Nitrate source identification using its isotopic ratios in combination with a Bayesian isotope mixing model in the Baltic Sea

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13 Abstract

14 Nitrate (NO_3) is the major nutrient responsible for coastal eutrophication worldwide and its production is related to intensive food production and fossil-fuel combustion. In the Baltic 15 16 Sea NO₃⁻ inputs have increased four-fold over the last decades and now remain constantly high. NO₃⁻ source identification is therefore an important consideration in environmental 17 18 management strategies. In this study focusing on the Baltic Sea, we used a method to estimate the proportional contributions of NO3⁻ from atmospheric deposition, N2 fixation, and runoff 19 from pristine soils as well as from agricultural land. Our approach combines data on the dual 20 isotopes of NO₃⁻ (δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻) in winter surface waters with a Bayesian isotope 21 mixing model (Stable Isotope Analysis in R, SIAR). Based on data gathered from 47 22 23 sampling locations over the entire Baltic Sea, the majority of the NO₃⁻ in the southern Baltic was shown to derive from runoff from agricultural land (33-100%), whereas in the northern 24 Baltic, i.e., the Gulf of Bothnia, NO₃ originates from nitrification in pristine soils (34–100%). 25 Atmospheric deposition accounts for only a small percentage of NO₃⁻ levels in the Baltic Sea, 26 except for contributions from northern rivers, where the levels of atmospheric NO_3^- are 27 higher. An additional important source in the central Baltic Sea is N₂ fixation by diazotrophs, 28 which contributes 49-65% of the overall NO₃ pool at this site. The results obtained with this 29 method are in good agreement with source estimates based upon δ^{15} N values in sediments and 30

a three-dimensional ecosystem model, ERGOM. We suggest that this approach can be easily
modified to determine NO₃⁻ sources in other marginal seas or larger near-coastal areas where
NO₃⁻ is abundant in winter surface waters when fractionation processes are minor.

4

5 1 Introduction

6 Throughout the world, anthropogenic reactive N currently exceeds natural production 7 (Galloway et al., 2003; Gruber and Galloway, 2008). Consequently, riverine nitrogen (N) 8 fluxes have doubled in recent years, which has strongly impacted the marine N cycle and 9 ecosystem health, both at regional and global scales. In coastal ecosystems, the adverse 10 effects of these excess N loads include eutrophication, hypoxia, loss of biodiversity, and habitat destruction (Galloway et al., 2003; Villnäs et al., 2013). For the shallow, brackish, 11 12 semi-enclosed Baltic Sea, where intense anthropogenic nutrient loadings have been documented since the 1950s (Elmgren, 2001), riverine and atmospheric nutrient inputs are 13 now at least four-fold higher than a century ago, when anthropogenic influence was low 14 (Schernewski and Neumann, 2005; Stålnacke et al., 1999). Furthermore, cyanobacterial 15 blooms, which can fix N₂, and thus add nutrients to the surface waters are regular large scale 16 phenomenon each summer (Finni et al., 2001; Vahtera et al., 2007) and the overall increase in 17 nutrient input has supported the expansion of hypoxic zones (Conley et al., 2009, 2011). 18

A main component of the N pool and the one most readily available is nitrate (NO_3) (Nestler 19 et al., 2011; Vitousek et al., 1997), which derives from a wide variety of sources. These can 20 be identified by analysis of the N and oxygen (O) isotopes (δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻) since 21 the isotopic ratios of NO₃⁻ from different sources fall within distinct ranges (Kendall, 1998; 22 Kendall et al., 2007). For example, NO_3^- inputs from forested catchments can be 23 discriminated from those coming from agricultural runoff, and the NO₃ signature of microbial 24 nitrification differs from that of atmospheric deposition (Kendall, 1998; Kendall et al., 2007; 25 Mayer et al., 2002). Source attribution is, however, complicated by N-transformation 26 27 processes such as denitrification, nitrification, and assimilation, each of which gives rise to significant isotope fractionation. Since heavier isotopes are sequestered more slowly than 28 lighter ones, the reaction product will be isotopically depleted compared to the original NO₃⁻ 29 source (Kendall, 1998). Alterations of isotope values because of microbial fractionation 30 processes can be minimized by collecting the samples in winter, when low water temperatures 31 32 reduce microbial activity (Pfenning and McMahon, 1997).

Nonetheless, source attribution is still complicated when there are more than three sources but 1 only two isotopes that describe them (Fry, 2013). SIAR (Stable Isotope Analysis in R), a 2 Bayesian isotope mixing model originally developed to infer diet composition from the stable 3 isotope analysis of samples taken from consumers and their food sources (Moore and 4 Semmens, 2008), was already successfully applied for NO_3^- source identification. Xue et al. 5 (2012, 2013) were able to estimate the proportional contributions of five potential NO_3^{-1} 6 7 sources in a small watershed in Flanders (Belgium). Based on their determinations of the isotopes of nitrogen and oxygen they could show that manure and sewage were the major 8 9 sources of NO₃⁻.

10 In the Baltic Sea the NO_3^- pool present in the surface waters in spring originates from the previous growth season and is consumed during the onset of the phytoplankton spring bloom, 11 12 in February/March. Stratification in summer hinders circulation down to the halocline, thus atmospheric deposition and N₂ fixation are the major N sources, whereas in coastal areas 13 riverine discharge dominates (Radtke et al., 2012; Voss et al., 2011). Yet, to what extent the 14 various NO₃⁻ sources add to the overall pool of NO₃⁻ in the Baltic as a whole is still a matter 15 of debate. In this study, a source attribution for four major sources is presented. Taking the 16 Baltic Sea as an example we will show, that the use of the isotopic composition of NO_3^{-1} 17 $(\delta^{15}$ N-NO₃⁻ and δ^{18} O-NO₃⁻) in combination with SIAR can be used elsewhere for source 18 identification on an ecosystem scale level. 19

20

21 2 Material and Methods

22 2.1 Field sampling

Surface water samples from the Baltic Sea were collected in February 2008 (n=22) and 2009 23 24 (n=17) before the onset of the phytoplankton spring bloom aboard the RV Alkor and in November 2011 (n=1) aboard the RV Meteor using a Seabird CTD system with attached 25 water bottles. Samples from the Nemunas River (55°18`5.5 N, 21°22`53.9 E; 55°41`25.6 N, 26 21°7`58.4 E; n=4) and Kalix River (65°56`4.2 N, 22°53`9.2 E; n=1) (Fig.1) were taken 27 between November 2009 and February 2010. Values for NO₃⁻ in which atmospheric 28 deposition was the source were obtained from wet deposition samples collected at three 29 stations around the Baltic Sea: Warnemünde, Germany (54°10' N, 12°5' E,); Majstre, Sweden 30 (57°30' N, 18°31' E); and Sännen, Sweden (56°13'N, 15°17'E) from December 2009 until 31 February 2010 (Table 1). In Warnemünde, precipitation was collected on an event basis, and 32 retrieved daily to limit microbial degradation, using a sampler consisting of a plastic funnel 33

(diameter: 24 cm) connected to a 1-L polyethylene bottle. At the two Swedish stations,
rainwater was sampled monthly by the Swedish Environmental Research Institute (IVL) as
part of the Swedish national long-term monitoring program. Here, the sampler consisted of a
plastic funnel (diameter 20.3 cm) connected to an 8-L polyethylene bag. All samples were
filtered through pre-combusted Whatman GF/F filters (4 h at 400°C) and stored frozen until
further analysis.

7

8 2.2 Nutrient concentrations and dual isotope analysis of NO₃⁻

Samples were analyzed following a standard protocol for the determination of NO₃⁻ and nitrite 9 (NO_2) (Grasshoff et al., 1983); the precision of the method is $\pm 0.02 \text{ }\mu\text{mol} \text{ }1^{-1}$. Dual isotope 10 analysis of NO₃⁻ (δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻) was carried out using the denitrifier method 11 (Casciotti et al., 2002; Sigman et al., 2001), in which NO₃⁻ and NO₂⁻ are quantitatively 12 converted to nitrous oxide (N₂O) by *Pseudomonas aureofaciens* (ATTC 13985), a bacterial 13 strain that lacks N₂O reductase activity. In brief, N₂O is removed from the sample vials by 14 purging with helium and then concentrated and purified in a GasBench II prior to analysis 15 with a Delta Plus mass spectrometer (ThermoFinnigan). NO_2^- was not removed since its 16 concentrations were always less than 2% (referring to the procedure described in Casciotti et 17 al., 2007). N and O isotope measurements of roughly 30% of the samples were replicated in 18 separate batch analyses. Two international standards, IAEA-N3 ($\delta^{15}N=4.7\%$ vs. N₂; $\delta^{18}O$ 19 25.6‰ vs. VSMOW) and USGS 34 (δ^{15} N -1.8‰ vs. N₂; δ^{18} O -27.9‰ vs. VSMOW) (Böhlke 20 et al., 2003), were measured with each batch of samples. Samples with NO_3^{-7}/NO_2^{-1} 21 concentrations as low as 1 μ mol l⁻¹were analyzed. The sample size for the actual stable 22 isotope measurements was 20 nmol for samples with concentrations >3.5 $\mu mol~l^{-1}$ and 10 23 nmol for those with concentrations $<3.5 \mu$ mol 1^{-1} . Isotope values were corrected after Sigman 24 et al. (2009) for δ^{18} O-NO₃; single point correction was referred to IAEA-N3 for δ^{15} N-NO₃. 25 The precision was <0.2 ‰ for δ^{15} N and <0.6 ‰ for δ^{18} O. Together with the samples, a culture 26 blank was analyzed to which no sample was added. The isotope ratios are reported using the 27 delta notation in units of per mil (%). 28

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30 2.3 NO₃ sources

To estimate the contribution of different NO₃⁻ sources, two isotopes δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻ (j=2) from the four major NO₃⁻ sources: (1) atmospheric deposition, (2) runoff from pristine soils, (3) runoff from agricultural land and (4) N₂ fixation were applied (Table 2). In this

context, N₂ fixation was defined as NO₃⁻ originating from the degradation and remineralization 1 of nitrogen fixers and therefore carried their low isotopic signal. Thus, for NO_3^- from N_2 2 fixation, δ^{15} N values of ~-2 to 0‰ were assumed, since N₂ fixation produces organic material 3 that is only slightly N depleted against air nitrogen (Carpenter et al., 1999, 1997; Montoya et 4 al., 2002). The δ^{18} O values were estimated to be between -3.8‰ and 2.0‰, based on 5 measurements in the subtropical northeast Atlantic where N₂ fixation was the main source of 6 N (Bourbonnais et al., 2009) (δ^{18} O-NO₃⁻ = 2‰) and the estimated δ^{18} O of NO₃⁻ deriving from 7 N_2 fixation by Sigman et al. (2009) (δ^{18} O-NO₃⁻ = -0.2‰) and Bourbonnais et al. (2012) (δ^{18} O-8 $NO_3^- = -3.8\%$). 9

10 To expand the dataset, we included NO_3^- isotope data from river water samples, ground water samples, and samples from tile drain outlets collected in 2003 and published in Deutsch et al. 11 12 (2006). In that study, the Warnow River (n=2) was sampled twice, in January and February 2003. These sources were likewise sampled in winter, since marked seasonal shifts in the 13 isotopic composition of NO_3^- can occur due to shifts in the origins of the sources (Knapp et 14 al., 2005). Samples from tile drain outlets were used to represent NO₃⁻ from agricultural 15 runoff and were obtained from the catchment of the Warnow River, whose waters are strongly 16 influenced by agricultural land use (Pagenkopf, 2001). High δ^{15} N-NO₃⁻ values of 9.9±1.5‰ 17 and lower δ^{18} O-NO₃⁻ values of 4.6±1.0‰ are typical for areas that are influenced by 18 agricultural activities and are comparable to studies of Wankel et al. (2006) and Johannsen et 19 20 al. (2008). Johannsen et al. (2008) found in the rivers Rhine, Elbe, Weser and Ems, with comparable high agricultural activities, δ^{15} N-NO₃⁻ values between 8.2 and 11.2‰ and δ^{18} O-21 NO₃⁻ values from 0.4 to 0.9‰ in winter. However, a differentiation between NO₃⁻ from 22 mineral fertilizers and sewage/manure was not done; rather a mixed signal from rivers that are 23 mainly influenced by agricultural activities was taken. Groundwater samples were used as the 24 source of NO₃⁻ from pristine land (Deutsch et al., 2006). Their δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻ 25 values significantly differed from those of agricultural runoff (p<0.05) but were similar to the 26 values of other areas, such as Biscuit Brook (Burns et al., 2009) and the San River (Koszelnik 27 and Gruca-Rokosz, 2013), where pristine soils were sampled and reflect nitrification activity 28 in soils unaffected by human activity. 29

30 The dual isotopes of NO_3 values presented in Deutsch et al. (2006) were analyzed according

- to Silva et al., (2000). In this method, NO_3^- is chemically converted via anion exchange resins
- 32 to AgNO₃⁻ and the δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻ values are measured via pyrolysis and isotopic
- 33 ratio mass spectrometry (for a detailed description, see Deutsch et al. 2006). A normal

distribution of the isotopic data from the four sources was confirmed by applying the ShapiroWilk normality test. δ¹⁵N-NO₃⁻ and δ¹⁸O-NO₃⁻ values from NO₃⁻ from atmospheric deposition
of 0.3±1.4‰ and 76.7±6.8‰, respectively, are also in line with literature values. The δ¹⁵N
values of atmospheric NO₃⁻ are usually between -15 to +15‰ and the δ¹⁸O between 63 and

5 94‰ (Kendall et al., 2007).

Six regions within the catchment of the Baltic Sea were investigated for their potential NO₃⁻ 6 7 sources (Fig. 1). According to the topography of the Baltic Sea, the samples were assigned to four major areas: Western Baltic Sea, Baltic Proper, Gulf of Finland, and Gulf of Bothnia. 8 Additionally, three rivers differing in their degree of anthropogenic impact were included in 9 10 this study and divided into two groups: northern and southern rivers. Rivers with high nutrient loads drain mainly into the southern Baltic Proper and were represented here by the Nemunas 11 and Warnow Rivers, whose NO_3^- concentrations in winter can be as high as 260 µmol l^{-1} 12 (Deutsch et al, 2006, Pilkaityte and Razinkovas 2006). The Gulf of Bothnia receives large 13 amounts of fresh water from rivers represented by the Kalix River. These rivers drain mainly 14 pristine, forested land and have maximum NO_3^- concentrations of around 20 µmol l^{-1} 15 (Sferratore et al, 2008). 16

17

18 2.4 SIAR mixing model

19 The applied mixing model is described by the following equations:

20
$$X_{ij} = \sum_{k=1} p_k (s_{jk} + c_{jk}) + \varepsilon_{ij}$$
 (1)

21
$$s_{jk} \sim N(\mu_{jk}, \omega_{jk}^2)$$
 (2)

22
$$c_{jk} \sim N(\lambda_{jk}, \tau_{jk}^2)$$
 (3)

23
$$\varepsilon_{ij} \sim N(0, \sigma_1^2)$$
 (4)

24 where X_{ij} is the observed isotope value j of the mixture i; i=1,2,3,...,I are individual observations; and j=1,2,3,...,J are isotopes. s_{ik} is the source value k of isotope j (k = 1,2,3,..., 25 26 K) and is normally distributed, with a mean of μ_{ik} and a standard deviation of ω_{ik} . p_k is the proportion of source k that needs to be estimated by the model. c_{ik} is the fractionation factor 27 for isotope j on source k and is normally distributed, with a mean of λ_{ik} and a standard 28 deviation of τ_{ik} . ε_{ii} is the residual error representing additional unquantified variations between 29 30 mixtures and is normally distributed, with a mean of 0 and a standard deviation of σ_i . Detailed descriptions of the model can be found in Jackson et al. (2009), Moore and Semmens (2008), 31 and Parnell et al. (2010). As noted above, by collecting samples between November and 32

February we minimized the influence of fractionation processes such as assimilation and denitrification that can alter the isotopic signal of NO_3^- . Therefore in Eq. (1) we assumed that $c_{jk} = 0$.

4 Two different runs of the SIAR model were performed. In the first, for the Western Baltic 5 Sea, Baltic Proper, and Gulf of Finland, all four sources were included in the calculation. In 6 the second, for the Gulf of Bothnia, the southern rivers, and the northern rivers, N_2 fixation as 7 a potential NO_3^- source was excluded since in these areas there is no N_2 fixation by 8 diazotrophs because the Gulf of Bothnia is phosphorus limited, in contrast to the Baltic Proper 9 (Graneli et al., 1990).

10

11 3 Results

12 3.1 NO₃⁻ concentrations and isotopes

Winter (Nov.-Feb.) surface NO_3^- concentrations ranged from a minimum of 2.6 μ mol l^{-1} in the 13 open Baltic Sea to a maximum of 259 μ mol l⁻¹ close to the estuaries of the most nutrient-rich 14 rivers, i.e., the Nemunas and Warnow Rivers (Fig. 2, Supplement 1). In most basins of the 15 Baltic Sea, the NO₃⁻ concentrations in winter were almost identical, with the exception of the 16 Gulf of Finland, where concentrations were about two-fold higher (7.6 \pm 0.9 µmol l⁻¹; Fig. 2). 17 In the western Baltic Sea, the Baltic Proper, and the Gulf of Bothnia NO₃⁻ concentrations were 18 similar with 3.3 \pm 0.6, 3.4 \pm 0.8, and 3.7 \pm 0.4 µmol 1⁻¹, respectively. 19 Highest nitrate concentrations in the Nemunas River also corresponded to the highest δ^{15} N-20

NO₃⁻ with 10.0 ‰ and vice versa, with lowest concentrations and nitrogen isotope values in the Baltic Proper (1.5 ‰). The δ^{18} O-NO₃⁻ values ranged from -2.8 ‰ in the Gulf of Bothnia to 10.6 ‰ in the Northern River, Kalix (Fig. 2, Supplement 1).

24

25 3.2 Sources of NO₃

SIAR calculated that in the southern Baltic Sea, agricultural runoff was the main NO₃⁻ source 26 with the highest contribution in the western Baltic Sea with up to 67 % (mean $53.5\pm3.2\%$) and 27 in the southern rivers with up to 100% (mean 93.5±4.1%) (Table 3, Fig. 1). NO₃⁻ from 28 atmospheric deposition was negligible with 3.5% (mean $1.1\pm0.5\%$) and NO₃⁻ from pristine 29 soils lower with up to 42% (mean 7.5±5.9%) in the western Baltic Sea (Table 3, Fig. 1). In the 30 Baltic Proper, NO_3^{-1} from N_2 fixation was the dominant NO_3^{-1} source with up to 65.3% (mean 31 58.8 \pm 2.0%) (Table 3, Fig. 1). In the northern Baltic Sea NO₃⁻ from atmospheric deposition is 32 only important in the northern rivers with a contribution of up to 23.4% (mean $11.8\pm1.5\%$) 33

(Table 3, Fig. 1). NO₃⁻ from pristine soils is mainly transported by the northern rivers
(75.3±7.9%) to the Gulf of Bothnia where SIAR calculated that 99.0±0.9% stems from the
runoff from pristine soils (Table 3, Fig. 1).

4

5 **3.3** Comparison of isotope patterns in the water column and sediments

The $\delta^{15}N$ values from surface water correlated significantly with those from surface 6 sediments, as reported in Voss et al. (2005) (p<0.001) (Fig. 3). Stations for sediment sampling 7 were in close vicinity to stations from water column sampling (Fig 4). In the Baltic Proper, 8 the $\delta^{15}N$ of the surface water NO₃⁻ was indistinguishable from the $\delta^{15}N$ of the sediment 9 surface (3.6±1.0‰ and 3.5±0.6‰, respectively; Table 4). In the near-coastal areas of the 10 Baltic Proper and the Gulf of Finland, the δ^{15} N of surface water NO₃⁻ was 7.9±1.8‰, slightly 11 higher than the surface sediment value for the same area of $7.3\pm2.1\%$ (data in Voss et al., 12 2005) but still not significant different (p<0.01) (Table 4). 13

14

15 4 Discussion

16 4.1 NO₃ in the Baltic Sea

The measured winter surface water concentrations of up to 259 µmol l⁻¹ are typical for 17 eutrophied systems and similar values have been reported from the Chesapeake Bay and the 18 coastal areas of the North Sea (Dähnke et al., 2010; Francis et al., 2013). The concentrations 19 of nutrients in the sub-basins of the Baltic Sea reflect the densities of the human populations 20 in the vicinity of the adjacent sub-catchments. Thus, in the near-coastal area of the southern 21 22 Baltic Proper, NO_3^{-1} concentrations were higher than in the northern parts, since the catchment areas of Germany, Poland, and the Baltic States are much more densely populated (>500 23 inhabitants km⁻²) and the land is intensively used for agricultural purposes. The northern 24 regions are dominated by boreal forests and less populated (<10 inhabitants km⁻²) (Lääne et 25 al., 2005; Stepanauskas et al., 2002; Voss et al., 2011). Consequently, for the southern Baltic 26 Proper a relationship between fluvial NO₃⁻ loads and NO₃⁻ concentrations in coastal waters 27 could be established that indicates a direct impact of riverine nutrients on coastal waters (Voss 28 et al. 2011, HELCOM, 2009). However, there was no similar correlation between riverine N 29 30 loads and nutrient concentrations either for the coastal areas of the Gulf of Bothnia or for the open waters of the Baltic Proper (Voss et al. 2011). The Gulf of Bothnia is the only sub-basin 31 in which the effects of eutrophication are so far minor, although Lundberg et al. (2009) and 32 Conley et al. (2011) reported a degradation in the water quality from north to south and from 33

the outer to the inner coastal area of the Gulf, with seasonal hypoxia at many sites. Trends of increasing nutrient levels should be interpreted as a warning signal for the future and highlight the need for management approaches based on sound knowledge of the many potential sources of NO₃⁻.

5 In the Gulf of Finland, which is regarded as the most heavily eutrophied sub-basin of the 6 Baltic Sea, a consequence of high receiving nutrient loads from the Neva River and the city of 7 St. Petersburg (Lundberg et al. 2005), NO_3^- concentrations were about two-fold higher 8 (7.6±0.9 µmol Γ^-) compared to the rest of the Baltic Sea sub-basins, where concentrations in 9 winter were almost identical. This shows that NO_3^- concentrations alone cannot be used to 10 identify NO_3^- sources for the sub-basins; rather, stable NO_3^- isotopes values allow for accurate 11 source determination, as we will show in the following sections.

12

13 4.2 Sources of NO₃

The use of NO_3^- stable isotopes for source identification is complicated when the mixing of 14 multiple N sources with overlapping isotopic ranges occurs together with microbial processes 15 such as nitrification, assimilation, and denitrification (Kendall, 1998; Wankel et al., 2006). In 16 this study, we assumed that the effects of fractionation by microbial processes were negligible 17 because all our samples were collected in winter, at a mean temperature of 3.1±1.3 °C (data 18 not shown), when microbial activity is low (Pfenning and McMahon, 1997), as confirmed in a 19 20 study of nitrification in the Baltic Sea by Jäntti et al. (2011). They showed that in the Gulf of Finland although nitrification potentials may be high during cold months, in situ nitrification 21 22 is undetectable whereas the rate increases progressively towards the summer.

We are aware that the variability of the source signals must be taken into account in source 23 24 attributions. Both Xue et al. (2012, 2013) and Yang et al. (2013) showed that SIAR can be applied in NO_3^- source identification, although the resolution of this model is largely 25 determined by the uncertainty of the isotopic composition of the sources. In the studies of 26 both groups, the means and variances of the sources were calculated mostly from literature 27 28 values, which were not obtained in the investigation areas, nevertheless they received consistent results. In contrast, in our study, the isotopic composition of the sources, except 29 NO3⁻ from N2 fixation, was determined from samples obtained within the study area. In our 30 calculations we considered the impact of the variability of the sources and report not only 31 mean values and error estimates but also minimum and maximum contributions, as suggested 32 by Fry (2013) (Table 3). 33

2 4.2.1 NO₃⁻ from agricultural runoff

The isotopic values of riverine NO_3^- were previously shown to be enriched when agricultural 3 land is the source of inputs (Johannsen et al., 2008; Mayer et al., 2002; Voss et al., 2006). 4 Catchments with high percentages of agricultural and/or urban land use export NO₃⁻ with 5 δ^{15} N-NO₃ values of around 7‰. In the same study, the oxygen isotope ratios of NO₃ were 6 almost uniformly 13±1‰ (Mayer et al. 2002). Johannsen et al. (2008) measured δ^{15} N-NO₃⁻ 7 values of 11.3‰ in highly eutrophied rivers draining into the North Sea, whereas the highest 8 δ^{18} O-NO₃ value was 2.2‰. In the Oder River outflow, a main NO₃ contributor to the Baltic 9 Sea, δ^{15} N-NO₃ of 7.6‰ and δ^{18} O-NO₃ of 2.9‰ were determined (Korth et al., 2013). Our 10 measurements for the Warnow and Nemunas Rivers fall in the expected range, with a mean 11 δ^{15} N-NO₃ of 9.2‰ and a mean δ^{18} O-NO₃ of 3.1‰, and are consistent with the high 12 percentages of agricultural land in the river catchment areas: 50% for the Warnow River 13 14 (Pagenkopf, 2001) and 50% for the Nemunas River (Christoph Humborg, personal communication, 2011). For both, SIAR calculations indicated that 75.2-100% (mean 15 16 93.5 \pm 4.2%) of the NO₃⁻ pool is from agricultural runoff. NO₃⁻ with this signature seems to be transported to the central Baltic Sea, since SIAR-based estimates showed significant 17 18 percentages of agriculturally derived NO_3^- in the Western Baltic Sea (41.0–66.5%; mean: 19 53.5±3.2%), the Baltic Proper (32.8–45.5%; mean: 39.0±1.6%), and the Gulf of Finland $(40.9-63.4\%; \text{ mean: } 51.9\pm3.0\%)$. However, high percentages were only expected for the Gulf 20 of Finland and the Western Baltic Sea, where large N loads from agricultural land have been 21 documented (Hong et al., 2012). Indeed, for the Baltic Proper, the sizeable contribution of 22 agricultural NO_3^- (39.0±1.6%) was surprising and contrasted with previous findings that 23 nearly excluded riverine NO_3^- as a major nutrient source for the central Baltic Sea (Voss et al., 24 25 2005, 2011). However, Neuman (2000) estimated that 13% of the N input of the Oder River is 26 transported to the central Baltic Sea, while Radtke et al. (2012) could show, using a source 27 attribution technique in the three-dimensional ecosystem model ERGOM (Ecological 28 ReGional Ocean Model), that at least a part of the dissolved inorganic nitrogen (DIN) load 29 from the Vistula River, the main NO_3^- contributor to the Baltic Sea (Wulff et al., 2009), enters the Baltic Proper. This 3D model comprises a circulation model, a thermodynamic ice model, 30 31 and a biogeochemical model and utilizes the Modular Ocean Model, MOM3.1 (Radtke et al., 32 2012).

Another explanation for the high estimated agricultural influence in our study could be the 1 intrusion of water containing NO_3^- with similar NO_3^- isotope values as our agricultural NO_3^- 2 source during mixing/advection from below the halocline. Deep-water NO₃⁻ in the Baltic Sea 3 has a δ^{15} N of about 7‰ (Frey et al. unpubl. data), which is higher than the average deep-water 4 ocean NO₃⁻ signature of 5‰ (Sigman et al., 2000). This elevated δ^{15} N in NO₃⁻ mainly comes 5 from water column denitrification in the oxic-anoxic interface in water at a depth of about 100 6 7 m (Dalsgaard et al., 2013). However, the year-to-year variations in DIN due to vertical mixing and advection from below the halocline are sensitive to hydrographic conditions. When the 8 9 halocline is weak and well ventilated, oxygen conditions improve, resulting in higher DIN concentrations in deep waters and greater advection and/or mixing (Vahtera et al, 2007) such 10 that the NO_3^- contribution from below the halocline is difficult to estimate. 11

Overall, the range of 32.8-45.5% (mean: $39.0\pm1.6\%$) determined for NO₃⁻ presumably originating from agricultural runoff has to be considered with caution, because the former imprint of deep water column denitrification and mixing/advection of this isotopically enriched NO₃⁻ from below the halocline with the residual winter surface NO₃⁻ pool could have resulted in an overestimation of the percentage of NO₃⁻ from agricultural runoff in the Baltic Proper.

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19 **4.2.2** NO₃ from N₂ fixation

The average δ^{15} N-NO₃⁻ value of 3.6‰ for the Baltic Proper is significantly lower than the ocean average of around 5% (Sigman et al., 2000) and presumably reflects the influence of N₂ fixation. This is because the δ^{15} N of newly fixed N is between -2 and 0‰ such that NO₃⁻ has slightly lower δ^{15} N values (Knapp et al., 2005; Liu et al., 1996). The δ^{18} O-NO₃⁻ value of -0.5‰ in the Baltic Proper is also slightly lower than the ocean average of 1.5‰, and close to our theoretical considered value of -0.7±2.9‰ after the degradation and remineralization of N₂ fixers.

N₂ fixers are abundant in summer, reflecting the stimulation of their growth by the low N/P ratios. N in the cyanobacterial biomass is remineralized over the winter months and the resulting NO₃⁻ remains in the water masses down to the halocline. Our results show that the contribution of N₂ fixation by diazotrophs to the NO₃⁻ pool is 49.3–65.3% (mean 58.8±2.0%). This is slightly higher compared to the data reported by Wasmund et al. (2001), who estimated that 39% (370 ktons yr⁻¹) of a total input of 955 ktons N yr⁻¹ (HELCOM, 2002) stems from N₂ fixations in the central Baltic Sea. Both Radtke et al. (2012) and Voss et al.

(2005) concluded that N₂ fixation was the main NO₃⁻ source in the Baltic Proper. Using an 1 independent approach, we were able to confirm the contribution of N₂ fixation in this area. In 2 addition, we found that N₂ fixation is also a major source of NO₃⁻ in the Western Baltic Sea 3 and the Gulf of Finland (respectively, 11.0-51.9% (mean 37.9±5.1%) and 32.7-59.0% (mean 4 5 $45.5\pm3.2\%$)). This finding is consistent with our current understanding of N₂ fixation in the Gulf of Finland (Vahtera et al, 2005) whereas the western Baltic Sea is rather perceived as an 6 7 area with no N₂ Fixation activity (Stal et al., 2003). In summary, our results provide important evidence that N₂ fixation by cyanobacteria is a significant N source not only in the Baltic 8 9 Proper but also in the Western Baltic Sea and Gulf of Finland.

10

11 **4.2.3** NO₃⁻ from atmospheric deposition

12 NO₃⁻ from atmospheric deposition is generally heavily enriched in ¹⁸O (>60‰) because of 13 reactions involving ozone (O₃), which is anomalously enriched in heavy oxygen isotopes 14 (Durka et al., 1994; Kendall et al., 2007). This is consistent with the δ^{18} O measurements at the 15 three stations around the Baltic Sea, where the averaged isotope value in winter was 77‰ 16 (Table 1).

Our results show that direct inputs of atmospheric deposition contribute significantly less 17 NO₃⁻ than all other sources. Indeed, among all basins of the Baltic Sea, that has a total area of 18 415.266 km², the maximum mean contribution was in the Western Baltic Sea 0 to 3.5% (mean 19 1.1±0.5%). Moreover, using a dataset from Michaels et al. (1993), Duce et al. (2008) 20 21 estimated that even an extremely rare and large atmospheric deposition event distributed over a 25-m mixed-layer depth would increase the reactive N concentration only by around 0.045 22 µmol l⁻¹. A study in the Kattegat estimated an input of 52 kt N yr⁻¹ from atmospheric 23 deposition, which implied rather limited nutritional support for phytoplankton (Spokes et al., 24 2006). Taking into account that in the Baltic Proper, with an area of 211.069 km², in winter 25 the mixed-layer depth is 80–100 m and that the residual NO₃⁻ pool, with a concentration of 26 3.6 μ mol l⁻¹, has a δ^{18} O-NO₃⁻ of -0.5‰, a comparable rain event with a δ^{18} O of 76.7‰ (Table 27 1) would increase the δ^{18} O-NO₃ of the residual NO₃ pool only by 0.2–0.3‰, which is within 28 our analytical error. Even though several rain events typically occur during winter, their 29 influence seems to be too low to leave a detectable isotopic imprint. Additionally, the NO_3^{-1} 30 from atmospheric deposition is presumably intensively cycled through the organic N pool in 31 32 spring and summer such that after several mineralization cycles its origin is difficult to recognize isotopically (Mayer et al., 2002). 33

In the Kalix River δ^{18} O-NO₃⁻ was clearly enriched (10.6%) compared to the values 1 determined for the Baltic Sea. We calculated that in this river up to 23.4% (mean 11.8±1.5%) 2 of the NO₃ originates from atmospheric deposition. Mayer et al. (2002) compared the isotopic 3 NO_3 signature of 16 watersheds in the USA and were able to show that riverine NO_3 derived 4 from atmospheric NO_3^- deposition and not from nitrification in soils is the dominant N input 5 in predominantly forested watersheds, when riverine NO_3^- concentrations are generally low. 6 7 Therefore only in the Kalix River, where up to 97% of the catchment with a size of 18.130 km² is covered by forests and NO₃⁻ concentrations are low during winter (Voss et al., 2011), 8 was the imprint of NO_3^- from atmospheric deposition visible; by contrast, in the southern 9 Baltic Sea and the rivers draining into it, the anthropogenic influence due to agriculture is 10 very high and therefore masks atmospheric contributions. However, NO₃ loads to the 11 northern Baltic Sea from the Kalix River and other, comparable boreal rivers are small, 12 13 comprising only about 20% of the sea's total N load (Voss et al., 2011). Thus, overall, we assume that atmospheric deposition is a very minor source of NO_3^- in the Baltic Sea. 14

15

16 **4.2.4** NO₃⁻ from pristine soils

In general, in rivers such as the Kalix River, whose catchments include pristine vegetation, 17 δ^{15} N-NO₃ values are low while those of δ^{18} O-NO₃ are high (Voss et al., 2006). This finding 18 was confirmed in the present study, in which δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻ values of 1.6‰ and 19 10.6%, respectively, were determined. In the Kalix River, the NO_3^- contribution from the 20 runoff of pristine soils as determined by SIAR is 33.8–92.8% (mean 75.3±7.9%). In pristine 21 soils the isotopic NO_3^- signal is mainly derived from nitrification, which is in agreement with 22 previous studies of small catchments, where much of the NO₃⁻ was shown to be of microbial 23 origin (Campbell et al., 2002; Kendall et al., 2007; Mayer et al., 2002). Similar δ^{15} N-NO₃⁻ 24 values were reported for areas where pristine soils were also sampled. For example, $\delta^{15}N\text{-}$ 25 NO₃⁻ and δ^{18} O-NO₃⁻ values of 1.9 and 2.8‰ were determined for Biscuit Bay (Burns et al., 26 2009) and 2.9 and 2.8‰ for the San River (Koszelnik and Gruca-Rokosz, 2013), respectively. 27 The higher δ^{18} O-NO₃ values of the Kalix River can, as discussed above, be attributed to 28 atmospheric deposition. 29

For the Gulf of Bothnia, where the catchment is dominated by pristine areas like forests (50%) and shrubs (20%), NO₃⁻ from pristine soils contributes 91.7–100% (99.0±0.9%). However, for the Baltic Proper the NO₃⁻ contribution from pristine soils is negligible, because the NO₃⁻ derived from nitrification is very low in concentrations and remains in the Gulf because of the cyclonic circulation in the Bothnian Sea and Bothnian Bay (Humborg et al.,
 2003) and the high residence time of the water (7.4 years) which results in a rather slow
 exchange with the rest of the Baltic Sea (Myrberg and Andrejev, 2006).

4

5

4.3 Comparison of isotope patterns in the water column and sediments

6 Correlations between $\delta^{15}N$ values from the water column and surface sediment is a common 7 feature in coastal basins, like Cariaco Basin (Thunell et al., 2004), Guaymas Basin, Monterey 8 Bay, and San Pedro Basin (Altabet et al., 1999). This occurs when NO₃⁻ in the surface mixed 9 layer is fully consumed, which is the case in the Baltic Proper during the spring bloom, when 10 the only significant loss comes from the sinking of particulate nitrogen (Altabet et al., 1999). 11 Moreover, high organic matter preservation seems to stimulate the similarity in the $\delta^{15}N$ in the 12 surface water and sediments as seen in other depositional environments (Thunell et al., 2004).

Overall, the comparison with the sediment data set from Voss et al. (2005) shows that the 13 isotopic signature of NO₃⁻ in the euphotic layer of the Baltic Sea is directly transferred to the 14 particulate organic nitrogen pool and is subsequently found in the sediment surface as 15 detritus, thus conserving information about the origin of this NO₃⁻ source. Additionally, we 16 could show how consistent the nitrogen input to the sediments is over the years. Even though, 17 our surface water samples were sampled from 2008 to 2011, the surface sediment samples 18 from 1993 to 2003 and deposited in the period of approximately 10 years before collection, 19 the comparison of the δ^{15} N values showed that there is no significant difference. Coastal areas 20 21 preserve the isotope signature of riverine sources while the open Baltic Sea sediments indeed 22 mirror the nitrogen input dominated by N₂ fixation. Moreover the data demonstrate that no change over time in the input of NO₃⁻ sources has occurred. 23

24

25 **5 Conclusions**

By combining dual isotope data of winter NO_3^- ($\delta^{15}N-NO_3^-$ and $\delta^{18}O-NO_3^-$) in surface waters with a Bayesian isotope mixing model (SIAR), we estimated the contribution of four major NO_3^- sources for the different basins of the Baltic Sea. A clear shift in the source of $NO_3^$ inputs, from agricultural sources in the south to runoff from pristine soils in the north, was identified. However, we could not fully determine how much of the agriculturally derived NO_3^- entering the Baltic Sea finally ends up in the open waters of its central region, where the addition of deep-water NO_3^- with similar isotope values might falsely indicate a higher 1 contribution. However, we were able to show that N_2 fixation is an important NO_3^- source in 2 the central Baltic Sea while the contribution of NO_3^- from atmospheric deposition is only a 3 minor one.

4 Because they are particularly sensitive to human pressure and global climate change, marginal seas, including the Baltic Sea, will no doubt be affected by the increases in temperature and 5 precipitation predicted for the near future (BACC, 2008). Indeed, increasing atmospheric 6 7 depositions of NO_3^- in the world's oceans have already been reported, by Duce et al. (2008) 8 and Kim et al. (2011) and, may impact northern catchments of the Baltic Sea to a larger extent. Additionally, in coastal waters under increasing eutrophication pressure the efficiency 9 10 of NO₃⁻ removal was shown to be reduced (Lunau et al., 2013; Mulholland et al., 2008), and this additional NO₃⁻ may alter the biogeochemical cycle. Therefore, the identification of NO₃⁻ 11 12 sources, especially as anticipated in response to global climate change, is important for future environmental management strategies for the Baltic Sea and other marine environments. We 13 suggest that with an adaption of the potential sources the approach used in this study can 14 easily be applied in other environments where NO_3^- is a major N contributor. 15

16

17 6 Supplement 1

Area	Date	Latitude	Longitude	Salinity	NO ₃ ⁻ +NO ₂ ⁻ [µMol]	δ ¹⁵ N (‰)	δ ¹⁸ Ο (‰)
	11.02.2008	12,4498	54,6503	8,87	3,79	4,8	3,4
	11.02.2008	12,7056	54,6959	9,22	3,81	4,3	3,5
	11.02.2008	13,0583	54,7950	8,51	4,26	4,7	4,4
Western	11.02.2008	13,2770	54,8598	8,20	2,71	5,0	5,2
Baltic Sea	12.02.2008	13,9460	54,7099	7,73	2,81	4,7	3,0
	12.02.2008	13,9886	55,0624	7,73	3,04	5,0	3,1
	12.02.2008	14,1578	54,0763	5,98	46,25	8,0	1,8
	12.02.2008	14,2827	54,6338	7,74	2,97	3,9	0,0
	25.02.2009	15,5695	55,5169	7,76	2,80	3,3	4,5
	25.02.2009	17,5808	57,2238	7,1	3,44	1,5	-0,9
	26.02.2009	17,3519	57,7003	7,01	3,52	3,1	0,1
	26.02.2009	18,2335	58,5837	6,45	3,79	2,0	-0,4
Baltic	26.02.2009	19,8832	59,7502	5,65	3,70	2,5	-0,8
гореі	12.02.2008	14,5376	55,4046	7,71	3,13	4,4	-0,8
	13.02.2008	14,7156	55,4659	7,68	3,23	3,9	0,3
	13.02.2008	15,3344	55,3835	7,51	3,07	3,6	-0,4
	13.02.2008	15,6326	55,4564	7,48	2,95	3,5	0,0
	13.02.2008	15,9834	55,2501	7,62	3,54	4,0	-1,0

18	Table S1:	Overview	of the	sampling	stations.
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	19.02.2008	15,9837	55,2498	7,64	3,27	4,3	-1,1
	19.02.2008	17,0665	55,2168	7,45	3,24	3,5	-1,0
	19.02.2008	18,2351	55,3266	7,37	2,91	3,4	-1,6
	19.02.2008	18,4013	55,5502	7,34	2,61	4,0	-1,6
	18.02.2008	18,6002	55,6339	7,37	2,76	3,8	-1,3
	18.02.2008	18,8658	55,8413	7,37	2,88	4,8	1,6
	18.02.2008	19,1677	56,0841	7,37	2,71	4,3	2,0
	18.02.2008	19,5833	56,6334	7,34	3,46	3,8	-1,7
	18.02.2008	19,8289	57,0713	7,34	3,34	3,4	-2,0
	16.02.2008	20,0506	57,3196	7,35	3,38	2,6	-2,7
	04.03.2009	24,8500	59,8036	5,24	6,82	4,1	-0,2
Gulf of	04.03.2009	25,6325	59,8659	5,09	8,12	6,4	-1,4
FILINATIO	04.03.2009	23,9989	59,6836	5,35	8,77	6,6	-1,4
	04.03.2009	22,9002	59,4836	6,11	6,95	6,4	-1,5
	27.02.2009	19,1177	60,1913	5,42	3,10	2,9	0,7
	27.02.2009	19,1464	60,6967	5,45	3,81	2,9	-1,1
	27.02.2009	19,2836	61,2265	5,54	3,96	5,8	-0,1
Gulf of	27.02.2009	19,4678	61,7032	5,57	3,40	2,4	-1,1
Dounnia	27.02.2009	19,7182	62,1441	5,45	3,49	3,2	-1,8
	28.02.2009	19,9682	62,5872	5,52	3,66	1,8	-2,2
	28.02.2009	20,4862	63,0417	5,54	4,31	4,0	-0,7
	20.11.2011	21,584	63,834	3,1	5,53	1,83	-2,79
	15.11.2009	21,38187	55,30128	0	145,03	10,0	1,5
	01.12.2009	21,38187	55,30128	0	179,18	9,0	1,3
Southern	07.12.2009	21,13293	55,69043	0	114,87	10,0	1,7
1/1/612	22.02.2010	21,13293	55,69043	0	140,47	8,8	1,3
	17.01.2003	12,1429	54,03246	0	259,00	8,2	6,5
	13.02.2003	12,1429	54,03246	0	135,00	9,3	6,5
Northern rivers	11.11.2009	22,8415	65,9335	0	7,41	1,6	10,6

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Location	Date	NO_3^- (umol L ⁻¹)	δ^{15} N-NO ₃	δ^{18} O-NO ₃
Warnemünde	21.12.2009	52.7	2.1	75.6
Warnemünde	04.01.2010	51.2	1.1	68.3
Warnemünde	19.01.2010	104.4	0.2	84.6
Warnemünde	01.02.2010	50.8	0.8	65.8
Warnemünde	19.02.2010	94.4	0.6	79.5
Warnemünde	22.02.2010	106.8	2.1	81.8
Sännen	Dec. 2009	12.1	-0.3	69.2
Sännen	Jan. 2010	60.4	-1.1	81.8
Sännen	Feb. 2010	69.3	-2.1	77.0
Majstre	Dec. 2009	30.7	-0.8	83.8

Table 1. NO₃⁻ concentrations and δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻ values of wet atmospheric deposition. Data are from Warnemünde (Germany), Sännen (Sweden), and Majstre (Sweden).

- 1 Table 2. Means and standard deviations of the δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻ values of the NO₃⁻
- 2 sources used in the SIAR mixing model. For further details, see Material and Methods, SIAR

3 mixing model.

Source	δ^{15} N-NO ₃ ⁻ (mean ± SD)	δ^{18} O-NO ₃ ⁻ (mean ± SD)	n	Origin	Reference
NO ₃ ⁻ from atmospheric deposition	0.3±1.4	76.7±6.8	10	Warnemünde (Germany), and Sännen and Majstre (Sweden)	This study
NO ₃ ⁻ from pristine soils	1.3±1.4	1.5±0.9	5	Groundwater	Deutsch et al., 2006
NO ₃ ⁻ from agricultural runoff	9.9±1.5	4.6±1.0	21	Tile-drain outlets, Warnow River	Deutsch et al., 2006
NO ₃ ⁻ from N ₂ fixation	-1.0±1.0	-0.7±2.9	0	Estimated	Carpenter et al., 1999, 1997; Bourbonnais et al., 2009, 2012; Montoya et al., 2002; Sigman et al. 2009

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1 Table 3. Source attribution results: Mean, standard deviation, and minimum and maximum values for the potential contributions of four potential

2 NO ₃ ⁻ sources for the areas Western Baltic Sea, Baltic Proper, Gulf of Finland, Gulf of Bothnia, southern rivers, and	d northern rivers.
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Area	NO ₃ ⁻ from atmospheric deposition		NO_3^- from pris	m pristine soils NO_3^- from agricultural		icultural runoff	NO_3^- from N_2 fixation	
	Mean±SD	Min-Max	Mean±SD	Min-Max	Mean±SD	Min-Max	Mean±SD	Min-Max
Western Baltic Sea	1.1±0.5	0.0–3.5	7.5±5.9	0.0–42.0	53.5±3.2	41.0–66.5	37.9±5.1	11.0–51.9
Baltic Proper	0.1±0.1	0.0–0.9	2.1±1.9	0.0–14.7	39.0 ± 1.6	32.8-45.5	58.8±2.0	49.3–65.3
Gulf of Finland	0.2±0.2	0.0–2.0	2.4±2.1	0.0–24.3	51.9±3.0	40.9–63.4	45.5±3.2	32.7–59.0
Gulf of Bothnia	0.1±0.1	0.0–0.5	99.0±0.9	91.7–100.0	1.0±0.9	0.0-8.2	-	-
Southern rivers	0.2±0.1	0.0–1.3	6.4±4.2	0.0–24.5	93.5±4.2	75.2–100.0	-	-
Northern rivers	11.8±1.5	6.6–23.4	75.3±7.9	33.8–92.8	12.9±8.1	0.0–57.2	-	-

1 Table 4. Comparison of δ^{15} N-NO₃⁻ values from surface water samples and δ^{15} N values from

	Baltic southern coastal areas/ Gulf of Finland	Central Baltic Proper	
δ^{15} N sediments (‰) δ^{15} N-NO ₃ surface water	7.3 ± 2.1	3.5 ± 0.6	Voss et al. 2005
column (‰)	7.9±1.8	3.6 ± 1.0	This study

2 sediments samples in sub-regions of the Baltic Sea.



Figure 1. Station Map of the Baltic Sea and percent contribution of the four nitrate sources,
NO₃⁻ from atmospheric deposition (blue), pristine soils (red), agricultural runoff (green), and
N₂ fixation (black), for the Western Baltic Sea, Baltic Proper, Gulf of Finland, Gulf of
Bothnia, southern rivers, and northern rivers. Stations are indicated as black dots.



Figure 2. Surface water column NO₃⁻ concentrations (A), δ^{15} N-NO₃⁻ values (B), and δ^{18} O-NO₃⁻ values (C) for the Baltic Sea. Stations are indicated as black dots. Additional NO₃⁻ concentrations were obtained from the Data Assimilation System (DAS) (http://nest.su.se/das/) in winter (Nov.–Feb.) of the years 2000 to 2012.



Figure 3. δ¹⁵N from sediment samples vs. δ¹⁵N- NO₃⁻ from surface water samples. δ¹⁵N
values from sediments were taken from Voss et al. (2005). The positive slope suggests a tight
coupling between δ¹⁵N- NO₃⁻ in surface waters and δ¹⁵N in sediment samples.



Figure 4. Station map for the comparison of isotope patterns in the water column and
sediments. Gray circles are the stations referred to in Voss et al. (2005) and black crosses are
those from this study. Isotope values were compared at stations with the same number.