

This is a self-archived version of an original article. This version may differ from the original in pagination and typographic details.

Author(s): Aalto, Sanni L.; Asmala, Eero; Jilbert, Tom; Hietanen, Susanna

Title: Autochthonous organic matter promotes DNRA and suppresses N2O production in sediments of the coastal Baltic Sea

Year: 2021

Version: Accepted version (Final draft)

Copyright: © 2021 Elsevier

Rights: In Copyright

Rights url: http://rightsstatements.org/page/InC/1.0/?language=en

Please cite the original version:

Aalto, S. L., Asmala, E., Jilbert, T., & Hietanen, S. (2021). Autochthonous organic matter promotes DNRA and suppresses N2O production in sediments of the coastal Baltic Sea. Estuarine, Coastal and Shelf Science, 255, Article 107369. https://doi.org/10.1016/j.ecss.2021.107369

Autochthonous organic matter promotes DNRA and suppresses N_2O production in sediments of the coastal Baltic Sea

Sanni L. Aalto, Eero Asmala, Tom Jilbert, Susanna Hietanen

PII: S0272-7714(21)00222-5

DOI: https://doi.org/10.1016/j.ecss.2021.107369

Reference: YECSS 107369

To appear in: Estuarine, Coastal and Shelf Science

Received Date: 14 May 2020

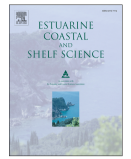
Revised Date: 31 January 2021

Accepted Date: 6 April 2021

Please cite this article as: Aalto, S.L., Asmala, E., Jilbert, T., Hietanen, S., Autochthonous organic matter promotes DNRA and suppresses N₂O production in sediments of the coastal Baltic Sea, *Estuarine, Coastal and Shelf Science* (2021), doi: https://doi.org/10.1016/j.ecss.2021.107369.

This is a PDF file of an article that has undergone enhancements after acceptance, such as the addition of a cover page and metadata, and formatting for readability, but it is not yet the definitive version of record. This version will undergo additional copyediting, typesetting and review before it is published in its final form, but we are providing this version to give early visibility of the article. Please note that, during the production process, errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

© 2021 Published by Elsevier Ltd.



1 Autochthonous organic matter promotes DNRA and suppresses N₂O

2 production in sediments of the coastal Baltic Sea

- 3 Sanni L. Aalto^{1,2*}, Eero Asmala³, Tom Jilbert^{3,4}, Susanna Hietanen^{3,4}
- ¹Department of Environmental and Biological Sciences, University of Eastern Finland, P.O. Box 1627,
 70211 Kuopio, Finland
- ²Department of Biological and Environmental Science, University of Jyväskylä, P.O. Box 35, 40014
 Jyväskylä, Finland
- 8 ³Tvärminne Zoological Station, University of Helsinki, 10900 Hanko, Finland
- 9 ⁴Ecosystems and Environment Research Program, Faculty of Biological and Environmental Sciences,
- 10 00014 University of Helsinki, Helsinki, Finland
- 11 Corresponding author: Sanni L. Aalto, sheaa@aqua.dtu.dk
- 12 *current address: Technical University of Denmark, DTU Aqua, Section for Aquaculture, The North
- 13 Sea Research Centre, P.O. Box 101, DK-9850 Hirtshals, Denmark

14 Abstract

Coastal environments are nitrogen (N) removal hot spots, which regulate the amount of land-15 16 derived N reaching the open sea. However, mixing between freshwater and seawater creates 17 gradients of inorganic N and bioavailable organic matter, which affect N cycling. In this study, we 18 compare nitrate reduction processes between estuary and offshore archipelago environments in the 19 coastal Baltic Sea. Denitrification rates were similar in both environments, despite lower nitrate and 20 carbon concentrations in the offshore archipelago. However, DNRA (dissimilatory nitrate reduction 21 to ammonium) rates were higher at the offshore archipelago stations, with a higher proportion of 22 autochthonous carbon. The production rate and concentrations of the greenhouse gas nitrous oxide 23 (N_2O) were higher in the estuary, where nitrate concentrations and allochthonous carbon inputs are 24 higher. These results indicate that the ratio between nitrate and autochthonous organic carbon governs the balance between N-removing denitrification and N-recycling DNRA, as well as the end-25 26 product of denitrification. As a result, a significant amount of the N removed in the estuary is 27 released as N₂O, while the offshore archipelago areas are characterized by efficient internal recycling 28 of N. Our results challenge the current understanding of the role of these regions as filters of land-29 to-sea transfer of N.

30 Keywords: denitrification; DNRA; DOM; estuary; N₂O; sediment organic matter

31 **1** Introduction

32 Coastal systems are transitional zones where riverine freshwater mixes with saline seawater. They 33 are important hot spots in the nitrogen (N) cycle, as N transformations in coastal ecosystems regulate the 34 amount of land-derived N reaching the open sea (Bouwman et al., 2013). Various coastal processes, 35 including assimilation to biomass and subsequent microbial degradation of organic matter, modulate 36 land-to-sea transfer of N. Crucially, N may be removed from biogeochemical cycling in estuaries by a 37 sequence of sedimentary microbial processes terminating in denitrification, which releases dinitrogen gas (N_2) into the atmosphere. Denitrification is a critical part of the 'coastal filter'; the set of biogeochemical 38 39 processes regulating the impact of riverine nutrient inputs on coastal eutrophication (Asmala et al., 40 2017).

Denitrification rates in coastal environments depend on nitrate concentrations, which typically 41 42 decrease from near-shore to offshore areas (Asmala et al., 2017). However, heterotrophic 43 denitrification also depends on the presence of bioavailable organic carbon (OC) in coastal sediments (Hellemann et al., 2017; Hietanen and Kuparinen, 2008). Higher OC bioavailability has 44 45 been suggested to promote denitrification in freshwater stream sediments (Barnes et al., 2012; 46 Stelzer et al., 2014), raising the question of whether the same is true in coastal marine systems. 47 Coastal systems often display strong gradients in both nitrate concentrations, and in sedimentary OC 48 sources and characteristics, with distance away from river mouths. Typically, the relative amount of 49 terrestrial OC in sediments decreases gradually along the coastal salinity gradient, while the amount 50 of fresh, autochthonous phytoplankton-derived OC increases (Fellman et al., 2011; Goñi et al., 2003; 51 Spencer et al., 2007). Combined, these observations suggest that coastal nitrate removal efficiency through denitrification could be related to the availability of both nitrate and bioavailable OC 52 53 (Asmala et al., 2017).

54 The balance in the availability of nitrate and bioavailable carbon may also influence rates of 55 alternative nitrate reduction pathways. Heterotrophic dissimilatory nitrate reduction to ammonium (DNRA), which retains N as biologically reactive ammonium in the aquatic system (e.g., Giblin et al., 56 57 2013), is the prominent pathway under conditions of high OC availability relative to nitrate (Hardison et 58 al., 2015; Kraft et al., 2014). This phenomenon may occur because under nitrate-limited conditions, DNRA 59 makes more efficient use of the available electron acceptors (6 electrons transferred per mole of N 60 reduced compared to 3 for denitrification), and therefore maximizes entropy production (Algar and 61 Vallino 2014). Furthermore, OC composition is as important as OC availability in controlling the nitrate 62 reduction end-product (Carlson et al., 2020). From this, it follows that the importance of DNRA in net 63 nitrate reduction may increase towards the open sea where terrestrial influence decreases (lower nitrate

and higher bioavailable carbon concentrations). Indeed, high contributions of DNRA to total nitrate reduction were recently observed in the Baltic Sea offshore region (Hellemann et al., 2020) and in Australian estuaries (Kessler et al., 2018). Therefore, outer coastal areas may recycle nitrate more efficiently than remove it, in comparison with near-shore areas with a lower bioavailable OC to nitrate ratio, which favours denitrification.

69 Incomplete denitrification leads to the production of nitrous oxide (N_2O). The proportion of N_2O 70 production from total denitrification can increase with DIN concentrations (Murray et al., 2015), and 71 decrease with increased bioavailable carbon (Zhao et al., 2014). This suggests that among other variables 72 (e.g. oxygen, temperature, salinity, and rates of nitrogen fixation and nitrification; Foster and Fulweiler, 73 2016; Silvennoinen et al., 2008; Zhao et al., 2014), OC bioavailability is an important factor controlling 74 denitrification-derived N₂O production in coastal ecosystems, and N₂O production the rates may be 75 higher in near-shore estuarine environments with low amounts of bioavailable OC and high nitrate 76 concentrations. Hence, OC characteristics and especially bioavailability may play a key role in many 77 aspects of coastal sedimentary N cycling. These factors must be deconvolved from the effects of 78 nitrate gradients to properly understand the coastal N cycle.

79 The overall bioavailability of aquatic OC can be assessed with optical proxies of dissolved organic 80 matter (DOM), derived from the absorbance and fluorescence properties of the colored dissolved 81 organic matter (CDOM) (Asmala et al., 2013). A range of optical proxies (e.g. the humification index 82 (HIX) and the index of recent autochthonous contribution (BIX)) have been derived to characterize 83 the DOM pool (Huguet et al., 2009; Murphy et al., 2008). We assume DOM in sediment porewaters 84 to reflect the broad overall organic matter composition of sediments, and optical analysis of 85 porewater DOM composition provides a tool for characterizing the source and bioavailability of sedimentary carbon. Porewater DOM characterization potentially provides additional information to 86 traditional approaches such as C/N ratios or δ^{13} C of bulk organic matter. 87

88 Here, we investigate the combined influence of nitrate availability and organic matter composition 89 on nitrate reducing processes in coastal sediments in the northern Baltic Sea. The Baltic is a semi-90 enclosed shallow brackish water basin with significant anthropogenic N loading. In 2010, the total N load 91 to the Baltic was 977 000 tons, of which 758 000 tons was waterborne (Helcom 2015), yielding a 92 waterborne N load from the catchment of 0.44 tonnes/km². The Baltic Sea coastal zone (29% of total 93 Baltic Sea area) was estimated to remove 16% of land-derived N inputs, the N removal efficiency varying 94 between different types of coastal ecosystems (Asmala et al. 2017). Denitrification dominates N₂ 95 production in Baltic Sea coastal ecosystems, with anammox playing only a minor role (Bonaglia et al., 96 2014; Hietanen, 2007; Thamdrup and Dalsgaard, 2002). Knowledge on the balance between

97 denitrification and DNRA is limited for this region, but results from an anthropogenically impacted Baltic 98 Sea estuary suggest that denitrification is the main process (Bonaglia et al., 2014) due to the high DIN 99 availability, and that the contribution of DNRA increases to 30-50% of total nitrate reduction in the 100 offshore region (Hellemann et al., 2020). The limited data from oligotrophic coastal sediments of the 101 Baltic Sea, where availability of labile organic carbon limits the denitrification process, also indicate that 102 N₂O production from benthic denitrification is low (N₂O:N₂ <0.02) (Hellemann et al., 2017).

103 In this study, we measured porewater DOM characteristics, nitrous oxide concentrations, and N 104 processes along a gradient encompassing near-shore (estuary) and offshore archipelago stations in a 105 coastal region of the Baltic Sea to examine the effects of both nitrate availability and OC characteristics on nitrate reduction processes. We hypothesized that higher nitrate availability and 106 107 terrestrial dominance of the carbon pool (i.e. low quantities of bioavailable carbon compared to 108 nitrate) would promote denitrification and possibly N₂O production at the near-shore estuarine 109 stations. Conversely, we hypothesized that the significance of DNRA as a nitrate reduction process 110 would increase at the offshore archipelago stations due to a higher amount of bioavailable carbon 111 and/or lower nitrate concentrations.

112 2 Materials and methods

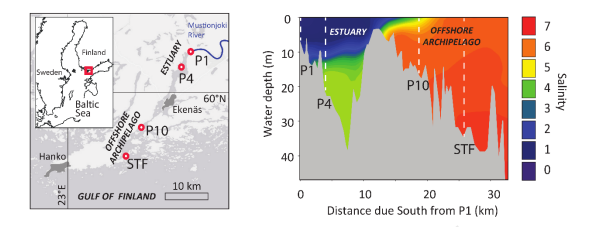
113 2.1 Study area, sampling, and water column analyses

114 The study was conducted in the Finnish coastal area of the Gulf of Finland, Baltic Sea. 115 Pohjanpitäjänlahti is a long and narrow embayment that receives freshwater input from the river 116 Mustionjoki and brackish water input from the adjacent coastal archipelago of the Baltic Sea (Fig. 1). 117 A shallow (2-3 m) sill area, with a dredged 6 m channel through it, separates the estuary from the 118 offshore region connecting to the open Baltic Sea, limiting the water exchange between the estuary and the offshore region. The inner basin is salinity-stratified, with a pronounced pycnocline at 10-15 119 120 m water depth, which leads to seasonal hypoxia in summer and autumn. Inflows of brackish water 121 over the sill usually occur in late autumn – early winter, leading to temporary ventilation of the basin 122 (Malve et al., 2000). The adjacent offshore region experiences temperature stratification in summer, leading to the development of hypoxia in isolated areas. However, much of that remains oxic 123 124 throughout the annual cycle due to sufficient vertical mixing and exchange of water masses. The 125 catchment of the Mustionjoki has a large proportion of lakes (11%; Mattsson et al., 2005) and 126 several hydropower plants regulating the flow. This characteristic leads to extensive processing of 127 the riverine nutrients and organic matter already within the lotic system and relatively low areaspecific loading of organic carbon to the estuary (Räike et al., 2012). 128

To monitor water column N₂O concentrations, water column sampling was conducted at stations P4 129 ("estuary", see Fig. 1) and STF ("offshore archipelago") at 5 m depth intervals using a 5L Limnos 130 sampler on multiple occasions during 2015-2017. Subsamples for determination of dissolved N₂O 131 132 were collected in triplicate by filling 60mL plastic syringes directly from a Limnos water sampler on 133 board. In the laboratory, the water volume in the syringe was reduced to 30 mL, and 31mL of 5.0 134 purity N2 gas was injected to create a headspace. Syringes were left at 20 C for 30 min and then vigorously shaken for 3 min, after which 25mL of the headspace was injected into a pre-evacuated 135 136 12mL gastight glass vial (LabCo Exetainer model 839W). Nitrous oxide concentrations in the headspace were determined using an Agilent Technologies 7890B gas chromatograph equipped with 137 138 electron capture detector (ECD) and the results calculated as in Myllykangas et al. (2017).

Sampling was carried out at two stations in the estuary (stations P1 and P4) on 6th of June 2017 and 139 15th–16th of August 2017 and at two stations in the offshore archipelago region (stations P10 and 140 STF; Fig. 1) only on 15th–16th of August 2017. Sampling occasions were chosen to represent situations 141 142 with high (June) and low (August) amount of fresh, recently deposited phytoplankton-derived 143 material on the sediment surface (Heiskanen and Kononen, 1994). Temperature, salinity and oxygen 144 were determined using a YSI CTD equipped with an optical oxygen sensor. Sediment cores were collected using a Gemax twin sampler (core diameter 9 cm, length of a core 30 - 50 cm) from each 145 sampling station. Water samples were collected using a 5L Limnos water sampler from 1 m depth to 146 147 1 m above the sediment at 2-5 m intervals, and from the overlying water of the sediment cores. Oxygen samples for Winkler titration (150 ml) were treated immediately with fixing reagents and 148 149 analyzed the following day. Dissolved inorganic nitrogen (ammonium, nitrite and nitrate) samples 150 were collected in acid-washed plastic bottles, filtered through 0.2µm polycarbonate filters and 151 stored dark at 4°C. Concentrations were measured using a discrete photometric analyzer (Thermo 152 Scientific Aquakem 250) the following day. Theoretical 3-sigma detection limits were as follows: 153 ammonium 0.11 μ M, nitrate and nitrite 0.08 μ M.

Figure 1. (left) Sampling locations in the Pohjanpitäjänlahti system on the Finnish coast of the Gulf of Finland, northern Baltic Sea. Stations P1 and P4 are classified as "estuary" stations, while P10 and STF are classed as "offshore archipelago". The Mustionjoki river discharges into the Pohjanpitäjänlahti estuary close to station P1. (right) Bathymetric detail of the transect through the sampling locations, showing typical salinity distribution (data shown here from June 2015, redrawn from Jilbert et al., 2018). A shallow sill close to the city of Ekenäs restricts exchange of brackish deeper waters between the offshore archipelago and estuary.





162 2.2 Sediment and porewater analyses

Sediment cores were collected using a Gemax twin sampler (core diameter 9 cm, length of core 30 -50 cm) from each sampling station. Sediment water content and porosity were determined from the upper portion of each core (0–6 cm) (Burdige, 2006). Sediment total C and N content (%C, %N) of the upper portion was determined by Thermal Combustion Elemental Analysis (TCEA) at Tvärminne Zoological Station with precision and accuracy of < 2.5% RSD. Sedimentary inorganic carbon and nitrogen are assumed insignificant in this setting, hence %C_{tot} and %N_{tot} are assumed equal to organic carbon and nitrogen, respectively (%C_{org} and %N_{org}).

170 Porewater DOC and CDOM samples were taken from the surface sediment layer (0-1 cm) of three 171 replicate cores. In the laboratory, pore water was extracted with centrifuging (1500 rpm for 10 min), 172 and filtered through a combusted (4 h 450 °C) glass fiber filter (47 mm, VWR collection GF/F). DOC 173 concentration in porewaters was measured with a Shimadzu TOC-V_{CPH} analyzer. The detection limit 174 for DOC analysis was 40 µmol L⁻¹. CDOM absorption was measured using a Shimadzu 2401PC 175 spectrophotometer with 1 cm quartz cuvette over the spectral range from 200 to 800 nm with 1 nm 176 intervals. Ultrapure water served as the blank for all samples. Excitation-emission matrices (EEMs) of 177 fluorescent DOM (FDOM) were measured and corrected as in Asmala et al., (2018). For assessing the 178 terrestrial signature of the porewater DOM, fluorescence peaks (peaks A, C, M, and T; Coble, 1996), 179 humification index (HIX; Zsolnay et al., 1999) and biological index (BIX; Huguet et al., 2009) were 180 calculated from the measured and corrected EEMs. Processing of the EEMs was done using the 181 eemR package for R software (Massicotte, 2018).

182 2.3 Sedimentary nitrogen process rates

Samples for benthic nitrate reduction rate measurements (n=8 per sampling station) were collected into acrylic cores (\emptyset 2.3 cm, length 15 cm), which were pushed gently into the sediment so that 1/3 of each core was filled with sediment and the rest with overlying water, capped and placed in a

water bath at in situ temperature. The four cores were immediately enriched with ¹⁵N-labelled 186 nitrate to a final concentration of 100 µM ¹⁵N-NO₃⁻ (K¹⁵NO₃ Sigma Aldrich, 98% ¹⁵N-atm), closed and 187 incubated under stirring at *in situ* temperature in dark for 3-4 h. Enrichment with 200 µM ¹⁵N-NH₄⁺ 188 (¹⁵NH₄Cl Cambridge Isotope Laboratories, 99% ¹⁵N-atm; 4 replicate cores) was used to exclude 189 anammox and measure nitrification (data not shown). After incubation, sediment and overlying 190 191 water in the samples were mixed and 12 mL subsamples were transferred into gas-tight glass vials 192 (Labco Exetainer model 739W) with 0.5 mL ZnCl₂ (100 % w/v, Merck) after a brief sediment settling period. Isotopic composition of N₂ and N₂O was analysed with a TraceGas preconcentrator system 193 194 interfaced with an IsoPrime 100 continuous flow isotope ratio mass spectrometer (CF-IRMS; 195 Isoprime Ltd, Cheadle Hulme, UK) at the Department of Environmental Sciences, University of Jyväskylä, Finland as in Hellemann et al., (2017). The detection limits were 320 nmol L⁻¹ for ²⁹N₂ 11 196 nmol L^{-1} for ${}^{30}N_2$, 397 pmol L^{-1} for ${}^{45}N_2O$, and 322 pmol L^{-1} for ${}^{46}N_2O$. 197

The remaining ¹⁵NO₃⁻-enriched slurry was mixed again, and 20 mL samples for ¹⁵NH₄⁺ analysis were 198 collected into 50 mL centrifuge tubes, treated with 1 mL of ZnCl₂, and frozen immediately. Before 199 ¹⁵NH₄⁺ analysis, NH₄⁺ attached to the sediment particles was desorbed using KCl extraction. The 200 isotopic composition of NH_4^+ in the samples was analyzed after conversion to N_2 using alkaline 201 202 hypobromite iodine solution (Risgaard-Petersen et al., 1995) as in Hellemann et al., (2020). A standard series of ¹⁵NH₄⁺ (5; 10; 15 µM, 5% ¹⁵N-atm from ¹⁵NH₄Cl Cambridge Isotope Laboratories, 203 98% ¹⁵N-atm) was prepared, treated and analyzed parallel with samples to calculate conversion 204 efficiency and ¹⁵N recovery, which was > 85 %. 205

The N₂ and N₂O producing denitrification rates were calculated from the production rates of $^{29}N_2$, 206 $^{30}N_2$ and $^{45}N_2O$, $^{46}N_2O$), and partitioned to denitrification based on water column nitrate (D_w) and 207 coupled nitrification-denitrification (D_n) (Nielsen, 1992). DNRA rates were calculated from the 208 production rates of ${}^{15}NH_4^+$ and the production rates of ${}^{29}N_2$, ${}^{30}N_2$ and ${}^{45}N_2O$, ${}^{46}N_2O$ in the same 209 incubation cores according to Christensen et al., (2000). It was assumed that DNRA takes place in the 210 same layers as denitrification, meaning that the ¹⁵N labeling of NO₃⁻ reduced to ammonia equals the 211 ¹⁵N labeling of NO₃⁻ reduced to N₂/N₂O. Total N₂ production (ΣN_2) was calculated as $\Sigma N_2 = D_w N_2 + N_2$ 212 $D_n N_2$ and total N_2O production ($\sum N_2O$) as $\sum N_2O = D_w N_2O + D_n N_2O$. The total denitrification was 213 then defined as $\Sigma N_2 + \Sigma N_2 O$ and total nitrate reduction as $\Sigma N_2 + \Sigma N_2 O$ + DNRA. The hourly rates were 214 215 scaled to day by multiplying with 24h. The N_2O produced in coupled nitrification-denitrification was divided into the rate of N₂O produced in the nitrification stage and the denitrification stage of the 216 217 coupled nitrification-denitrification according to Dong et al., (2006).

218 2.4 Statistical analysis

The data analysis was conducted using R (version 3.6.3; R Core Team, 2020). The differences in the porewater DOM characteristics, and N processes between estuary and offshore archipelago region were examined with one-way ANOVA, or if the assumptions on the normality and equal variances were not met, with Mann-Whitney U test. The relationship between DOM variables and N processes were examined with Pearson correlation analysis, and relative DNRA (%DNRA) and N₂O (%N₂O) and DOC and bioavailable carbon fraction (protein-like DOM fluorescence) were further examined with linear regression.

226 **3 Results**

227 3.1 Hydrography

In both estuary and offshore archipelago, the water column was well oxygenated during the sampling campaigns despite being stratified, with a thermocline present at all stations between 3.5-10 m depth (Table 1; Suppl. Fig. 1). At the estuary stations, closer to the direct influence of the Mustionjoki River, a pronounced halocline was present (Suppl. Fig. 1).

Table 1. Temperature (T), salinity, oxygen concentration (O_2) , and DIN concentrations (NO_x^-, NH_4^+) in near-bottom water and sediment C:N at the estuary and offshore archipelago sampling stations.

	Station	Sampling time	т∘с	salinity	Ο ₂ μΜ	NO _x ⁻ μM	${\sf NH_4}^+ \mu {\sf M}$	C:N
Offshore Estuary archipelago	P1	June 2017	5.8	4.0	234	11.1	0.7	18.7
	P4	June 2017	3.3	5.1	236	13.0	1.5	12.4
	P1	August 2017	13.8	3.3	155	1.7	3.2	21.6
	P4	August 2017	4.7	5.0	126	14.1	6.1	12.7
	P10	August 2017	10.1	6.2	176	1.6	5.8	11.2
	STF	August 2017	8.8	6.4	216	1.4	4.0	10.1

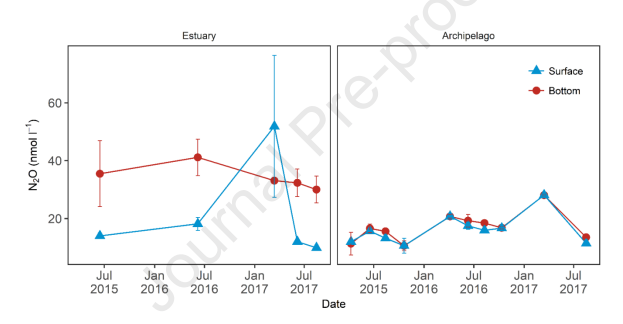
234

235 **3.2** Dissolved inorganic nitrogen and nitrous oxide

The near-bottom combined nitrite+nitrate (NO_x^-) concentrations decreased as expected from nearshore estuary to offshore archipelago stations. At the estuary stations P1 and P4, near-bottom NO_x^-

- concentrations varied between 3-14 μ M (Table 1; Suppl. Fig. 2). Near-bottom NO_x⁻ concentrations were consistently low ($\leq 1.6 \mu$ M) at the offshore archipelago stations P10 and STF. Near-bottom ammonium (NH₄⁺) concentrations (1-6 μ M) were similar at all sampling stations. Dissolved nitrous oxide (N₂O) concentrations were consistently high (25-50 nM at P4, Fig. 2) below the halocline in the estuary. Surface waters at the offshore archipelago stations, P4 and STF, and deeper waters at STF, had lower N₂O concentrations (10-30 nM), except for a high value in the surface waters of P4 under
- ice cover in March 2017.

Figure 2. Nitrous oxide (N_2O) concentration in water column above (blue triangles) and below halocline (red circles) between April 2015 and August 2017 at the near-shore estuary (station P4) and offshore archipelago (station STF) stations. Points indicate mean value and error bars \pm 1 standard deviation. Number of observations per each mean value in the figure ranges between 3 and 24, the median number of observations being 10.



250

251 3.3 Organic carbon source proxies

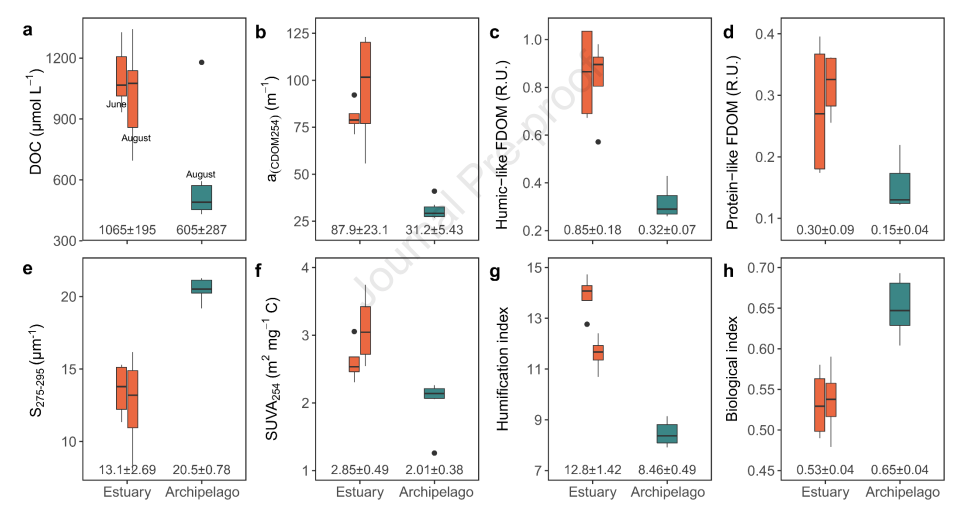
252 All the sampled sediments were muddy, with surface (0-1 cm) porosities ranging from 0.94 to 0.97. 253 Sediment C:N ratio decreased from the estuary (16 ± 5) to the offshore archipelago stations (11 ± 1) 254 (Table 1). The amount of bulk dissolved organic matter in the porewater, as indicated by the DOC 255 concentration, was almost twice as high at the estuary stations as at the offshore archipelago 256 stations (Fig. 3a). The ratio (mean ± SD) between DOC concentrations in the uppermost sediment 257 layer (0-1 cm) and near bottom NO_x (DOC:NO_x⁻) was 2.5 \pm 2.9 at the estuary stations and 4.8 \pm 1.3 at the offshore archipelago stations. Organic matter characteristics were on average more terrestrial-258 259 like at the estuary than at the offshore archipelago stations (one-way ANOVA, p < 0.05), as indicated 260 by optical proxies: higher CDOM absorption at 254 nm ($a_{(CDOM254)}$), DOC-specific UV absorbance (SUVA254), humic- and protein-like DOM fluorescence (peak C and T, respectively) and higher 261

humification index (HIX). Also, at the offshore archipelago stations, UV absorption slope ($S_{275-295}$) and biological index (BIX) were higher than at the estuary stations (p < 0.05; Fig. 3), indicating higher contribution of autochthonous bioavailable carbon with smaller molecular size.

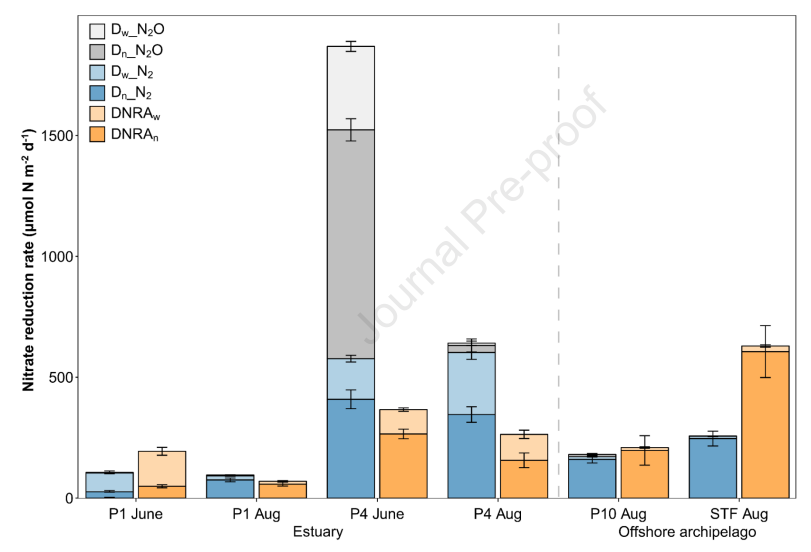
265 **3.4** Nitrogen transformation rates in estuary and offshore archipelago sediments

266 All nitrate reduction rates varied substantially between the sampling stations, as both the highest 267 and lowest rates were measured at the estuarine stations (Fig. 4, Suppl. Fig. 3). Total denitrification $(\Sigma N_2 + \Sigma N_2 O)$ rates and total nitrate reduction $(\Sigma N_2 + \Sigma N_2 O + DNRA)$ rates did not differ significantly 268 269 between the estuary and offshore archipelago stations (Mann-Whitney U test, p > 0.05, Fig. 4, Suppl. 270 Fig. 3). No anammox was detected (data not shown). Denitrification rates based on water column 271 nitrate ($D_w N_2$, $D_w N_2$) were higher at the estuary stations ($D_w N_2$: one-way ANOVA, p = 0.004; 272 $D_w N_2O$: p < 0.001), but the coupled nitrification-denitrification process rates ($D_n N_2$, $D_n N_2O$) were 273 dominant and equal between the estuary and offshore archipelago stations (p > 0.05, Fig. 4, Suppl. 274 Fig. 3). Similarly, DNRA rates based on water column nitrate (DNRA_w) were higher at the estuary 275 stations (p = 0.006), while total DNRA rates (p = 0.024) and the proportion of DNRA of total nitrate 276 reduction (%DNRA; p = 0.03) and nitrification-fed DNRA (DNRA_n; p = 0.003) rates were higher at the 277 offshore archipelago stations (Fig. 5, Suppl. Fig. 3). The proportion of N₂O produced in nitrate reduction (N_2O) as well as the proportion of N_2O produced from the denitrification stage of 278 279 coupled nitrification-denitrification were higher at the estuary stations than at the offshore 280 archipelago stations (N_2O : p < 0.001, N_2O from denitrification: p = 0.006), being especially high at 281 P4 in August (Fig. 5). Significant relationships between organic carbon characteristics (source 282 proxies) and both %DNRA (decreasing with higher terrestrial OM share) and %N₂O (increasing with 283 higher terrestrial OM share) were observed (Suppl. Table 1), while no relationship was found with 284 total denitrification rates. Notably, the variance of either %DNRA or %N₂O was not explained by bulk 285 carbon concentration (DOC) (Fig. 6a–b). Rather, protein-like DOM fluorescence (a common proxy for 286 biologically labile organic carbon) had a strong negative relationship with %DNRA and strong positive 287 relationship with %N₂O (Fig. 6c–d).

- Figure 3. Porewater (0–1 cm) DOM quantity and quality characteristics at the estuary stations in June (left orange bar, n = 6) and August (right orange bar, n
- = 6) and at the offshore archipelago stations in August (n = 6): a) dissolved organic carbon (DOC), b) CDOM absorption coefficient at 254 nm ($a_{(CDOM254)}$, c)
- humic-like DOM fluorescence (Peak C), d) protein-like DOM fluorescence (Peak T), e) CDOM spectral slope between 275–295 nm (S_{275–295}), f) DOC-specific
- UV absorbance at 254 nm (SUVA₂₅₄), g) humification index (HIX) and h) biological index (BIX). Mean values ± standard deviation for estuary and offshore
- archipelago groups are also given. The two groups are significantly different for each variable (one-way ANOVA, p < 0.05).



- Figure 4. DNRA and denitrification rates at the estuary (P1, P4) and offshore archipelago (P10, STF) stations. D_w denotes water column nitrate based process
- and D_n process based on the nitrate produced through sediment nitrification. Bars represent mean values ± standard error for four sediment core
- 297 replicates.



- 299 Figure 5. Differences in the a) absolute and b) relative rates of DNRA, and the proportion of N₂O of c)
- 300 total nitrate reduction, and d) originating from denitrification stage of total N₂O production during
- 301 coupled nitrification-denitrification process between the estuary stations in June (left orange bar, n
- = 8) and August (right orange bar, n = 8) and at the offshore archipelago stations in August (n = 8).
- 303 Mean values ± standard deviation for estuary and offshore archipelago groups are given. The two

304 groups are significantly different for each variable (one-way ANOVA/Mann-Whitney U test, p < 0.05).

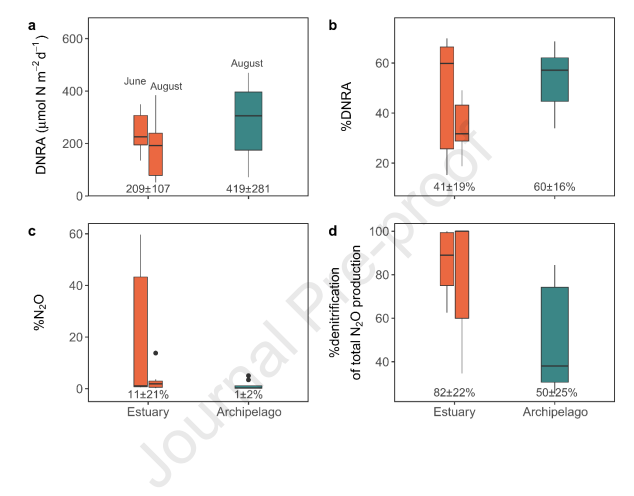
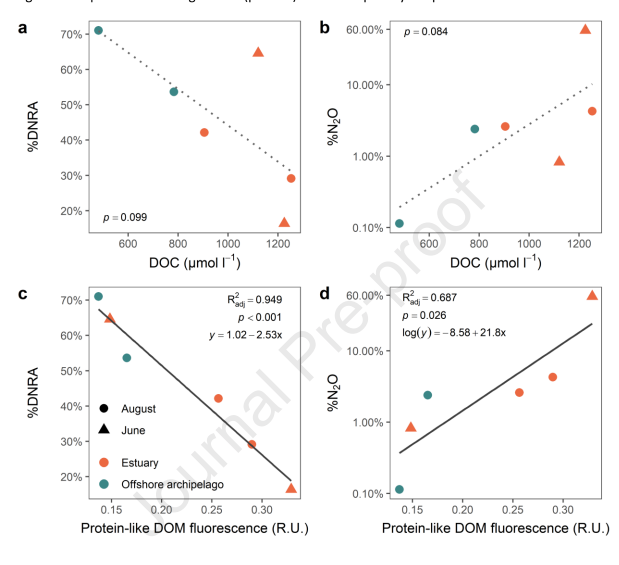


Figure 6. Relationships between dissolved organic carbon (DOC) and relative a) DNRA and b) N_2O production, and between bioavailable organic matter fraction (protein-like fluorescence; peak T) and relative c) DNRA and d) N_2O production at the estuary and offshore archipelago stations. The linear regression equations of the significant (p <0.05) relationships only are presented.



311 312

313 4 Discussion

314 Our results show that the dominant microbial nitrate reduction process switched from N-removing 315 denitrification to N-recycling DNRA when moving from the terrestrially-dominated estuary to 316 offshore archipelago region. This can be explained by changes in both DIN concentrations and 317 organic carbon bioavailability. As expected, nitrate concentrations were generally higher at the 318 estuarine than at the offshore archipelago stations, due to the diminishing impact of high-DIN 319 riverine water (Asmala et al., 2017). In parallel, we observed strong contrasts in the DOM 320 characteristics between estuary and offshore archipelago. High humic-like fluorescence, 321 humification index and SUVA₂₅₄ in porewater DOM at the estuarine stations indicate a pronounced 322 terrestrial contribution to the DOM pool (Asmala et al., 2013). These proxies suggest low DOM 323 bioavailability in these areas, while high S275-295 and BIX values at the offshore archipelago stations 324 indicate a higher contribution of recently produced autochthonous, likely more bioavailable DOM 325 (Lee et al., 2018). A similar gradient in the source of sedimentary particulate OM was observed by 326 Jilbert et al., (2018), where sedimentary N:C values of 0.05-0.06 (C:N of 17-20) observed in the 327 estuary indicated a higher contribution of terrestrially sourced material, while in the offshore region, 328 the N:C of 0.13-0.14 (C:N of 7-8) reflected the dominance of phytoplankton-derived material.

329 In previous studies, denitrification has been shown to decrease with decreasing water-column 330 nitrate concentrations in the coastal Baltic Sea (Asmala et al., 2017). Our data show that rates of all 331 nitrate reduction processes using water column nitrate (Dw N2, Dw N2O, DNRAw) decrease from 332 estuary to offshore archipelago (Suppl. Fig. 3). However, because total nitrate reduction was mainly 333 based on the nitrate provided through nitrification rather than water column nitrate, total nitrate 334 reduction rates ($\Sigma N_2 + \Sigma N_2 O + DNRA$) were not significantly different between estuary and offshore 335 archipelago stations. We suggest that the low amount of bioavailable carbon was limiting 336 denitrification in the estuary, whereas decreasing nitrate availability started to limit the process 337 offshore archipelago. The low bioavailable organic carbon-to-nitrate ratio at the estuarine stations 338 was reflected in the higher denitrification-to-DNRA ratio, whereas DNRA dominated nitrate 339 reduction under high bioavailable carbon-to-nitrate ratio at the offshore archipelago stations. A 340 preference of the sediment microbial community for DNRA under nitrate-limited conditions has previously been explained in terms of the efficiency with which DNRA makes use of nitrate as an 341 342 electron acceptor, with a higher rate of electron transfer per mole of N reduced despite the higher 343 free energy yield of denitrification (Algar and Vallino, 2014). Interestingly, the DOM characteristics 344 was directly related to N processes, while the amount of bulk organic carbon (as indicated by the 345 porewater DOC concentration) was not (Fig. 6). We acknowledge that several alternative factors may

346 influence rates and pathways of nitrate reduction processes in coastal sediments. For instance, the 347 presence of hydrogen sulfide (H₂S) close to the sediment-water interface promotes %DNRA 348 (Plummer et al., 2015). However, upper-sediment sulfide concentrations in the range of 1–3 mM are 349 required for a clear impact on N processes, while sulfide in the upper sediments of our study area 350 were consistently < 0.1 mM (Jilbert et al., 2018). These low concentrations result from the titrating 351 effect of sedimentary Fe oxides in the coastal Baltic Sea, suggesting that sulfide is a minor driver of the observed changes in %DNRA in our dataset. Furthermore, the presence of abundant Fe oxides 352 producing Fe²⁺, an alternative electron donor, may promote DNRA (Kessler et al., 2018; Robertson et 353 al., 2016). Again, our study area shows only mild enrichments of porewater Fe²⁺ in the upper 354 sediments (up to 0.2 mM, Jilbert et al., 2018) in comparison to the sites studied by Robertson et al., 355 356 (2016) (up to 0.8 mM), decreasing the potential significance of Fe. The anomalously high rates of 357 DNRA at P1 in June may however relate to porewater Fe²⁺, since this is the most Fe-rich of our sampling stations (see Station A in Jilbert et al., 2018). 358

359 In addition, our results demonstrate that the overall difference in potential organic carbon bioavailability between estuary and offshore archipelago regions is likely to influence the end-360 product of denitrification. At the near-shore estuarine stations, denitrification produced high 361 proportions of N₂O (1-58% of total nitrate reduction; 3-1230 μ M N m⁻² d⁻¹). This result implies that 362 363 nitrate was preferred over N₂O as an electron acceptor under conditions of high nitrate to 364 bioavailable carbon the nitrate-replete conditions of the estuary (Richardson et al., 2009), allowing N₂O to accumulate in bottom waters. In contrast, the share of N₂O in denitrification was lower in the 365 offshore archipelago stations (0.1-2%, 1-9 μ mol N m⁻² d⁻¹), where the bulk carbon concentrations 366 367 were low but the contribution of bioavailable autochthonous carbon to the carbon pool was high 368 and nitrate concentration low. In accordance, N₂O concentrations in the bulk water column samples 369 collected between 2015 and 2017 were higher at the estuarine stations than in the offshore 370 archipelago, agreeing with the previous results in coastal environments with high freshwater impact 371 and fluctuating environmental conditions (e.g. Foster and Fulweiler, 2016; Nielsen et al., 2009; 372 Silvennoinen et al., 2008). While part of the accumulated N₂O can originate from nitrification or 373 coupled nitrification-denitrification (Foster and Fulweiler, 2016), we measured rather equal rates of sediment nitrification at the estuary and offshore archipelago stations (estuary: 841±378, offshore 374 archipelago: 1089±193 μ mol N m⁻² d⁻¹; data not shown), arguing against an important role for 375 nitrification in N₂O production in the estuary. Furthermore, our data show that N₂O produced in 376 377 coupled nitrification-denitrification was mainly derived from denitrification. Although part of the 378 water-column N₂O pool in the estuary is likely advected with riverine water (Bange et al., 1998), the

majority appears to derive from sediment processes, since N_2O concentrations were generally higher in the bottom water than at the surface (Fig. 2).

381 Coastal systems are considered as important nutrient filters, reducing N loading from catchment 382 areas towards the open sea. Although our results confirm that the main N removal process in the 383 studied coastal environment is N₂-producing heterotrophic denitrification, they also highlight the 384 importance of N-recycling DNRA. In the outer offshore archipelago region with decreasing influence 385 of riverine water, DNRA can produce substantial amounts of bioavailable ammonium, enhancing the 386 N recycling between sediments and surface water, especially in summer with the highest 387 autochthonous biomass production and sedimentation. Intensifying eutrophication increases 388 bioavailable carbon availability through higher algal biomass production, which in turn may promote 389 DNRA and increase the role of estuaries as hotspots for N recycling, over N removal. This 390 phenomenon has already been observed in some eutrophied systems (Bernard et al., 2015; Song et 391 al., 2014), and could delay the recovery of water quality of the open sea in the Baltic Sea region.

The future role of eutrophic coastal systems as sources of N₂O to the atmosphere depends on the 392 393 balance of N processes in coastal sediments. In systems such as Pohjanpitäjänlahti, the DIN pool of 394 the estuary is dominated by nitrate, favouring production of N_2O during denitrification under 395 nitrate-replete conditions. Hence, further increases in nutrient loading to this system is likely to enhance N₂O-producing denitrification, especially under scenarios of increased annual runoff and 396 397 higher summer temperature, which will enhance stratification and hypoxia throughout the Baltic Sea 398 (Meier et al., 2011), contributing to the predicted rise in emissions of this greenhouse gas in the 399 future (Murray et al., 2015). Our results highlight the need to consider the intricate balance of 400 processes in the nitrogen cycle along coastal gradients, especially in relation to organic carbon 401 characteristics. Also their spatial variation and temporal evolution needs to be further clarified in 402 order to properly understand the role of coastal ecosystems as filters of land-to-sea transfer of N.

403 5 Acknowledgements

We are grateful to the technical staff of Tvärminne Zoological Station and the Ecosystems and Environment Research Program at University of Helsinki for assistance during fieldwork and laboratory analyses. This work was supported by the Academy of Finland (projects 267112, 309748, 310302, and 317684)

408 6 References

Algar, C.K., Vallino, J.J., 2014. Predicting microbial nitrate reduction pathways in coastal sediments.
Aquat. Microb. Ecol. 71, 223–238. https://doi.org/10.3354/ame01678

- 411 Asmala, E., Autio, R., Kaartokallio, H., Pitkänen, L., Stedmon, C.A., Thomas, D.N., 2013. Bioavailability
- 412 of riverine dissolved organic matter in three Baltic Sea estuaries and the effect of catchment
- 413 land use. Biogeosciences 10, 6969–6986. https://doi.org/10.5194/bg-10-6969-2013
- 414 Asmala, E., Carstensen, J., Conley, D.J., Slomp, C.P., Stadmark, J., Voss, M., 2017. Efficiency of the
- 415 coastal filter: Nitrogen and phosphorus removal in the Baltic Sea. Limnol. Oceanogr. 62, S222–
 416 S238. https://doi.org/10.1002/lno.10644
- Asmala, E., Haraguchi, L., Markager, S., Massicotte, P., Riemann, B., Staehr, P.A., Carstensen, J., 2018.
 Eutrophication Leads to Accumulation of Recalcitrant Autochthonous Organic Matter in Coastal
 Environment. Global Biogeochem. Cycles 32, 1673–1687.
- 420 https://doi.org/10.1029/2017GB005848
- 421 Bange, H.W., Dahlke, S., Ramesh, R., Meyer-Reil, L.A., Rapsomanikis, S., Andreae, M.O., 1998.
- 422 Seasonal study of methane and nitrous oxide in the coastal waters of the southern Baltic Sea.
- 423 Estuar. Coast. Shelf Sci. 47, 807–817. https://doi.org/10.1006/ecss.1998.0397
- Barnes, R.T., Smith, R.L., Aiken, G.R., 2012. Linkages between denitrification and dissolved organic
 matter quality, Boulder Creek watershed, Colorado. J. Geophys. Res. Biogeosciences 117, 1–14.
 https://doi.org/10.1029/2011JG001749
- Bernard, R.J., Mortazavi, B., Kleinhuizen, A.A., 2015. Dissimilatory nitrate reduction to ammonium
 (DNRA) seasonally dominates NO3- reduction pathways in an anthropogenically impacted subtropical coastal lagoon. Biogeochemistry 125, 47–64. https://doi.org/10.1007/s10533-0150111-6
- Bonaglia, S., Deutsch, B., Bartoli, M., Marchant, H.K., Brüchert, V., 2014. Seasonal oxygen, nitrogen
 and phosphorus benthic cycling along an impacted Baltic Sea estuary: Regulation and spatial
 patterns. Biogeochemistry 119, 139–160. https://doi.org/10.1007/s10533-014-9953-6
- 434 Bouwman, A.F., Bierkens, M.F.P., Griffioen, J., Hefting, M.M., Middelburg, J.J., Middelkoop, H.,
- 435 Slomp, C.P., 2013. Nutrient dynamics, transfer and retention along the aquatic continuum from
- 436 land to ocean: Towards integration of ecological and biogeochemical models. Biogeosciences
- 437 10, 1–23. https://doi.org/10.5194/bg-10-1-2013
- 438 Burdige, D.J., 2006. Geochemistry of Marine Sediments, 1st ed. Princeton Univ. Press, Princeton.
- 439 Carlson, H.K., Lui, L.M., Price, M.N., Kazakov, A.E., Carr, A. V., Kuehl, J. V., Owens, T.K., Nielsen, T.,
- Arkin, A.P., Deutschbauer, A.M., 2020. Selective carbon sources influence the end-products of
 microbial nitrate respiration. ISME J. https://doi.org/10.1038/s41396-020-0666-7

- 442 Christensen, P.B., Rysgaard, S., Sloth, N.P., Dalsgaard, T., Schwærter, S., 2000. Sediment
- 443 mineralization, nutrient fluxes, denitrification and dissimilatory nitrate reduction to ammonium
- in an estuarine fjord with sea cage trout farms. Aquat. Microb. Ecol. 21, 73–84.
- 445 https://doi.org/10.3354/ame021073
- 446 Coble, P.G., 1996. Characterization of marine and terrestrial DOM in seawater using excitation-
- 447 emission matrix spectroscopy. Mar. Chem. 51, 325–346. https://doi.org/10.1016/0304-
- 448 4203(95)00062-3
- Dong, L.F., Nedwell, D.B., Stott, A., 2006. Sources of nitrogen used for denitrification and nitrous
 oxide formation in sediments of the hypernutrified Colne, the nutrified Humber, and the
 oligotrophic Conwy estuaries, United Kingdom. Limnol. Oceanogr. 51, 545–557.
- 452 https://doi.org/10.4319/lo.2006.51.1_part_2.0545
- Fellman, J.B., Petrone, K.C., Grierson, P.F., 2011. Source, biogeochemical cycling, and fluorescence
 characteristics of dissolved organic matter in an agro-urban estuary. Limnol. Oceanogr. 56,
 243–256. https://doi.org/10.4319/lo.2011.56.1.0243
- Foster, S.Q., Fulweiler, R.W., 2016. Sediment nitrous oxide fluxes are dominated by uptake in a
 temperate estuary. Front. Mar. Sci. 3, 1–13. https://doi.org/10.3389/fmars.2016.00040
- 458 Giblin, A.E., Tobias, C.R., Song, B., Weston, N., Banta, G.T., Rivera-Monroy, V.H., 2013. The
- 459 importance of dissimilatory nitrate reduction to ammonium (DNRA) in the nitrogen cycle of
- 460 coastal ecosystems. Oceanography 26, 124–131. https://doi.org/10.5670/oceanog.2013.54
- 461 Goñi, M.A., Teixeira, M.J., Perkeya, D.W., 2003. Sources and distribution of organic matter in a river-
- dominated estuary (Winyah Bay, SC, USA). Estuar. Coast. Shelf Sci. 57, 1023–1048.
- 463 https://doi.org/10.1016/S0272-7714(03)00008-8
- 464 Hardison, A.K., Algar, C.K., Giblin, A.E., Rich, J.J., 2015. Influence of organic carbon and nitrate
- 465 loading on partitioning between dissimilatory nitrate reduction to ammonium (DNRA) and N2
- 466 production. Geochim. Cosmochim. Acta 164, 146–160.
- 467 https://doi.org/10.1016/j.gca.2015.04.049
- Heiskanen, A.S., Kononen, K., 1994. Sedimentation of vernal and late summer phytoplankton
 communities in the coastal Baltic Sea. Arch. fur Hydrobiol. 131, 175–198.
- 470 Hellemann, D., Tallberg, P., Aalto, S.L., Bartoli, M., Hietanen, S., 2020. Seasonal cycle of benthic
- 471 denitrification and DNRA in the aphotic coastal zone, northern Baltic Sea. Mar. Ecol. Prog. Ser.
 472 637, 15–28.

- 473 Hellemann, D., Tallberg, P., Bartl, I., Voss, M., Hietanen, S., 2017. Denitrification in an oligotrophic
- 474 estuary: A delayed sink for riverine nitrate. Mar. Ecol. Prog. Ser. 583, 63–80.
- 475 https://doi.org/10.3354/meps12359
- 476 Hietanen, S., 2007. Anaerobic ammonium oxidation (anammox) in sediments of the Gulf of Finland.
 477 Aquat. Microb. Ecol. 48, 197–205. https://doi.org/10.3354/ame048197
- 478 Hietanen, S., Kuparinen, J., 2008. Seasonal and short-term variation in denitrification and anammox
- 479 at a coastal station on the Gulf of Finland, Baltic Sea. Hydrobiologia 596, 67–77.

480 https://doi.org/10.1007/s10750-007-9058-5

- 481 Huguet, A., Vacher, L., Relexans, S., Saubusse, S., Froidefond, J.M., Parlanti, E., 2009. Properties of
- 482 fluorescent dissolved organic matter in the Gironde Estuary. Org. Geochem. 40, 706–719.

483 https://doi.org/10.1016/j.orggeochem.2009.03.002

- 484 Jilbert, T., Asmala, E., Schröder, C., Tiihonen, R., Myllykangas, J.P., Virtasalo, J.J., Kotilainen, A.,
- 485 Peltola, P., Ekholm, P., Hietanen, S., 2018. Impacts of flocculation on the distribution and
- 486 diagenesis of iron in boreal estuarine sediments. Biogeosciences 15, 1243–1271.
- 487 https://doi.org/10.5194/bg-15-1243-2018
- Kessler, A.J., Roberts, K.L., Bissett, A., Cook, P.L.M., 2018. Biogeochemical Controls on the Relative
 Importance of Denitrification and Dissimilatory Nitrate Reduction to Ammonium in Estuaries.
 Global Biogeochem. Cycles 32, 1045–1057. https://doi.org/10.1029/2018GB005908
- 491 Kraft, B., Tegetmeyer, H.E., Sharma, R., Klotz, M.G., Ferdelman, T.G., Hettich, R.L., Geelhoed, J.S.,
- 492 Strous, M., 2014. The environmental controls that govern the end product of bacterial nitrate 493 respiration. Science (80-.). 345, 676–679. https://doi.org/10.1126/science.1254070
- Lee, M.H., Osburn, C.L., Shin, K.H., Hur, J., 2018. New insight into the applicability of spectroscopic
- 495 indices for dissolved organic matter (DOM) source discrimination in aquatic systems affected
- 496 by biogeochemical processes. Water Res. 147, 164–176.
- 497 https://doi.org/10.1016/j.watres.2018.09.048
- 498 Malve, O., Virtanen, M., Villa, L., Karonen, M., Aakerla, H., Heiskanen, A.S., Lappalainen, K.M.,
- 499 Holmberg, R., 2000. Artificial oxygenation experiment in hypolimnion of Pojo Bay estuary in
- 500 1995 and 1996: Factors regulating estuary circulation and oxygen and salt balances. Finnish
- 501 Environ. 377, 1-163 (In Finnish with English summary).
- Massicotte, P., 2018. eemR: Tools for Pre-Processing Emission-Excitation-Matrix (EEM) Fluorescence
 Data. R package version 1.0.1. https://CRAN.R-project.org/package=eemR.

- Mattsson, T., Kortelainen, P., Räike, A., 2005. Export of DOM from boreal catchments: Impacts of
 land use cover and climate. Biogeochemistry 76, 373–394. https://doi.org/10.1007/s10533 005-6897-x
- Meier, H.E.M., Andersson, H.C., Eilola, K., Gustafsson, B.G., Kuznetsov, I., Mller-Karulis, B., Neumann,
 T., Savchuk, O.P., 2011. Hypoxia in future climates: A model ensemble study for the Baltic Sea.
 Geophys. Res. Lett. 38, 1–6. https://doi.org/10.1029/2011GL049929
- Murphy, K.R., Stedmon, C.A., Waite, T.D., Ruiz, G.M., 2008. Distinguishing between terrestrial and
 autochthonous organic matter sources in marine environments using fluorescence
- 512 spectroscopy. Mar. Chem. 108, 40–58. https://doi.org/10.1016/j.marchem.2007.10.003
- Murray, R.H., Erler, D. V., Eyre, B.D., 2015. Nitrous oxide fluxes in estuarine environments: Response
 to global change. Glob. Chang. Biol. 21, 3219–3245. https://doi.org/10.1111/gcb.12923
- Myllykangas, J.P., Jilbert, T., Jakobs, G., Rehder, G., Werner, J., Hietanen, S., 2017. Effects of the 2014
 major Baltic inflow on methane and nitrous oxide dynamics in the water column of the central
- 517 Baltic Sea. Earth Syst. Dyn. 8, 817–826. https://doi.org/10.5194/esd-8-817-2017
- Nielsen, L.P., 1992. Denitrification in sediment determined from nitrogen isotope pairing. FEMS
 Microbiol. Lett. 86, 357–362. https://doi.org/10.1111/j.1574-6968.1992.tb04828.x
- 520 Nielsen, M., Gieseke, A., De Beer, D., Revsbech, N.P., 2009. Nitrate, nitrite, and nitrous oxide
- 521 transformations in sediments along a salinity gradient in the Weser Estuary. Aquat. Microb.
- 522 Ecol. 55, 39–52. https://doi.org/10.3354/ame01275
- 523 Plummer, P., Tobias, C., Cady, D., 2015. Nitrogen reduction pathways in estuarine sediments:
- 524 Influences of organic carbon and sulfide. J. Geophys. Res. Biogeosciences 120, 1958–1972.
- 525 https://doi.org/10.1002/2015JG003004.Received
- 526 R Core Team, 2020. R: A language and environment for statistical computing.
- Räike, A., Kortelainen, P., Mattsson, T., Thomas, D.N., 2012. 36year trends in dissolved organic
 carbon export from Finnish rivers to the Baltic Sea. Sci. Total Environ. 435–436, 188–201.
- 529 https://doi.org/10.1016/j.scitotenv.2012.06.111
- 530 Richardson, D., Felgate, H., Watmough, N., Thomson, A., Baggs, E., 2009. Mitigating release of the
- 531 potent greenhouse gas N2O from the nitrogen cycle could enzymic regulation hold the key?
- 532 Trends Biotechnol. 27, 388–397. https://doi.org/10.1016/j.tibtech.2009.03.009
- 533 Risgaard-Petersen, N., Revsbech, N.P., Rysgaard, S., 1995. Combined microdiffusion-hypobromite

- oxidation method for determining nitrogen-15 isotope in ammonium. Soil Sci. Soc. Am. J. 59,
 1077–1080.
- Robertson, E.K., Roberts, K.L., Burdorf, L.D.W., Cook, P., Thamdrup, B., 2016. Dissimilatory nitrate
 reduction to ammonium coupled to Fe(II) oxidation in sediments of a periodically hypoxic
 estuary. Limnol. Oceanogr. 61, 365–381. https://doi.org/10.1002/lno.10220
- 539 Silvennoinen, H., Liikanen, A., Torssonen, J., Stange, C.F., Martikainen, P.J., 2008. Denitrification and
- 540 N2O effluxes in the Bothnian Bay (northern Baltic Sea) river sediments as affected by
- 541 temperature under different oxygen concentrations. Biogeochemistry 88, 63–72.
- 542 https://doi.org/10.1007/s10533-008-9194-7
- 543 Song, B., Lisa, J.A., Tobias, C.R., 2014. Linking DNRA community structure and activity in a shallow
- 544 lagoonal estuarine system. Front. Microbiol. 5, 1–10.
- 545 https://doi.org/10.3389/fmicb.2014.00460
- 546 Spencer, R.G.M., Ahad, J.M.E., Baker, A., Cowie, G.L., Ganeshram, R., Upstill-Goddard, R.C., Uher, G.,
- 547 2007. The estuarine mixing behaviour of peatland derived dissolved organic carbon and its
- 548 relationship to chromophoric dissolved organic matter in two North Sea estuaries (U.K.).
- 549 Estuar. Coast. Shelf Sci. 74, 131–144. https://doi.org/10.1016/j.ecss.2007.03.032
- 550 Stelzer, R.S., Thad Scott, J., Bartsch, L.A., Parr, T.B., 2014. Particulate organic matter quality
- 551 influences nitrate retention and denitrification in stream sediments: Evidence from a carbon
- 552 burial experiment. Biogeochemistry 119, 387–402. https://doi.org/10.1007/s10533-014-9975-0
- 553 Thamdrup, B., Dalsgaard, T., 2002. Production of N2 through anaerobic ammonium oxidation
- coupled to nitrate reduction in marine sediments. Appl. Environ. Microbiol. 68, 1312–1318.
 https://doi.org/10.1128/AEM.68.3.1312
- 556 Zhao, Y., Xia, Y., Li, B., Yan, X., 2014. Influence of environmental factors on net N2 and N2O
- 557 production in sediment of freshwater rivers. Environ. Sci. Pollut. Res. 21, 9973–9982.
- 558 https://doi.org/10.1007/s11356-014-2908-6
- 559 Zsolnay, A., Baigar, E., Jimenez, M., Steinweg, B., Saccomandi, F., 1999. Differentiating with
- 560 fluorescence spectroscopy the sources of dissolved organic matter in soils subjected to drying.
- 561 Chemosphere 38, 45–50. https://doi.org/10.1016/S0045-6535(98)00166-0
- 562

Highlights

- The availability of bioavailable carbon defines nitrate reduction end-product
- Estuaries with low bioavailable organic carbon can release high amounts of N2O
- Nitrogen is recycled through DNRA in the archipelago areas

ournal pre-proof

Aalto et al.

Author statement

Sanni L. Aalto: Methodology, Formal analysis, Investigation, Writing - Original Draft, Writing - Review & Editing, Visualization

Eero Asmala: Methodology, Formal analysis, Investigation, Writing - Review & Editing, Visualization

Tom Jilbert: Methodology, Investigation, Resources, Writing - Review & Editing, Visualization

Susanna Hietanen: Conceptualization, Methodology, Investigation, Resources, Supervision, Funding acquisition

Declaration of interests

 \boxtimes The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

□The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: