Riverine anaerobic ammonium oxidation across contrasting geologies

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Rivers are an important global sink for excess bioavailable nitrogen: they convert approximately 40% of terrestrial N-runoff per year (~47 Tg) to biologically unavailable N₂ gas and return it to the atmosphere.⁠¹ Currently, riverine N₂ production is conceptualised and modelled as denitrification.²⁻⁴ The contribution of anaerobic ammonium oxidation (or anammox), an alternate pathway of N₂ production important in marine environments, is not well understood.⁵,⁶ Here we use in situ and laboratory measurements of anammox activity using ¹⁵N tracers and molecular analyses of microbial communities to evaluate anammox in clay, sand, and chalk-dominated river beds in the Hampshire Avon catchment, UK during summer, 2013. Anammox hzo gene abundance varied across the contrasting geologies. Anammox rates were similar across geologies but contributed different proportions of N₂ production because of variation in denitrification rates. In spite of requiring anoxic conditions, anammox, most likely coupled to partial nitrification, contributed up to 58% of in situ N₂ production in oxic, permeable riverbeds. In contrast, denitrification dominated in low permeability clay-bed rivers, where anammox contributes roughly 7% to the production of N₂ gas. We conclude that anammox can represent an important nitrogen loss pathway in permeable river sediments.

Humans have greatly altered the global nitrogen cycle through industrial N₂ fixation and application of this fixed-N to the land, disturbing the balance between N₂ fixation and N₂ production.¹ Almost half of global terrestrial N₂ production occurs within freshwaters (rivers, lakes, groundwater)² which to date has been conceptualised as a simple function of labile organic matter availability, i.e. canonical denitrification.⁴,⁷ Anammox alters the fundamental stoichiometry of the complete mineralisation of organic matter as, for every mole of organic-N converted to N₂, only half of the N-bearing compounds need partial oxidation to nitrite, and for each mole of nitrate/nitrite (NOₓ⁻) reduced, one mole of more bioavailable
ammonium is also removed.\textsuperscript{8} Rivers may not appear the most suitable environments for anammox – labile carbon,\textsuperscript{9} supplied from the catchment, and variable redox environments\textsuperscript{10} in the sediments, should, in theory, favour heterotrophic denitrification by facultative anaerobes.\textsuperscript{11,12} If active however, anammox alters our perception of how riverbeds function and increases a river’s capacity to attenuate nitrogen.

Much of what is known about anammox in the environment comes from estuaries and coastal seas where anammox varies in response to sediment reactivity. The relative contribution of anammox to marine N\textsubscript{2} production (\textit{ra}) decreases with proximity to the shore as supply of carbon stimulates denitrification over anammox.\textsuperscript{12,13} Extrapolating this trend further inshore suggested anammox activity would be insignificant in estuaries but anammox potential actually increased.\textsuperscript{14,15} In both estuaries and coastal seas, however, anammox is important in low permeability sediments (\textit{ra} $<$1 to 11 \%)\textsuperscript{9,16}, where oxygen penetration is restricted\textsuperscript{12,15} and it is these muddy sediments that the few studies of riverine anammox have occurred.\textsuperscript{5,6} In addition, anammox is widespread in marine sediments but the affiliated bacteria are phylogenetically constrained. In contrast, freshwater environments have been shown to possess highest anammox diversity, purportedly containing many novel anammox bacteria.\textsuperscript{17}

The geology of the Hampshire Avon catchment (United Kingdom) is dominated by permeable chalk from the Upper Chalk formation underlain by less permeable Upper Greensand and smaller outcrops of impermeable Gault clay (see Supplementary Information). Using a combination of \textit{in situ} and laboratory-based\textsuperscript{15}N tracer techniques\textsuperscript{12,18} and molecular assays we characterised both the anammox community and its activity within rivers from clay, sand and chalk-dominated sub-catchments under summer, base flow conditions (Table S1). For rivers in which \textit{in situ} measurements were performed we indexed catchment permeability by calculating the base-flow index (BFI, Table S2), the proportion of river flow...
derived from deep groundwater sources. In clay catchments, low soil permeability leads to routing of rainfall overland or through shallow, more permeable soils into the river (low BFI). Whilst in chalk or sand catchments, the higher soil permeability allows infiltrated water to percolate deeper into the aquifer and follow much longer flow paths to towards the river (high BFI).

We began by characterising the anammox hzo functional gene that encodes hydrazine oxidoreductase which catalyses the oxidation of hydrazine to N₂. The hzo gene was detected in all sediments confirming that anammox bacteria were present (Table S3). Anammox activity was then confirmed by production of ²⁹N₂ following addition of ¹⁵NH₄⁺ and ¹⁴NO₃⁻ to anoxic sediment slurries (Table S4). We can attribute this oxidation of ammonium to anammox rather than reduction of metal oxides,¹⁹ for example, as no ¹⁵N-N₂ was produced in ¹⁵NH₄⁺ only controls (Table S4). Anammox potential varied across the riverine gradient with fastest rates and greatest anammox contribution to N₂ production observed in the permeable sands and chalk-gravels (Figure 1a, see Table S5). Anammox potential was also positively correlated with hzo gene abundance ($r_s (7) = 0.867, p = 0.005$). The absolute abundance of the hzo gene was significantly higher in chalk-gravels ($F_{(2,6)}=8.64; p=0.017$) and the proportion of hzo to 16S rRNA was even greater (Table S3), given that 16S rRNA copies were highest in the clays.

The anammox functional hzo gene was sequenced and phylogenetic analysis revealed four clades (Clade I-IV) that differed in their relative distributions between the three geologies. In general, there was a broad diversity of hzo sequences that were distinct from known hzo sequences (Figure S1a; Table S6; Supplementary discussion 1). In addition, we sequenced the 16S rRNA gene, and clustered 951,000 sequences into 28,000 OTUs. All Planctomycete sequences represented only 0.5-0.9% of the total 16S rRNA sequences, yet none were assigned to anammox genera (RDP classifier), or grouped phylogenetically with
any of the currently known anammox bacteria suggesting that anammox bacteria were not
detected due to their low relative abundance in the bacterial communities (Table S3).

Somewhat surprisingly, $^{15}$N-$\text{N}_2$ production was not limited to anoxic slurries as we
also measured $^{15}$N-$\text{N}_2$ production after addition of $^{15}\text{NH}_4^+$ to slurries with an air headspace
(i.e. $\text{O}_2$ saturated; Figure 2, Figure S2). Sediments at the start of the incubation contained
considerable $^{14}\text{NH}_4^+$ but little $^{14}\text{NO}_\chi^-$ (e.g. 350-960 $\mu$M and $\leq 3$ $\mu$M, respectively; Table S7)
which, in combination with $^{15}\text{NH}_4^+$, could result in $^{29}\text{N}_2$ and $^{30}\text{N}_2$ through either anammox or
denitrification (or both) coupled to nitrification. As production of $^{15}$N-$\text{N}_2$ happened
immediately upon addition of $^{15}\text{NH}_4^+$ (Figure 2), nitrification must be rapid and the coupling
to pathway(s) of $\text{N}_2$ production very tight. The potential for aerobic nitrification to fuel
anammox has been demonstrated in oceanic waters with no measurable oxygen$^{20}$ and inferred
in riparian soils.$^{21}$ Here we confirmed nitrifications’ direct involvement in oxic $\text{N}_2$ production
in sediments by addition of allylthiourea, an inhibitor of aerobic ammonium oxidation, which
turned off $\text{N}_2$ production completely (Figure 2).

To apportion oxic $\text{N}_2$ production to coupled nitrification-anammox or nitrification-
denitrification we modelled the distribution of isotopes within the $\text{N}_2$ produced via either
pathway (see Methods). Despite clear evidence of nitrification within the slurries (see above),
the majority of samples (31 of 45 incubations) did not have measurable $^{15}\text{NO}_\chi^-$ after the
$^{15}\text{NH}_4^+$ addition (Table S7) and, therefore, coupling between nitrification and $\text{N}_2$ production
was 100 % efficient. Without measurable $\text{NO}_\chi^-$ we can only assume that nitrification and $\text{N}_2$
production are so closely affiliated that the $^{15}$N-content of the $\text{NO}_\chi^-$ and $\text{NH}_4^+$ pools are equal
which, by definition, prevents separation of anammox from denitrification.$^{22}$ In the remaining
incubations ($n = 14$; 5 and 9 for chalk-gravel and sand, respectively), $^{15}\text{NO}_\chi^-$ was detectable
and the $^{15}$N-labelling of the $\text{NO}_\chi^-$ and $\text{NH}_4^+$ pools was different. Production of $^{15}$N-$\text{N}_2$ within
the $^{15}\text{NO}_\chi^-$-bearing subset of incubations was representative of the entire dataset (Figure S2);
although no $^{15}$NO$_2^-$ was detected within any clays ($n = 15$ time series, consisting of 75 discrete sediment samples).

We could apportion the production of N$_2$ gas to either anammox or denitrification in some of the $^{15}$NO$_2^-$-containing experiments ($n = 8$ of 14), with 38±2 % and 65±15 % of potential N$_2$ production occurring via anammox in the chalk-gravels and sands, respectively (mean ± 1 s.e.; Figure 2). Within the remaining $^{15}$NO$_2^-$-bearing experiments ($n = 6$ of 14), the N$_2$ pool was more enriched in $^{15}$N than could be explained by either denitrification or anammox (deviation between measured and predicted $^{15}$N-content of produced N$_2$ was 26±4 %, mean ± 1 s.e.; Table S8). This observation violates the assumption of a random combination of isotopes which is fundamental to $^{15}$N-assays and, for the N$_2$ to be more enriched in $^{15}$N than predicted, suggests heterogeneity in the $^{15}$N labelling of substrate pool(s). Here, the heterogeneity probably exists because the NO$_2^-$ pool actually being reduced is partially physically isolated from the bulk NO$_2^-$ pool, further supporting a tight coupling of nitrification to N$_2$ production. Heterogeneity in the NH$_4^+$ pool is less likely as ammonium was plentiful – both as ambient $^{14}$NH$_4^+$ (Table S7) and added $^{15}$NH$_4^+$ (500 µM 98 % $^{15}$N).

Supply of nitrite rather than ammonium has been suggested as the limiting factor for anammox in aquatic sediments – potentially coupling anammox to either nitrification and/or denitrification.$^{15}$ In anoxic marine and estuarine sediments, anammox can be fuelled by denitrification-derived nitrite,$^{15}$ even forming a symbiotic relationship with some nitrate reducing / sulphur oxidising bacteria.$^{24}$ Association between nitrifiers and anammox bacteria may be weak in low permeability sediments (clays, estuarine mud) because much of the nitrite produced in the upper few millimetres of the bed can diffuse into the overlying water or be fully oxidised to nitrate before reaching the sub-oxic layer.$^{15}$ In the presence of oxygen however, affinity between nitrifiers and anammox bacteria can exist in aggregates and is
indeed the fundamental principle of CANON waste-water treatment reactors operating at reduced oxygen.\textsuperscript{25} The ability for anammox bacteria to couple to both aerobic and anaerobic pathways of nitrite production could be very advantageous in permeable riverbeds, where groundwater-surface exchange facilitates advective transport of solutes,\textsuperscript{26} creating a mosaic of redox micro-environments\textsuperscript{27} within which both nitrification and denitrification can occur.\textsuperscript{18}

Rates of \textit{in situ} anammox activity were similar across riverbeds of differing geology (Figure 1b), despite large differences in porewater oxygen – clays were essentially anoxic whilst oxygen was present in both the sand and chalk-gravels (8 to 110 \% of air-equilibration; mean ± 1 s.e. = 134±14 µM; Figure S3). In contrast, anammox did make a markedly higher contribution to N\textsubscript{2} production in the permeable sediments compared to the clays (Figure 1b, Table S5); not because anammox activity increased, rather that denitrification activity declined (Table S9). Similar increases in the significance of anammox at the expense of denitrification are well documented in marine sediments.\textsuperscript{12} Differences in \textit{ra} between clays, sands and chalk-gravels were consistent across seasons (Figure S5) and are related to the chemical gradient inherent to porewaters of these different riverbed types. Within clays, porewaters were typically reduced (high in ammonium, iron (II) and phosphate) whereas chalk-gravel porewaters were more oxidised (high in nitrate, intermediate in O\textsubscript{2}; Figure S4) with \textit{ra} increasing as porewaters become more oxidised (Figure 3a; \textit{r}, (38) = -0.73, \(p<0.001\)); hinting at a potential coupling between anammox and nitrification. Anammox activity was strongly associated with nitrite and O\textsubscript{2}, increasing as nitrite accumulates in partially oxygenated sediments (Figure 3b) – again mirroring the coupling between partial nitrification and anammox in aggregates as exploited in CANON reactors.\textsuperscript{25} In contrast, anammox activity had essentially no association with ammonium, as where ammonium accumulates in these riverine sediments, labile organic carbon (the source of the ammonium) must also be plentiful (Figure 3b), fuelling denitrification at the expense of anammox.\textsuperscript{12,13} Overall we found a very
strong increase in the contribution of anammox to N\textsubscript{2} production and hzo abundance (as a fraction of total bacteria, hzo:16S rRNA sequences) with increasing BFI ($r_s (6) = 1.0, p = 0.003$ and $r_s (6) = 1.0, p < 0.001$ for anammox contribution and hzo abundance, respectively; Figure 3c), suggesting, in the long-term, that anammox is favoured with stable conditions (nutrients, temperature, pH).\textsuperscript{11}

Here, we have shown how anammox is making a significant contribution to the removal of fixed-N in oxic, permeable riverbeds; a pattern completely at odds with current knowledge. Supply of nitrite to anammox from partial nitrification removes the stoichiometric constraint of denitrification-anammox coupling ($ra \leq 29\%$),\textsuperscript{28} allowing anammox to potentially be as important an N-sink as denitrification in permeable riverbeds (maximum in situ $ra = 58\%$, median $= 37\%$ for chalk-gravels; Figure 3a). In the clays, anammox proceeds as per muddy estuarine sediments, making only a minor contribution to N\textsubscript{2} production ($ra \leq 7\%$) and being fuelled by canonical denitrification.\textsuperscript{15} It is important to appreciate that the stoichiometry of anammox, requiring only partial oxidation of some N-substrates, increases the efficiency of rivers to remove fixed N as both NO\textsubscript{x}\textsuperscript{-} and ammonium, changing our understanding of the ecosystem services they provide.

**Data sources**

URL and DOI for activity data to be provided once deposition into the Environmental Information Data Centre is complete. hzo gene sequences from this study are deposited in GenBank (NCBI) under the accession numbers ---

**References**


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Author contributions

MT with CMH, AB and KL conceived the original project. KL performed \(^{15}\)N-related work and with MT interpreted the process data and drafted the original manuscript. BAM designed and performed all the molecular work and phylogenetic analysis. AJD constructed the bioinformatic pipeline and performed the NGS analysis. CW directed the molecular component of project. LO assisted with fieldwork and performed sediment characterisation. AB and CMH performed hydrologic measurements and calculated base-flow indices. All authors contributed to writing the paper and approved the final manuscript.

Competing financial interests

The authors declare no competing financial interests.
Figure Captions

**Figure 1:** Anammox activity, both rate and contribution to N$_2$ production ($ra$), differs across a riverine gradient. Activity was measured as total potential in anoxic slurries (a) and ambient rates by direct, *in situ* measurements (*see* Supplementary Methods) (b). Grey bars indicate significant differences between groups. Data are mean values ±1 standard error ($n = 5$ and 10 for a and b, respectively).

**Figure 2:** Production of $^{15}$N-labelled N$_2$ following addition of $^{15}$NH$_4^+$ to chalk-gravels (a) and sand (b) incubated under air-saturated conditions. Notice that in the presence of allylthiourea (white circles), an inhibitor of nitrification, there was no production of $^{15}$N-N$_2$ but without the inhibitor (black circles) there was immediate conversion of $^{15}$NH$_4^+$ to $^{15}$N-N$_2$ confirming the tight coupling between nitrification and N$_2$ production. Grey boxes are dissolved O$_2$. The pathways of N$_2$ production (anammox or denitrification, red versus white column sections, respectively) were determined via modelling (*see* Supplementary Methods). Data are mean values ±1 standard error ($n = 5$).

**Figure 3:** Anammox varies with both patch scale and sub-catchment river characteristics. Differences in anammox contribution to *in situ* N$_2$ production within clays (red, $n=10$), sands (blue, $n=20$) and chalk-gravels (yellow, $n=20$) result from fine-scale chemical variation (a) (lower scores are more oxidised porewaters, *see* Figure S4). Anammox activity is most strongly associated with nitrite and O$_2$ (b). Squares represent geology averages in the redundancy-analysis triplot. At the sub-catchment scale, both anammox contribution and bacterial abundance ($hzo$ copy number: total bacteria) increase markedly with increasing base-flow index (c). Data are means ±1 standard error ($n = 3$ for $hzo$ fractions).
Methods

**Measurement of potential anammox activity**

**Collection of sediment.** Sediment was collected from nine rivers in the Hampshire Avon catchment in summer 2013 (19-20 August) under base flow conditions. Rivers were in sub-catchments of predominantly clay, sand or chalk (n = 3 per geology). Full site descriptions are provided as supplementary information. At each river, surficial sediments (<5 cm) were removed from five un-vegetated patches of the main channel by hand with Perspex cores (internal diameter = 9 cm). After sediment disturbed within the core settled, the overlying water was gently decanted and sediment for activity measurements was placed in ziplock bags and stored at ~4 °C.

**Preparation of slurries.** In the laboratory, each bag of sediment was homogenised by gentle stirring and particles > 9 mm (the internal diameter of the vials) were removed. Sediments were then placed in pre-weighed gas-tight vials (Exetainer, Labco) with replicates from the rivers treated as discrete samples. Slurries were prepared with synthetic river water (see supplementary information) in a 1:1 sediment-to-water ratio.

The potential for anammox and denitrification was measured in anoxic slurries \(^{29-31}\) (oxygen-free \(N_2\) headspace, British Oxygen Company) prepared with de-oxygenated synthetic river water in an anoxic hood (CV24, Belle Technologies). Anoxic slurries were pre-incubated in the dark on an orbital shaker (80 r.p.m., Stuart SSL1) for at least 18 hours to remove any ambient \(^{14}NO_x^-\). \(^{15}N\) tracers (100 µL, de-oxygenated) were injected through the septa of the vials in the following combinations: \(^{15}NH_4^+\) only, \(^{15}NH_4^+\) and \(^{14}NO_3^-\) and \(^{15}NO_3^-\) only. All \(^{15}N\)-salts were 98 atom % \(^{15}N\) (Sigma-Aldrich). Tracers increased ammonium concentrations by 500 µM and nitrate concentrations to 100 or 300 µM in the clay and permeable sediments, respectively. Each sediment and treatment combination consisted of a reference (no tracer added) and a killed control (100 µL 7M ZnCl\(_2\) injected prior to the
tracer). The $^{15}$NO$_3^-$ treatment consisted of 5 additional slurries per sample which were incubated for 0.5, 1, 2, 3 and 6 h on an orbital shaker (as above). The $^{15}$NH$_4^+$ and $^{15}$NH$_4^+$ and $^{14}$NO$_3^-$ treatments were end point only experiments with 1 additional slurry per sample (i.e. $T_{final}$) incubated for 6 h. At the end of the incubation period, biological activity was stopped by injection of ZnCl$_2$ (as above) and gas-tight vials were stored upside down until analysis. Once headspace analysis was complete (see below) vials were opened and water extracted after centrifugation. Sediment was re-suspended twice with ultrapure water (same volume as aqueous phase of slurry) and the supernatant reserved. Water samples were filtered (0.45 µm polypropylene, Gilson Scientific) into plastic tubes (polypropylene, VWR International) and frozen until analysis (see below). The mass of sediment within each vial was then determined after sediments had been dried.

A parallel set of oxic slurries (air headspace) were prepared to investigate nitrification potential by addition of air-equilibrated synthetic river water to sediments on the lab bench with no pre-incubation. Air-equilibrated $^{15}$NH$_4^+$ tracer was injected through the septa of the vials (as above) and slurries were treated as per the anoxic $^{15}$NO$_3^-$ treatment. A second experiment was devised to examine if N$_2$ production observed in the oxic slurries was linked to nitrification (Rivers Ebble and Nadder only, 8/11/2013, sampling procedures as above). In addition to the $^{15}$NH$_4^+$ treatment, a second treatment containing both $^{15}$NH$_4^+$ and allylthiourea (concentration in slurry = 100 µM), a nitrification inhibitor, was also included. To determine if oxygen depletion occurred within the oxic slurries a set of scaled-up slurries were prepared in 20 mL gas-tight vials (Chromacol). At each time point the vial was opened and the dissolved O$_2$ concentration of the slurry was measured by inserting a calibrated, fast response micro-electrode (50 µm, Unisense).
Measurement of in situ anammox activity

Impermeable sediments. Ambient rates of anammox and denitrification were estimated in unvegetated clays by incubation of $^{15}$NO$_3^-$ in intact sediment cores (Perspex cores with rubber bungs, internal diameter = 3.4 cm, experiments performed between 03/08 and 10/08/2013).

Cores were collected by hand from the River Sem ($n=34$) and a tributary of the River Sem (Clay 2 in Table S1, $n=29$) and incubated on site in a tank full of river water. The amount of $^{15}$NO$_3^-$ added and the duration of the incubation varied between cores, ranging from 0.05 to 2.5 mL of 78 mM $^{15}$NO$_3^-$ (98 atom % $^{15}$N) and 31 to 252 minutes respectively. The range in $^{15}$NO$_3^-$ amendments aided separation of anammox from denitrification and different incubation times were used to verify $^{15}$N-N$_2$ production was linear. Following $^{15}$NO$_3^-$ injection into the overlying water column, cores were immediately capped with a bung fitted with a magnetic stirrer and placed in the incubation tank. The overlying water column was gently stirred to prevent stratification and light was excluded from the incubation tank. At the end of the incubation the bung was removed from the core and a water sample was quickly withdrawn with a syringe (polypropylene, BD Plastipak). The core was then homogenised by gentle stirring and decanted into a gas-tight vial (12 mL Exetainer, Labco) which was allowed to overflow before being capped. Biological activity was stopped by injection of 100 µL of formaldehyde through the septum. The water sample was then filtered and frozen (as above) until later analysis. Four additional sediment cores were retrieved on each day of fieldwork and a water sample and slurried sample collected (as above) to determine ambient $^{15}$N-N$_2$ concentrations.

Ambient chemistry within the clays was determined on porewaters recovered using rhizon samplers (0.2 µm mesh, 10 cm screen, Rhizosphere) inserted into the riverbed. Rhizon samplers were allowed 24 h to pre-equilibrate before porewater was extracted by applying a vacuum to the rhizon sampler via a syringe held open with a spacer bar. Water
samples for nutrient analysis were processed as described above. Water samples for iron (II) determination were preserved by dispensing porewater directly into a solution of buffered phenanthroline\textsuperscript{35} (3.5:1 1M pH 4.5 Acetate buffer: 0.2 % (w/v) 1-10-phenanthroline monohydrate). The dissolved O\textsubscript{2} concentration of the recovered porewater was measured by placing the O\textsubscript{2} microelectrode (as above) into an empty syringe barrel and gently transferring porewater into the vessel with a 2-way valve.\textsuperscript{36} We estimate that sample collection and transfer adds approximately 10 µM O\textsubscript{2} to the actual dissolved O\textsubscript{2} concentration and corrected all values accordingly. Following measurement of dissolved O\textsubscript{2} the pH was determined (pH100, VWR International). Additional sediment cores (Perspex, 9 cm diameter) were collected and transported back to the laboratory for fine scale oxygen profiling using a Clark-type oxygen microsensor (OX50, Unisense) within an automated micromanipulator controlled by microprofiling software (SensorTracer PRO, Unisense). Readings from the microelectrode were displayed on a picoammeter (PA 2000; Unisense) and logged after 4 s when the signal had stabilized.

Permeable sediments. Ambient rates of anammox and denitrification were estimated in unvegetated sediments of the Rivers Ebble, Wylye, Nadder and Avon (“Sand 2” in Table S1) by injection of \textsuperscript{15}NO\textsubscript{3}\textsuperscript{-} into the riverbed and collection of samples over time (i.e. “push-pull” sampling; sampling occurred between 31/07 and 15/08/2013). Ten bespoke stainless steel mini-probes were installed between 4 and 20 cm depth in the bed of the main channel on the day prior to the injection. We modified the system from previous work\textsuperscript{36} by attaching an extension (1 m length of 0.75 mm internal diameter Polyetheretherkeytone (PEEK) tubing, Polyflon Technology Ltd.) to the luer connector of the mini-probe to improve speed of sampling. Prior to the injection of \textsuperscript{15}NO\textsubscript{3}\textsuperscript{-}, porewater (15 mL total) was withdrawn from each mini-probe for dissolved O\textsubscript{2} and pH measurement, nutrient and iron (II) analysis and natural
abundance $^{15}$N-$\text{N}_2$ (as above). Gas samples were collected in 3 mL gas-tight vials and
poisoned with ZnCl$_2$ (25 µL, as above).

A tracer solution consisting of 300 µM $^{15}$NO$_3^-$ (98 atom % $^{15}$N) in a synthetic river
water/ KCl (4 mM) matrix, was de-oxygenated (as above) and 25 mL aliquots were drawn
into luer-lock syringes. Tracer was injected into the riverbed via the mini-probes, with each
injection lasting ~20 seconds. Porewater was recovered from each mini-probe immediately
post injection and a dissolved gas and water sample was collected (as above). Porewater was
then recovered at ~5, 10 and 30 minutes post injection and sampled as above.

Analytical methods for activity measurements

Nitrate (Limit of detection (LOD) 0.4 µM, precision 1 %), nitrite (LOD 0.1 µM, precision
1 %), ammonium (LOD 0.8 µM, precision 3 %) and soluble reactive phosphate (SRP, LOD
0.1 µM, precision 1 %) were quantified by automated colorimetric analysis using standard
methods (San++, Skalar). Iron(II) concentrations were quantified on samples preserved with
buffered phenanthroline$^{35}$ by absorbance measurement at 520 nm on a UV/Visible
spectrophotometer (LOD 1 µM, precision 1 %; Evolution 100, Thermo Fisher). The dissolved
oxygen electrode was calibrated with a zero solution (0.1 M sodium ascorbate in 0.1 M
sodium hydroxide) and 100% air-equilibrated water (laboratory measurements) or river water
(field-based measurements), the dissolved O$_2$ concentration of which was later determined by
Winkler titration. Samples for $^{15}$N-$\text{N}_2$ quantification that did not contain a headspace were
prepared for analysis by addition of helium (commercially pure grade, British Oxygen
Company) with a syringe and a two-way valve (0.5 or 2 mL headspaces were added to
porewater and slurred core samples, respectively) and were equilibrated at 22 °C overnight
on an orbital shaker (as above). The isotopic composition of N$_2$ was determined by injection
of 50 or 100 µL of headspace (porewater and core/slurry samples, respectively; CombiPAL,
CTC Analytics) into a continuous flow isotope-ratio mass spectrometer (Delta Plus, ThermoFinnigan) and measurement of mass-to-charge ratios 28, 29 and 30. Further details of calibration are provided as supplementary information. Samples for N₂O determination were prepared by withdrawing a sub-sample of the headspace described above (100 µL for porewater samples and 1-10 µL for slurried core samples) and injecting it into a gas-tight vial containing 2 nmoles of N₂O (prepared by dilution of 100 % N₂O in a N₂ matrix, British Oxygen Company). The entire contents of these vials was swept into a trace-gas pre-concentrator module (Cryo-Focusing, Precon, ThermoFinnigan) and mass-to-charge ratios 44, 45 and 46 were measured on the mass spectrometer described above. Samples for \(^{15}\)NO⁻ determination were prepared by reduction of nitrate to nitrite with spongy cadmium (modified from ref. 37 - 5 mL of sample and 0.2 mL of 1 M Imidazole were used and samples were incubated for 2 h on an orbital shaker, as above). Samples were then transferred to gas-tight vials (3 mL Exetainer, Labco) and a 0.5 mL helium headspace was added (as above). Nitrite was reduced to N₂ by injection of sulphamic acid through the septa (100 µL 4 mM sulphamic acid in 4 M HCl; B. Thamdrup, personal communication) and, after overnight equilibration, the headspace was analysed for \(^{15}\)N-N₂ as above. The amount of \(^{15}\)NO⁻ within each vial was determined by preparation of a calibration curve of differing amounts of \(^{15}\)NO₃⁻ (treated as above) versus the mass-to-charge ratio 29: sum of all areas.

Calculations for activity measurements

Production of \(^{15}\)N-N₂, anammox and denitrification potential in anoxic slurries were calculated using standard procedures. Rates of ambient anammox and denitrification were calculated using methods previously applied to intact sediment cores with differences in the \(^{15}\)N-labelling of the N₂ and N₂O pools used to determine the contribution of anammox. In oxic slurries, anammox and denitrification were apportioned by comparing the proportion of
15N in the produced N₂ to anammox and denitrification endmembers in a mixing model. All calculations used to derive rates, contribution of anammox to N₂ production and other parameters (e.g. base flow index) are provided as supplementary information.

**Statistical methods for activity measurements**

All statistics were performed in R\(^3\) (version 3.1.1) using RStudio\(^3\) (version 0.98.1091). Differences in anammox activity between groups was tested with linear mixed effects models using the nlme package\(^4\) where geology or permeability were fitted as fixed effects and site was a random effect.\(^4\) Model fit was improved by adding variance structure to the model allowing variance to differ between groups. Significance of fixed effects (p < 0.05) were determined by log likelihood ratio tests between the model of interest and a reduced model,\(^4\) i.e. with no fixed effect but just a random intercept (see Table S5).

The effect of porewater chemistry on anammox and denitrification was examined using multivariate techniques. First we used principal component analysis (PCA) to investigate correlations between chemical variables and differences in porewater chemistry between rivers. The PCA reduced 7 chemical variables (nitrate, nitrite, ammonium, dissolved O₂, SRP, iron (II) and pH) to two principal components (total variance explained = 74%). Principal component 1 accounted for 56 % of the variance and comprised strong positive loadings for ammonium, Iron (II) and SRP, strong negative loadings for nitrate and pH and an intermediate negative loading for O₂ (Figure S4). We have interpreted this axis as a chemical gradient moving from reduced porewaters, where mineralisation products such as ammonium and SRP accumulate (high scores), to oxidised porewaters (low scores) high in nitrate and intermediate in O₂. The chemistries captured within PC1 separate data into their respective geologies (Figure S4), essentially converting our categorical “gradient” of permeability (i.e. clay, sand or chalk) into a true riverine gradient. PC2 was most strongly
associated with oxygen (positive) and nitrite (negative, Figure S4), however, this axis only explained 19% of the variance.

We then performed a redundancy analysis using the vegan package with the same chemical dataset, geology as a grouping factor and response variables ambient anammox and denitrification rates and the contribution of anammox to N₂ production (ra). We attempted to determine the most parsimonious model by performing stepwise addition of the variables, however, after inclusion of the factor “geology” there were no significant improvements to the model Akaike Information Criterion. Geology alone, i.e. sand or clay, is not very useful for determining chemical controls on riverine anammox but when removed from the model the goodness of fit was considerably reduced (>11% reduction observed). We therefore determined the most parsimonious model by manually comparing adjusted R² values following the addition of chemical variables. The explanatory variables of the simplest model were found to be geology, ammonium, nitrate, nitrite and O₂ (Figure 3b, Adjusted R² = 0.40). In this simplest model 78% of the variance was explained by the 1st canonical axis which had similar chemical loadings as PC1 in the original PCA.

Relationships between anammox and other variables (e.g. hzo gene copy number) were quantified using Spearman’s rank correlation on untransformed data with p<0.05 used as the criteria for significance.

Molecular analyses

Collection of sediment. Sediment collected for potential anammox activity (see above) was sub-sampled for molecular analysis (n=3 for the 9 rivers sampled). Sediment was placed in sterile tubes and preserved cryogenically at -150°C.

qPCR gene abundance. DNA was extracted from 0.25 g wet weight sediment using PowerSoil® DNA Isolation Kit (MO BIO Laboratories, Inc). Gene abundance was
quantified by qPCR with SensiFAST SYBR No-ROX Kit (Bioline) on a CFX96 Real-Time PCR Detection System (BioRad) using the 16S rRNA primer pair Bakt_341F (CCTACGGGNGGCWGCAG) and Bakt_805R (GACTACHVGGGTATCTAATCC) and the hzo primer pair HZO-1F (AAGACNTGYCAYTGGGGWAAA) and HZO-1R (GACATAACCATACTKGTRTANACNGT). Gene abundances were quantified by absolute quantification method against an internal standard calibration curve of DNA standards of the target gene from 10^2 to 10^6 copies in 20 µl reactions containing 400 nM of primers and 1 µl of DNA template. Cycle conditions were 95 °C for 2 min followed by 40 cycles at 95 °C for 10s then 60 °C for 30 seconds. Amplification of a single product was confirmed by melting curve analysis.

**Amplicon sequencing.** Amplicon libraries were prepared by a 28-cycle PCR using primers containing the same target region as the qPCR primers but flanked with Illumina Nextera overhang sequences. Amplicons were purified using AMPure XP (Agencourt) SPRI bead protocols, before adding Illumina flowcell adapter sequences, and one of 96 unique combinations of Nextera paired-end Indexes via a 8-cycle PCR. Amplicons were again purified using AMPure XP beads, quantified using a Quant-iT Picogreen dsDNA assay kit (Life Technologies) on a Nanodrop 3300 fluorospectrometer (Thermo Scientific) and then pooled in equimolar concentrations. The amplicon libraries were quality checked using a DNA 1000 kit on at 2100 Bioanalyzer (Agilent) before sequencing was performed on the Illumina Miseq platform using a MiSeq reagent kit V3 (2 × 300 bp) at TGAC (The Genome Analysis Centre, Norwich). The sequencing reads were analysed using the QIIME pipeline and associated modules. Sequences were de-multiplexed using the Nextera Indexes and quality filtered to remove sequences below Q20 or that contained, any errors in the primer region, above 6 ambiguous bases, and chimeras. The quality filtered reads were clustered into operational taxonomic units (OTUs) using the USEARCH algorithm at the 0.95 level (hzo)
or 0.97 level (16S rRNA). 16S rRNA representative sequences from each OTU were assigned
taxonomic identities with the RDP classifier.\textsuperscript{48} Statistical analysis was performed in the R
statistical language version 3.1.3 using the R base libraries\textsuperscript{38} and the community ecology
analysis- specific package ‘vegan’.\textsuperscript{42} hzo gene multiple sequence alignment was performed
on the 100 most abundant OTUs (representing 92-93 % of all sequences in each geology) and
codon aligned deduced amino acid sequences using MUSCLE (MUltiple Sequence
Comparison by Log- Expectation)\textsuperscript{47} and phylogenies were constructed in MEGA6\textsuperscript{49} The
nucleotide sequence evolutionary history was inferred by using the Maximum Likelihood
method based on the General Time Reversible model.\textsuperscript{50} Initial trees for the heuristic search
were obtained by applying the Neighbor-Joining method to a matrix of pairwise distances
estimated using the Maximum Composite Likelihood (MCL) approach. A discrete Gamma
distribution was used to model evolutionary rate differences among sites. The amino acid
evolutionary history was inferred by using the Maximum Likelihood method based on the Le
and Gascuel 2008 model.\textsuperscript{51} Initial tree(s) for the heuristic search were obtained automatically
by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated
using a JTT model, and then selecting the topology with superior log likelihood value. A
discrete Gamma distribution was used to model evolutionary rate differences among sites.
Significance of branching order was determined by bootstrap analysis (1000 replicates).\textsuperscript{52}

\textbf{References for methods}

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(a) N\textsubscript{2} Production (nmol $^{15}$N g\textsuperscript{-1})

(b) Dissolved O\textsubscript{2} (µM)