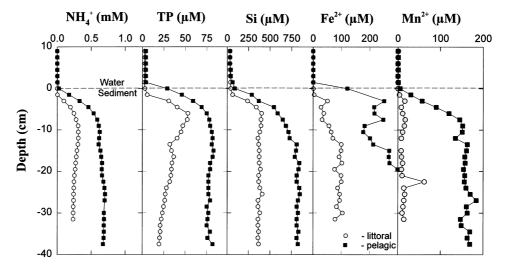


Figure 2. Comparison of littoral and pelagic porewater profiles. Concentrations of all species released from decomposition of biomass (DIC, P, NH₄⁺, Si, CH₄) are higher in pelagic sites (L. Sempach, 86-m water depth, April 1991) compared to littoral sites (L. Sempach, 15-m water depth, March 1991). Profiles of DIC and CH₄ are not shown. Higher concentrations of Fe and Mn also are observed in the pelagic sites. Concentrations of NO₃⁻ and SO₄²⁻ (not shown) are similar in littoral and pelagic sediments

either 5- or 15-mm. On seven occasions, parallel profiles of some solutes were collected with the two peeper styles. For any given solute, no more than four such replicate profiles were measured. For 14 of the 25 paired profiles diffusive fluxes were higher in the profiles with the finer depth resolution (viz., 5 mm). However, for all individual solutes (except NO_3) as well as for all solutes collectively there was no statistically significant difference (paired t-test) in the fluxes calculated from profiles with 5- and 15-mm resolution.

A further test of the accuracy of the calculated diffusive fluxes across the sediment surface may be made by comparing fluxes calculated from porewater profiles with those measured in benthic chambers. As reported elsewhere (Wehrli et al., 1997), benthic chambers (Lander) after the design of Devol (1987) were deployed on multiple occasions in L. Sempach. These chambers allowed measurement of fluxes with a controlled rate of water mixing within the chamber and within a short enough time period (12–24 hrs) that oxygen depletion did not occur in the chambers. Linear increases or decreases in solute concentrations with time were observed in the Lander chambers. Comparison of diffusive fluxes from the deepest area of the lake (Table 3) indicates that porewater equilibrators and benthic chambers yielded significantly different results for all solutes except CH₄ and Fe²⁺. For solutes with low net fluxes (Ca, Mg, Cl, SO₄²⁻) and large concentrations in the lake water, the large uncertainties in fluxes measured with the Lander preclude meaningful comparison with those calculated from porewater profiles.

Table 2. Comparison of diffusive fluxes (mmol $m^{-2}d^{-1}$) in littoral and pelagic zones of lakes Sempach and Baldegg.

Solute	Lake	Littoral flux	Pelagic flux
HCO ₃	Baldegg ^a Sempach ^b	+ 2.24 + 2.4 ± 0.3 (2)	+ 3.09 + 2.5 ± 0.8 (7)
NH_4^+	Baldegg Sempach	$+ 1.5 + 0.7 \pm 0.01 (2)$	+ 2.5 + 1.3 ± 0.5 (8) *
P	Baldegg Sempach	$+ 0.95 + 0.2 \pm 0.15$ (2)	+0.10 + 0.08 ± 0.05 (6)*
Fe^{2+}	Baldegg Sempach	$+ 0.06 + 0.1 \pm 0.04$ (2)	$+ 0.15 + 0.34 \pm 0.32 (7)*$
Mn^{2+}	Baldegg Sempach	$+ 0.015 + 0.017 \pm 0.002$ (2)	$+ 0.14 + 0.16 \pm 0.08 (7)*$
SO ₄ ²⁻	Baldegg Sempach	-0.15 $-0.32 \pm 0.08 (2)$	-0.14 $-0.14 \pm 0.02 (8)*$
NO_3^-	Baldegg Sempach	-0.14 -0.19 ± 0.05 (2)	-0.18 $-0.14 \pm 0.11 (8)$
$\mathrm{CH_4}$	Baldegg Sempach	na° + 1.44 ± 0.09 (2)	na + 3.1 ± 2.7 (6)
Si	Baldegg Sempach	na + 0.61 ± 0.15 (2)	na $+ 0.58 \pm 0.14 (7)$

^a Measurements made in November.

Table 3. Comparison of diffusive fluxes (mmol m^{-2} d^{-1} \pm S.D.) across the sediment surface at the deepest location in L. Sempach as measured with benthic chambers and calculated from porewater profiles

Solute	Average flux as me	Ratioa	
	Lander (n) ^b	Porewater profiles (n)	
HCO ₃	$13.7 \pm 4.8 (7)$	2.5 ± 0.8 (7)	5.5
CH ₄	$1.6 \pm 2.1 \; (6)$	$3.1 \pm 2.7 (6)$	0.5
NH ⁺ ₄	$2.7 \pm 1.0 (6)$	$1.3 \pm 0.5 (8)$	2.0
$NO_{3}^{\frac{1}{3}}$	$-1.9 \pm 0.6 (8)$	$-0.14 \pm 0.11(8)$	13.8
SO_4^{2-}	$+0.4 \pm 0.6 (5)$	-0.14 ± 0.02 (8)	-2.8
PO ₄ 3-	$0.3 \pm 0.4 (7)$	$0.08 \pm 0.05 (6)$	3.7
Si	$3.7 \pm 0.4 (5)$	$0.6 \pm 0.2 (7)$	6.1
Fe^{2+}	$0.8 \pm 0.6 \ (8)$	$0.34 \pm 0.3 (7)$	2.5
Mn^{2+}	$1.5 \pm 1.9 (8)$	$0.16 \pm 0.08(7)$	9.5

^a Ratio = Lander flux/Peeper flux.

b Mean ± standard deviation based on all measurements at given sites (all months). Number of measurements given in parenthesis; * following numbers indicates that littoral and pelagic rates are significantly (p < 0.05; non-paired t-test) different.

c na – not available.

b n = number of measurements.

Discussion

Depths of solute consumption and release

Numerous recent studies have pointed to the sediment surface as a site of intense diagenetic activity. Comparison of sedimentation rates with sediment accumulation rates has shown that much mineralization (as much as 85% of inputs) occurs within a few millimeters of the surface (e.g., Baker et al., 1989; Höhener, 1990). Use of microelectrodes has demonstrated that oxygen penetration is limited to less than two millimeters in eutrophic lakes (e.g., Sweerts et al., 1989, 1991). Bacterial mats such as found on the surface of the sediments in L. Sempach have been shown to be sites of rapid oxygen consumption, sulfide oxidation, and nitrate reduction (e.g., Cohen et al., 1977; Jorgensen and Revsbech, 1983; Sweerts et al., 1990).

This study demonstrates that porewater profiles with a depth resolution of 5- to 15-mm do not reflect the rapid processes occurring at the sediment surface. This conclusion is supported by several lines of evidence. First, diffusive fluxes across the sediment surface as calculated from porewater profiles were much lower than those measured with benthic chambers (Table 3). Although this could be an artifact of the Lander technique (Wehrli et al., 1997), such discrepancies have been interpreted before as evidence of rapid surface reactions (e.g., Johnson et al., 1992; Thamdrup et al., 1994).

A second line of evidence is that the difference between sedimentation and burial is much larger than the diffusive flux out of the sediments of numerous substances (Table 4). The diffusive fluxes for C, N, and P are only 30–90% of those required to balance the sedimentation and burial fluxes. The discrepancy is greater for P (40% of sedimented P is not measured in either the diffusive or burial fluxes) and N (28%) than for C (4%) as a result of the more rapid mineralization of organic N and P as well as the rapid release of P from Fe and Mn oxides that are reduced at the sediment surface (Hupfer et al., 1995). Diffusive fluxes of Fe²⁺ are higher than predicted due to the rapid oxidation of Fe(II) in the water column; the sedimentation rate for Fe is probably underestimated even by the trap immediately (50 cm) above the sediment surface. The discrepancy between predicted and observed diffusive fluxes is most pronounced for Mn; diffusion (0.16 mmol m⁻² d⁻¹) plus burial (0.1 mmol m⁻² d⁻¹) account for only 17% of the amount deposited on the sediment surface at the deepest site in L. Sempach. Furthermore, the Mn present even in the uppermost 2 mm of sediment at this site is already reduced (Friedl et al., 1997); clearly, the porewater profiles fail to reflect the initial, rapid reduction and efflux of Mn²⁺ from the sediments. For N, P and Mn, the discrepancies between diffusive, sedimentation, and burial fluxes are greater than the uncertainties in any of the fluxes.

A third indication that surface processes are not captured in the porewater profiles is that the diffusive fluxes across the sediment surface of oxidized mineralization products ($\rm CO_2 + HCO_3^- = 3.7~mmol~m^{-2}~d^{-1}$) plus products potentially oxidized in the hypolimnion ($\rm NH_4^+$, $\rm CH_4$, $\rm Mn^{2+}$, $\rm Fe^{2+}$; 9 mmol m⁻² d⁻¹ combined as oxygen equivalents) equal only 87% of the sum (15 mmol m⁻² d⁻¹ as oxygen equivalents) of the hypolimnetic oxygen demand (11.4 mmol m⁻² d⁻¹; Urban et al., unpub.), the net reduction of Fe (0.9, Table 4), Mn (0.1, Table 4), $\rm SO_4^{2-}$ (0.14, Table 3) and

Table 4. Comparison of sedimentation, diffusive, and burial fluxes (mmol $m^{-2} \, d^{-1}$) at the deepest location in L. Sempach

Substance	Sedimentation ^a	Burial ^b	Sedimentation – Burial	Diffusive flux ^c
Inorg. C	12 ^d	18.2	- 6.2	
Org-C	23.5 e	9.6	13.8	
Total C	35.5	27.8	7.7	6.8 ^f
N	3.2 e	1.0	2.2	1.3
P	0.49e	0.21	0.28	0.08
Fe	0.90°	0.94	-0.04	0.34
Mn	1.60°	0.10	1.50	0.16

- ^a Taken from Urban et al. (1997).
- b Taken from Urban et al. (1997). Sediment accumulation rate of 1070 g m⁻² yr⁻¹ based on ²¹⁰Pb dating of Wieland et al. (1993).
- ^c Average based on all pore water profiles from 86-m depth (Table 4).
- ^d Based on Ca fluxes for periods 1988–1992 at 86-m depth.
- ^e Based on flux (1988–1992) at 86-m depth.
- ^f Sum of HCO₃, H₂CO₃, and CH₄ diffusive fluxes.

denitrification (1.9 mmol m⁻² d⁻¹; Table 3). If mineralization rates are highest at the deepest site of a lake (Carignan and Lean, 1991), then the discrepancy between consumption of electron acceptors and donors would be even larger for the entire hypolimnion. As indicated by the difference between fluxes of bicarbonate measured with benthic chambers and those calculated from porewater profiles (Table 3), the apparent electron imbalance results from a failure of the porewater profiles to indicate the large consumption of electron donors at the sediment surface.

A further indication that surface processes are not measured is that, as discussed below, the diffusive fluxes across the sediment surface failed to exhibit either the spatial or temporal trends expected. Spatial and temporal trends are expected to mirror trends in sedimentation of organic matter; the absence (or weakness) of such trends is taken as evidende that the porewater profiles fail to reflect all processes occurring in the sediments. Taken together these lines of evidence indicate that porewater profiles collected with peepers cannot be used to study processes at the sediment surface.

Although it must be regarded as a major limitation that porewater equilibrators cannot be used to measure surface phenomena, this result points to a second important conclusion of this study. If surface mineralization is not recorded by porewater profiles, then the calculated diffusive fluxes across the sediment surface must result from the slower decomposition that proceeds below the sediment surface. The porewater profiles themselves may be used to determine the locus of this decomposition. Because lake conditions have not changed over the past 2–3 years (the time required for porewater profiles to reach steady state to a depth of 20 cm), the first derivative of the concentration profiles is proportional to the diffusive flux, and the second derivative is proportional to the rate of production (or consumption) at each depth. Depth profiles of the second derivatives of porewater profiles of NH⁺₄

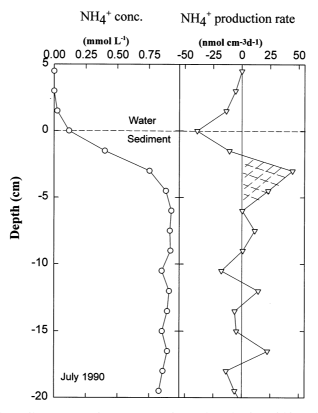


Figure 3. Depth profiles of ammonium concentration and production within sediments. From the second derivative of the concentration profile (left panel) in pore waters the production (or consumption) rate (right panel) of NH_4^+ is calculated. Shown here is the profile for July, 1990 at the deepest site (86-m) in L. Sempach. The production profile shows that mineralization occurred primarily in the upper 5 cm of sediments (hatched area); the negative rate at the sediment surface reflects both oxidation and enhanced transport above the sediments. Analysis of all NH_4^+ profiles yielded similar results

(Fig. 3) demonstrate that production of ammonium continues to a depth of 5–7 cm. A depth of 5–7 cm corresponds to a sediment age of 10–15 years. Evidently, decomposition of sediments deposited 10–15 years ago is still contributing to the diffusive fluxes of nutrients crossing the sediment surface. Carignan and Lean (1991) reported that release of solutes to pore waters continued to a depth greater than 50 cm in a mesotrophic lake. Our results indicate that sediments deposited under eutrophic lake conditions will continue to release nutrients for one to two decades after they reach the lake bottom.

The subsurface mineralization revealed by the peeper profiles is a major contributor to the total solute flux across the sediment surface (Table 5). Assuming that the fluxes into benthic chambers (Lander; Wehrli et al., 1997) represent the sum of surface and subsurface mineralization, the difference between Lander and peeper fluxes equals the surface mineralization rate. For N subsurface mineralization is

Table 5. Comparison of the magnitude of nutrient release from surface mineralization and from mineralization below the sediment surface

Nutrient	$Surface^a \\ mmol \ m^{-2} \ d^{-1}$	Subsurface ^b mmol m ⁻² d ⁻¹	Total % of deposition
carbon	19.0°	6.8 ^d	81 %
nitrogen	1.4°	1.3	83 %
phosphorus	0.22	0.08	61 %
silica	3.1	0.6	

- ^a Lander flux minus diffusive flux calculated from porewater profiles.
- ^b Diffusive flux calculated from porewater profiles.
- ^c Sum of HCO₃ and H₂CO₃.
- ^d Sum of HCO₃, H₂CO₃ and CH₄.
- e Ammonium flux into Lander may be underestimated due to nitrification within the chamber.

about equal to that at the surface. Desorption of P from Fe and Mn oxides at the sediment surface causes the Lander P flux to exceed P mineralization (Hupfer et al., 1995); similarly, it is incorrect to equate the diffusive fluxes of P calculated from the porewater profiles with mineralization of P because much of the mineralized P is trapped within the sediments. Nevertheless, subsurface release of P equals nearly 40% of that at the surface. Clearly, this slow mineralization of organic matter is quantitatively important and must be considered in efforts to decrease nutrient concentrations in lakes. Similarly, to predict the time required for lake recovery from eutrophic conditions, the kinetics of this slow decomposition must be determined.

Spatial variation

Systematic spatial variations in porewater concentrations and diffusive fluxes were expected as a result of sediment focusing. Higher wind energies delivered to littoral sediments cause resuspension of fine, organic-rich material that is focused to deeper areas of the lake where it accumulates (e.g., Davis, 1968; Bloesch, 1982; Bloesch and Uehlinger, 1986). Hence, inventories and net delivery of organic matter are higher at pelagic than littoral sites. That these processes operate in lakes Sempach and Baldegg is shown by the distribution of ²¹⁰Pb and organic matter in the sediments (Fig. 4A). Concentrations of organic N at the sediment surface increase with increasing water depth as a result of the increasing fluxes of organic matter caused by particle focusing (Fig. 4). The increase in deposition and accumulation of organic matter is greatest between 10- and 50-m depth. Both lakes are characterized by steep slopes and relatively flat central basins (mean depth ≅ one half of the maximum depth for both lakes). If rates of mineralization were proportional to the rate of delivery of organic matter, the diffusive fluxes of nutrients and C as well as of electron acceptors should increase with increasing water depth, particularly over the transition (10-50 m) between littoral and pelagic sediments. Increasing diffusive fluxes of nutrients and inorganic carbon with increasing water depth have been reported previously by Carignan and Lean (1991) for a mesotrophic lake, but

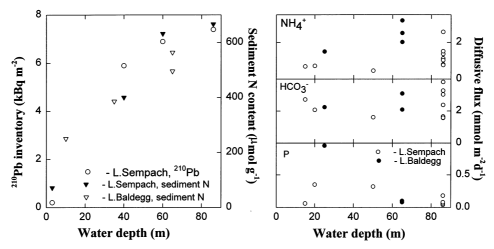


Figure 4. Spatial trends in organic matter deposition and mineralization. Concentrations of organic matter in surface sediments (Höhener, 1989) and sediment inventories of 210 Pb (Wieland et al., 1993) increase with increasing water depth as a result of focusing of fine particulate material. However, diffusive fluxes of NH $_4^+$, HCO $_3^-$ and P do not show parallel trends. Even when fluxes of CO $_2$ and CH $_4$ are added to the fluxes of HCO $_3^-$, no significant difference exists between littoral and pelagic areas. The increase in NH $_4^+$ fluxes with increasing water depth is small in comparison to the increase in organic matter in the sediments

Sherman et al. (1994) did not observe such a phenomenon in an oligotrophic lake. In L. Vechten oxygen consumption rates are higher in pelagic than in littoral sediments (Sweerts et al., 1991), but diffusive fluxes of sulfate and nitrate have been reported to be similar in littoral and pelagic sediments (Cook et al., 1986, Rudd et al., 1986; Urban et al., 1994).

In this study, systematic changes with water depth were observed in solute concentrations in porewater profiles. Littoral porewater profiles were characterized by lower concentrations of Mn²⁺, Fe²⁺, P, Si, NH₄ and alkalinity compared to pelagic profiles (Fig. 2). These differences were observed in both late summer and winter and in both lakes of this study. Lower concentrations in the dissolved phase are mirrored by lower concentrations of organic C, N, P, Fe and Mn in littoral compared to pelagic sediments of these lakes (Höhener, 1990; Urban, unpub.; Hupfer, unpub.). Low concentrations in littoral sediments of organic C, N and P are caused largely by focusing of fine particles as discussed above. Biogeochemical as well as physical processes contribute to the focusing of Fe and Mn and the depletion of these elements in littoral sediments (e.g., Davison et al., 1980; Hilton and Gibbs, 1984). The parallel between concentrations in solid and dissolved phases may indicate a kinetic rather than equilibrium control of porewater concentrations (e.g., Davison, 1991).

Despite the documented focusing of organic matter in both lakes and the higher solute concentrations in pelagic porewaters (Fig. 2), the diffusive fluxes across the sediment surface of carbonate species and Si showed little or no increase with increasing water depth (Table 2, Fig. 4). Diffusive fluxes of NH₄ across the sediment surface were approximately twofold higher in pelagic vs. littoral sites. The small or absent differences between littoral and pelagic sediments may result, in

part, from the failure of the porewater profiles to reflect the rapid mineralization at the sediment surface. However, the inventories and concentrations of organic matter below as well as at the sediment surface are higher in pelagic than in littoral sites. If production of mineralization products (inorganic carbon, NH₄⁺, P, Si) at littoral sites with small inputs and pools of organic matter equals production at pelagic sites with large inputs and pools, then the specific mineralization rate (i.e., rate per unit of organic matter) must be higher in the littoral sites. Our data would, therefore, support the hypothesis of Hilton and Gibbs (1984) that mineralization in littoral sediments is more rapid than in pelagic sediments. If mineralization is to be considered as first order with respect to organic matter (e.g., Berner, 1980), then the rate constants for littoral and pelagic sites must be different. Different rate constants are thought to result from the different oxygen regimes in littoral and pelagic sediments; greater oxygen availability in littoral areas enhances the activity of bacterial grazers (Lee, 1992) and the regeneration of other electron acceptors within the sediments and thereby accelerates sediment respiration.

Diffusive fluxes of NO₃, SO₄², and CH₄ across the sediment surface also did not show the expected increase with increasing water depth (Table 2). Three factors may contribute to this unexpected finding. First, given that the fluxes of inorganic carbon, Si, and P did not increase with increasing water depth, it is not surprising that fluxes of electron acceptors and fermentation products also did not increase. Second, as discussed below, profiles of NO₃ and SO₄² often show depletion above the sediment surface in response to variations in mixing intensity of the water column. Variations in water mixing intensities in epilimnion and hypolimnion may affect the fluxes of these ions and counteract the expected spatial trend. Third, internal cycling of electron acceptors could obviate any relation between net diffusive fluxes and gross rates of electron acceptor consumption. Rates of sulfate reduction are 20–100 times higher than rates of diffusion of SO₄- into sediment of L. Sempach as a result of sulfide oxidation within the sediments (Urban, 1994). Similar internal cycles exist for Mn (Thamdrup et al., 1994), Fe (Canfield et al., 1993), and N (Höhener and Gächter, 1994). The cycling of electron acceptors within the sediments or at the sediment-water interface causes the net fluxes of electron acceptors (except for O_2) to be lower than the total rates of reduction and could obscure the spatial pattern in total reduction rates.

The difference between diffusive fluxes of Fe²⁺ and Mn²⁺ in littoral and pelagic sites was highly significant (Table 2). Fluxes of Fe²⁺ were threefold higher at the deepest site compared to littoral sites, and fluxes of Mn²⁺ were 10-fold higher. This would not be surprising if the differences in fluxes could be attributed to different rates of reductive dissolution of Fe and Mn oxides; much higher rates are expected at the pelagic sites due to focusing of these substances (e.g., Davison et al., 1980). However, the calculated diffusive fluxes of Mn and possibly of Fe appear to be a result of dissolution of carbonate phases. No Mn oxides could be detected by EXAFS (Extended X-ray Absorption Fine Structure Spectroscopy) in the upper 2 mm of sediments from the deepest site in L. Sempach; EXAFS, scanning electron microscopy, microprobe analysis, and porewater saturation indices all indicated that Mn was present as a mixed Mn-Ca carbonate (Friedl et al., 1997). As mentioned above, the reductive dissolution of Mn oxides occurring at the sediment-water interface is not recorded in the porewater profiles. Similarly, sequential extractions

indicated that more than 90% of Fe in the surface sediments of L. Sempach was present as Fe(II) phases (Urban, unpub. data). As discussed below, the similarity in seasonal patterns of Fe²⁺ and Mn²⁺ diffusive fluxes as well as the similar depths of Fe and Mn dissolution within the sediments suggests that both elements are controlled by the same processes. It would appear that the large fluxes of Fe and Mn oxides to pelagic sediments (Wehrli et al., 1995) results in rapid reductive dissolution at the sediment-water interface, and some fraction of the reduced metals ($\sim 10\%$ for Mn, 90-100% for Fe; Table 4) are retained as carbonates, phosphates and sulfides (Friedl et al., 1997). Dissolution of these phases controls the porewater concentrations and fluxes measured in this study.

Temporal patterns in porewater profiles and diffusive fluxes

Seasonal variations in deposition of organic matter (Fig. 5) are an important driving force for temporal changes in porewater profiles (Fig. 6). Unlike in shallow lakes where temperature may be a driving variable (Rudd et al., 1990; Urban et al., 1994), temperature remains constant (5 °C) year-round in the pelagic sediments in the

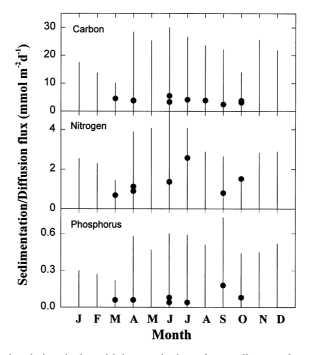


Figure 5. Seasonal variations in deposition to and release from sediments of nutrients. Deposition of organic C, N and P (bars) as measured with sediment traps above the deepest area of L. Sempach shows strong seasonal variations in response to seasonal variations in primary production. In contrast, diffusive fluxes (filled circles) of HCO $_3^+$ + CO $_2$, NH $_4^+$, and total P calculated from porewater profiles (86-m water depth, L. Sempach) show no seasonal variations. Diffusive fluxes represent all measurements at one site over a 3-yr period

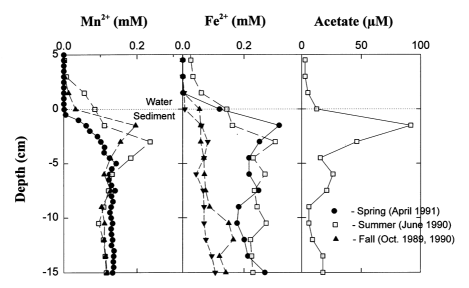


Figure 6. Seasonal variations in porewater concentrations of Fe^{2+} , Mn^{2+} , and acetate. All profiles were collected at 86-m water depth in L. Sempach. Transient peaks in concentrations of Fe^{2+} and Mn^{2+} occurring at depths of 1.5–3 cm are thought to result from seasonal dissolution of carbonate phases induced by enhanced rates of fermentation in summer. The solid triangles represent a profile of Fe^{2+} measured in October one year after the profile depicted by the open triangles

deep lakes of this study. However, increased deposition of organic matter in spring and summer months causes higher rates of organic matter hydrolysis and fermentation during these seasons. Rates of acetate production via fermentation are higher than rates of consumption in early summer, leading to development of a peak in acetate (as well as propionate and formate) concentrations within the first two centimeters of sediments (Fig. 6). The high rates of acetogenesis during this season imply high rates of CO₂ production. Increased concentrations of acetate and CO₂ cause a decrease in pH in the upper sediments. A decline in pH immediately below the sediment surface was observed in all porewater profiles; aerobic respiration is confined to the upper 5 mm of sediments and is unlikely contribute to the pH decline in the peeper profiles. The pH decline indicates that production of CO₂ via fermentation exceeded consumption of protons during anaerobic respiration.

The seasonal changes in profiles of Ca and Mg (Fig. 7) are believed to result from the seasonal change in pH. The pH decline below the sediment surface present in all months induces some dissolution of carbonate minerals. The dominant carbonate mineral in L. Sempach is low-magnesium calcite (Kelts and Hsue, 1979). Hence a slight increase in concentrations of Ca and Mg occurs just below the sediment surface in all months. This concentration gradient is considerably steeper (Fig. 7), and diffusive fluxes are higher in summer, presumably as a result of higher rates of fermentation. Enhanced carbonate dissolution in summer also might be related to the higher rates of calcite deposition during May through July (Fig. 7); however, given that surface mineralization of organic matter is not reflected in the porewater profiles, it is unlikely that dissolution of calcite at the sediment surface

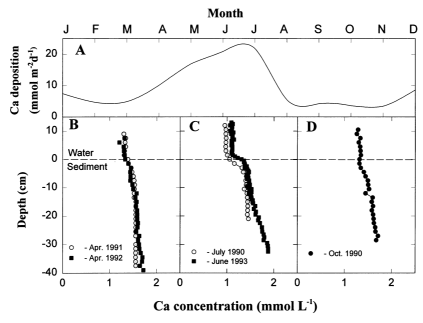


Figure 7. Seasonal deposition and dissolution of CaCO₃. Deposition (A) of CaCO₃ (measured with sediment traps at the center of L. Sempach) shows a marked seasonality with peak deposition in summer months. Porewater profiles of Ca (and Mg; not shown) from the deepest site in L. Sempach also show some seasonal features. Profiles are flatter in spring and fall, but show a marked kink at the sediment surface in summer that reflects higher rates of carbonate dissolution

is recorded either. Carbonate dissolution in these eutrophic lakes appears to be driven by fermentation as is observed in sediments of productive marine regions (e.g., Froehlich et al., 1979; Walter and Burton, 1990).

Seasonal changes in profiles of Mn and Fe also appear to be driven by the seasonal variations in fermentation. As mentioned above, both Fe and Mn are present primarily in reduced phases. Some dissolution or alteration of these phases evidently occurs in all months as evidenced by the increase in Mn concentrations within the upper 5 cm of sediments in all months (Fig. 6). Dissolution would appear to be enhanced in summer months leading to production of peaks in Mn concentrations at depths of 1.5 to 3 cm. In pelagic sediments of L. Sempach, Fe is divided nearly equally between sulfides and carbonates (Urban, unpub.). The similarity between seasonal patterns of Fe²⁺ and Mn²⁺ concentrations in pore waters suggests that the seasonal variations in Fe²⁺ also may be driven by dissolution of Fe carbonate phases (cf. Wersin et al., 1991). Some caution must be exercised in interpretation of profile shapes, however. First, although low concentrations of Fe²⁺ were observed twice in autumn, this could result from exposure of the peeper to oxygen (e.g., Carignan et al., 1994). Furthermore, seasonal changes in porewater profiles imply that the assumption of steady state may not be valid for all profiles. The peaks in Fe²⁺ and Mn²⁺ observed at 3-4 cm depth in the summer profiles may not reflect production of dissolved Fe²⁺ and Mn²⁺ at these depths at the time of sampling,

but rather the remnant of a larger peak that had existed previously at shallower depths.

The foregoing discussion is not intended to deny the existence of a seasonal cycle of Mn and Fe reduction. Such cycling does occur in lakes Sempach and Baldegg as evidenced by the seasonal variations in deposition of Fe and Mn oxides (Wehrli et al., 1995; Friedl et al., 1997) and the discrepancies between burial and deposition rates (Table 4). As discussed above, the fact that diffusive fluxes of Mn²⁺ across the sediment surface are much lower than sedimentation fluxes of Mn oxides provides one line of evidence that the measured porewater profiles fail to capture those processes occurring within millimeters of the sediment-water interface.

The failure of the porewater profiles to reflect surficial processes must be born in mind when examining the seasonal variations in diffusive fluxes of all components across the sediment surface. Despite the large seasonal variations in inputs of organic matter, no seasonal variations were observed in the diffusive fluxes of inorganic carbon, NH⁺₄, P, or Si (Fig. 5). The absence of seasonal variations together with the disrepancies between rates of input, output and accumulation (Table 4) constitute strong evidence that surface mineralization is not reflected in the porewater profiles. The calculated diffusive fluxes of nutrients across the sediment surface result from decomposition of organic matter deposited prior to the current year. There is little reason why this accumulated reservoir of organic matter should exhibit seasonal variations in rates of mineralization given that temperatures remain constant in the profundal sediments throughout the year.

The porewater profiles of NO_3^- and SO_4^{2-} do change seasonally (Fig. 8). In early spring and late fall, both NO_3^- and SO_4^{2-} appear to penetrate slightly deeper into the

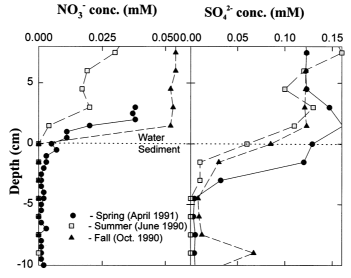


Figure 8. Seasonal variations in porewater profiles of NO_3^- and SO_4^{2-} . Depletion of both electron acceptors begins higher above (NO_3^-) or within (SO_4^{2-}) the sediments during summer months. All profiles shown were measured at 86-m water depth in L. Sempach

sediments than in summer months. In summer months, NO_3 depletion begins higher above the sediment surface than in spring or fall. Such seasonal variations have been reported for several other lakes (e.g., Rudd et al., 1990; Sinke et al., 1992; Sherman et al., 1994), and generally are attributed to increased microbial activity in summer months in response to higher inputs of organic matter. The variations also could indicate temporal variations in the intensity of mixing of the water immediately above the sediments.

Curiously, the diffusive fluxes across the sediment surface of NO_3^- and SO_4^{2-} do not show any seasonal variations (Fig. 9). Sherman et al. (1994) also did not observe any seasonal variations in diffusive fluxes of SO_4^{2-} or NO_3^- in an intensive study of Little Rock Lake. Although measured porewater profiles underestimate the concentration gradients of these ions at the sediment surface, it also is clear that the diffusive fluxes of both of these species underestimate the gross reduction rates that

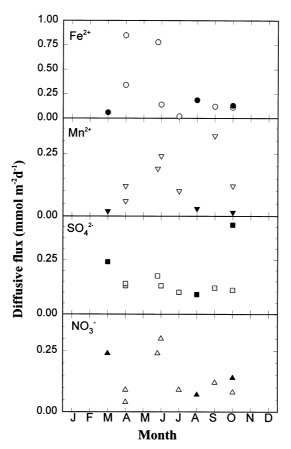


Figure 9. Seasonal variations in diffusive fluxes of electron acceptors. The points represent all measurements made in L. Sempach over a 4-year period (9/89–6/93). No clear seasonal trends are apparent. (Filled symbols indicate measurements from littoral sites; all other points were measured at the deepest location.)

occur within the sediments. Rates of sulfate reduction in L. Sempach 100-fold greater than rates of SO_4^{2-} diffusion into the sediments are made possible by high rates of sulfide oxidation (sulfate regeneration in the mat of Beggiotoa on the sediment surface). Similarly, Sweerts et al. (1991) reported that 27-36% of the NH_4^+ flux out of the sediments of L. Vechten was oxidized to NO_3^- at the sediment surface. The concentrations of both SO_4^{2-} and NO_3^- in sediment pore waters and immediately above the sediment surface must be regarded as representing the balance between rates of reduction and regeneration (i.e., oxidation or reduced S and N). The concentration profiles are not, therefore, a satisfactory means for assessing the dynamics of microbial processes of sulfate reduction and denitrification (cf. Lovley and Klug, 1986; Brandl and Hanselmann, 1991).

Summary

Both methodological lessons as well as insights into biogeochemical dynamics of lake sediments may be drawn from this study. In L. Sempach the small-scale spatial variability of the porewater was small in a limited set of replicate profiles. Variability in diffusive fluxes across the sediment surface is probably element-specific, but averaged 35%. This study clearly pointed out that porewater profiles with resolution of 0.5 to 1.5 cm do not record the processes occurring at the sediment surface. In both of the eutrophic lakes of this study, mineralization at the surface and mineralization below 0.5 cm depth contributed significantly to the recycling of nutrients to the water column. Consequently, any changes in nutrient deposition will require more than a decade to have their full effect; that is, the sediment's memory of previous conditions will require some time to be erased.

Systematic changes in porewater chemistry and diffusive fluxes across the sediment surface occurred with increasing water depth. Focusing of Fe and Mn oxides led to large differenes between fluxes of these metals from littoral and pelagic regions. Because the porewater profiles did not record surface mineralization, there were not systematic, spatial differences in the diffusive fluxes across the sediment surface of nutrients or inorganic carbon. Seasonal variations in diffusive fluxes across the sediment surface were minor for most solutes despite some marked changes in profile shapes. However, the seasonal variations in seston deposition did drive seasonal variations in rates of fermentation. These seasonal variations in microbial activity in turn caused seasonal variations in mineral dissolution as revealed in the porewater profiles of Ca²⁺, Mg²⁺, Mn²⁺, and Fe²⁺. The small or absent seasonal changes in diffusive fluxes of Fe²⁺ and SO₄²⁻ despite changes in rates of reduction of both substances point to the fact that porewater profiles reflect only net reactions. The gross rates of opposing reactions can only be calculated from porewater profiles if the opposing reactions are spatially distinct in the profile. Whereas this is seldom the case in eutrophic lake sediments, porewater profiles generally will underestimate gross rates of microbial activity.

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