Tracing lake pollution, eutrophication and partial recovery from the sediments of Windermere, UK, using geochemistry and sediment microfabrics

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Abstract

Many lakes undergo anthropogenically driven eutrophication and pollution leading to decreased water and sediment quality. These effects can enhance seasonally changing lake redox conditions that may concentrate potentially toxic elements. Here we report the results
of a multi-method geochemical and sediment microfabric analysis applied to reconstruct the 
history of cultural eutrophication and pollution of the North and South Basins of 
Windermere, UK. Eutrophication developed from the mid-19th to the earliest 20th centuries. 
Enhanced lake productivity is indicated by increased sedimentary δ¹³C, and increased 
pollution by a higher concentration of metals (Pb, Hg, and As) in the sediment, likely 
enhanced by incorporation and adsorption to settling diatom aggregates, preserved as 
sedimentary laminae. In the South Basin, increasing sediment δ¹⁵N values occur in step with 
Zn, Hg, and Cu, linking metal enrichment to isotopically heavy nitrate (N) from 
anthropogenic sources. From around 1930, decreases in Mn and Fe-rich laminae indicate 
reduced deep-water ventilation, whereas periods of sediment anoxia increased, being most 
severe in the deeper North Basin. Strongly reducing sediment conditions promoted Fe and 
Mn reduction and Pb-bearing barite formation, hitherto only described from toxic mine 
wastes and contaminated soils. From 1980 there was an increase in indicators of bottom 
water oxygenation, although not to before 1930. But in the South Basin, the continued 
impacts of sewage is indicated by elevated sediment δ¹⁵N. Imaging and X-ray microanalysis 
using scanning electron microscopy has shown seasonal-scale redox mineralisation of Mn, 
Fe, and Ba related to intermittent sediment anoxia. Elevated concentrations of these metals 
and As also occur in the surficial sediment and provide evidence for dynamic redox 
mobilisation of potentially toxic elements to the lake water. Concentrations of As (up to 80 
ppm), exceed international Sediment Quality Standards. This process may become more 
prevalent in the future with climate change driving lengthened summer stratification.

Key Words: paleolimnology, multi-method, cultural eutrophication, Itrax XRF core 
scanning, isotopic analysis, scanning electron microscopy.

1. Introduction
Freshwater lakes represent a critical resource providing a wide range of services, such as a municipal water source (Fowler et al., 2007), flood mitigation (Thampapillai and Musgrave, 1985), habitats for rare and protected species (Dudgeon et al., 2006; Maberly and Elliott, 2012), and are often the basis for a multi-million-pound tourism industry (Maltby et al., 2011). Despite their value, many are poorly managed and subject to anthropogenic stressors, such as cultural eutrophication caused predominantly by phosphorus (P) and nitrogen (N) enrichment (Richardson and Jørgensen, 1996), and toxic metal enrichment, from industrial and municipal pollution (Förstner and Wittmann, 2012). The consequent economic impact is estimated to be £75 -115 M yr$^{-1}$ in England and Wales alone (Harvey et al., 2012; Pretty et al., 2003). Furthermore, anthropogenic climate change has been linked to amplification of these stressors (Williamson et al., 2009), making the restoration of freshwater lakes a priority (Gulati et al., 2012). To this end, current legislation (the European Union Water Framework Directive, WFD; 2000/06/EC) legally obliges stakeholders to return waterbodies to “a good ecological and chemical status” with reference to pre-anthropogenic chemical, ecological and environmental conditions (Carvalho et al., 2019).

Numerous lakes in industrialised areas or near developing population centres have undergone similar histories over the past 1-2 centuries with progressive eutrophication and pollution driven by population increase, agricultural intensification and industrialisation (Dudgeon et al., 2006; Hodell and Schelske, 1998). Despite more recent remediation measures, anthropogenic activities have often left a legacy of sediments enriched in metals and trace elements (Thevenon et al., 2011). Many lakes vary seasonally with summer stratification allowing depletion of oxygen in the hypolimnion that may lead to anoxia in the bottom sediments. Such seasonal-scale redox changes may affect the absorption or release of pollutants, potentially producing relatively high concentrations on the lake bed or in the lake waters (Itai et al., 2012; Wu et al., 2017). Climate change may reinforce such processes
since raised temperatures may lead to a longer and more intense stratification period driving enhanced isolation of bottom waters (Jenny et al., 2014). In order to reconstruct the history of lake eutrophication and to assess the efficacy of lake restoration programmes, it is necessary to obtain detailed geochemical records of lake sediments to quantify the history of inputs and to investigate the controls on mineral formation and dissolution (Thevenon et al., 2011; Wu et al., 2017).

In this context, Windermere, England’s largest natural lake, has undergone a typical trajectory of pollution and eutrophication over the past 150 years (McGowan et al., 2012; Moorehouse et al., 2014; 2018). This has been recorded, in part, by regular monitoring of the water column since the 1940s (Pickering, 2001; Maberly et al., 2011), however, the extent to which anthropogenic pollution has affected the lake beyond the time scale of monitoring and how the lake sediments are affected are not yet fully understood. Windermere forms a particularly interesting case study as it comprises two basins which are variably susceptible to eutrophication, which together, constitute a sensitive barometer to the effects of anthropogenic and climatic influences (Pennington, 1973; McGowan et al. 2012).

This study has applied a range of geochemical and sediment fabric techniques to reconstruct the history of anthropogenic impacts on Windermere. By combining radiochronology, major and trace element geochemistry and high resolution scanning electron microscope sediment microfabric imaging on new gravity cores, we reconstruct the sedimentary history including the input of metals and toxic elements and the relation to biological activity and redox changes in the lake sediment. Sediment fabric analysis specifically enables us to relate anthropogenically-induced pollution and eutrophication to biological activity, including the reconstruction of seasonal-scale algal blooms in the lake surface waters and of the activity of zoobenthos on the lake bed.
2. Study location, materials and methods

2.1. Study site

Windermere, in the Lake District National Park, (Fig. 1), is of particular significance as it acts as a refugium for locally rare fish species such as the Arctic Char (Winfield et al. 2008; Miller et al., 2015), a municipal water supply for the North West of England during periods of drought, and is host to a thriving tourist industry. The lake is 17 km long with a maximum width of 1.5 km and is divided into a North Basin (maximum depth 64 m) and a South Basin (maximum depth 42 m) separated by an area of basement high, small islands and shallows with an average depth of 10 m (Pearsall and Pennington, 1973; Pickering, 2001; Miller et al., 2013). The larger North Basin has a surface area of 8.1 km² and volume of 201.8 x 10⁶ m³ while the South Basin has a surface area of 6.7 km² and volume just over half that of the North at 112.7 x 10⁶ m³ (McGowan et al., 2012). The main freshwater inflows are from the rivers Rothay and Brathay (Fig. 1). Primary inflow to the South Basin comes from the North Basin and from Cunsey Beck that drains the small seasonally anoxic lake, Esthwaite Water. Outflow is at the south end via the River Leven (Fig. 1), with retention times in the North and South Basin of 180 and 100 days respectively (Moorhouse et al., 2018).

The lake is currently classified as warm monomictic (Hutchinson and Löffler, 1956), becoming stratified in early summer with re-mixing usually in late October or early November in the South Basin and a later (November or December) in the North Basin (Pickering, 2001). The water column has been regularly monitored and sampled for its physical properties, chemistry and biology since 1945 by the Freshwater Biological Association (FBA) and the Centre for Ecology and Hydrology (CEH) (Reynolds and Irish, 2000; Pickering, 2001; McGowan et al., 2012). These records show a progressive mid- to late-20th century increase in eutrophication, especially in the South Basin (Reynolds and Irish,
The current trophic status of the North Basin is mesotrophic and that of the South Basin, eutrophic (Moorhouse et al., 2018). Palaeolimnological studies of the sediments of Windermere indicate a mid- to late-19th century onset for eutrophication driven primarily by population increase (Pennington, 1973; Sabater and Howarth, 1995, McGowan et al., 2012), marked in the sediment by the transition from “brown mud” to a “black ooze” (Pennington, 1973; Sabater and Howarth, 1995). Enrichments in some metals (Pb, Zn, Cu) were previously reported within the “black ooze” from cores taken in the South Basin in 1975 (Hamilton-Taylor et al., 1984). A more recent geochemical study of piston cores taken from Windermere in 2012 found a 20th century enrichment in metals (Miller et al., 2014), but did not recover the topmost 10–20 cm of sediment.

### 2.2. Core location, sampling and analysis

Following the bathymetric surveys and coring reported in Miller et al. (2013), the four gravity cores of the present study were taken in 2014 from mid-depths and deep basinal locations (Fig. 1), recovering the sediment-water interface and up to 46 cm of sediment (Table 1). The cores were split at the British Ocean Sediment Core Facility (BOSCORF), then processed and sampled using standard methods for high resolution sediment fabric analysis (Dean et al., 1999; Kemp et al., 2001). Split cores underwent digital imaging, x-radiograph imaging and geochemically analysis using an Itrax Energy Dispersive X-Ray Fluorescence (ED-XRF) core scanner (Croudace et al., 2006). To calibrate and complement the ED-XRF analyses, continuous sediment samples (1 cm resolution) were also analysed by Wavelength Dispersive XRF Spectrometer. Mercury was measured using a Milestone DMA-1 system (Milestone SRL, Italy) employing sample thermal decomposition, mercury amalgamation and atomic absorption detection providing a typical detection limit in the ppt range. Further details of the methods are found in the Supplementary Material.
2.2.1. **Microfabric analysis**

One centimetre-thick slab samples were taken continuously with overlapping sections for microfabric studies. These slabs were X-rayed, and following further subsampling, underwent fluid-displacive resin embedding after which covered thin sections (CTS) for optical microscopy and polished thin sections (PTS) for backscatter electron imagery were prepared. PTS were imaged and analysed using a Scanning Electron Microscope (SEM). Sediment fabrics, including bioturbation and mineral and organic components, were documented using a combination of photomosaic and higher magnification backscatter electron images (BSEI), together with optical thin section microscopy. For further analysis of microfossil remains and mineral habits and structures, SEM stub samples were prepared from the un-treated sediment counterparts to the embedded sediment sections.

To match broader scale variations in geochemistry with changes in sediment microfabric and mineralogy, elemental maps, line scans, and spot analyses of key intervals within PTS were undertaken in the SEM by energy dispersive X-ray spectroscopy microanalysis (EDS).

2.2.2. **TOC, TN, δ¹³C**

One gram samples, taken continuously at 2 cm intervals from cores SC68 and SC57 were analysed for TOC, TN, δ¹³C and δ¹⁵N using a Elementar Vario Isotope Cube Elemental Analyser equipped with a TCD (Thermal conductivity detector), interfaced with an isotope ratio mass spectrometer (IRMS). Six samples representing sediment endmember types in both basins were tested for calcium carbonate (CaCO₃), and all yielded results below the detection limit (4 counts).

2.3. **Chronology**

2.3.1. **²¹⁰Pb and ¹³⁷Cs**
One centimetre thick samples were taken at regular intervals and analysed for $^{210}\text{Pb}$ and $^{137}\text{Cs}$ in all cores. $^{137}\text{Cs}$ was measured by direct gamma spectrometry using a Canberra well-type HPGe gamma-ray spectrometer counting for 100,000 seconds and evaluating the 661 keV photopeak (Croudace et al., 2012; Ritchie and McHenry, 1990). Gamma spectra were processed using FITZPEAKS gamma deconvolving software (JF Computing, Stanford in the Vale, UK). Each activity was corrected for sample mass and volume. Lead-210 was measured (via its granddaughter Po-210) using a method based on Flynn (1968) involving double acid leaching, $^{209}\text{Po}$ spiking and auto-deposition onto silver discs to determine $^{210}\text{Pb}$ activities using alpha spectrometry (Croudace et al., 2012). The linear sedimentation accumulation rates (LSR) and subsequent age estimates reported below are generated from $^{210}\text{Pb}$ activity and validated using $^{137}\text{Cs}$.

A chronology was generated from excess $^{210}\text{Pb}$ ($^{210}\text{Pb}_{\text{xs}}$) activity data for the top of each gravity core by applying the constant flux: constant sedimentation (CF:CS) model (Robbins, 1978). The model generates an LSR that is extrapolated down core (Fig. 2). Age error was calculated by applying the CF:CS model to the upper and lower counting statistical uncertainty of $^{210}\text{Pb}_{\text{xs}}$ values to produce LSRs which were extrapolated down core to give an upper and lower age depth error estimate. All $^{210}\text{Pb}$ ages stated in this paper are given in years A.D. and are the median probable ages followed by instrument uncertainty—in brackets. LSR generated from $^{210}\text{Pb}$ were only applied to depths at which $^{210}\text{Pb}_{\text{xs}}$ activity exceeded the background concentration. Below this, age estimates were generated by linear interpolation of the model to the topmost radiocarbon age in the corresponding piston core. Accumulation rates based on $^{137}\text{Cs}$ used the depths of initial increase of the 1963 bomb peak from atmospheric nuclear weapons testing and the Chernobyl nuclear reactor incident (1986) (Ritchie and McHenry, 1990).

2.3.2. Radiocarbon dating
In 2012 piston cores were taken at the same locations as the gravity cores (Avery et al., 2017). From these split core surfaces a series of samples were taken for $^{14}$C analysis following an inspection of the core for macrofossils, and analysed by the Natural Environment Research Council Radiocarbon Facility (NRCF) in East Kilbride. In order to extend the age-depth model beyond the oldest date achievable by $^{210}$Pb and $^{137}$Cs the uppermost $^{14}$C analyses from the piston cores (Table 2) were used. Integration of the gravity and piston core age depth, and overlap at corresponding sites was achieved by matching lithostratigraphy, $^{210}$Pb and $^{137}$Cs data from both cores. Samples from the piston cores were for radiocarbon analysis were converted to graphite and processed at the Scottish Universities Environmental Research Centre (SUERC) laboratory for accelerator mass spectrometry (AMS) $^{14}$C analysis (Damon et al., 1989).

Radiocarbon dates were processed and calibrated using Calib 7.1 (Stuiver and Reimer, 1993) and the Intcal 13 calibration curve (Reimer et al., 2013). The radiocarbon results are reported with ages given as the calibrated median probability age, and $2\sigma$ calibrated ages.

3. Results

3.1. Age-depth model

The age-depth model and LSR generated from $^{210}$Pb and $^{137}$Cs data for the upper part of the cores (SC68 = 27cm, SC64 = 20.5cm, SC67 = 18.5cm, SC57 = 22.5cm) is described in detail in Fielding et al. (2018). In all cores linear interpolation between the deepest $^{210}$Pb activity and the upper most radiocarbon date in the corresponding piston cores (Table 2) yielded LSR an order of magnitude lower than those generated by the CF:CS model in the upper part of the core. This increase in sedimentation since the 19th century is consistent with other studies in the area, and could have a number of causes including increases in land erosion due to
agriculture and increases in biogenic sediment as a result of increased primary productivity in the lake (Sabater and Haworth, 1995; Schillereff et al., 2019).

3.2. Lithostratigraphy and Geochemistry

The North Basin cores comprise a lower pale brown mud (Unit I) overlain by a dark grey mud (Unit II), a thin pale clay rich interval (Unit III) and an upper pale brown mud (Unit IV) (Fig. 3). The South Basin stratigraphy is similar, except that Unit III is missing and the upper muds of Unit IV are darker in Core 57 (Fig. 4). This stratigraphy is broadly consistent with earlier cores taken from Windermere with the “brown mud” of Pennington (1973) and Sabater and Howarth (1995) corresponding to Unit I and their “black ooze” to our Unit II. The main difference is the thicker upper pale brown mud, our Unit I, reflecting an additional 3–4 decades of recent sediment accumulation, the transition to Unit I having been around 1970–1975. The thin clay-rich interval in the North Basin (Unit III) is interpreted as an event bed generated by a slope failure restricted to the North Basin caused by the 1979 Carlisle earthquake (Fielding et al., 2018; Musson and Henni, 2002). The transition from Unit I to Unit II occurs earlier in the deep South Basin cores (1870 in SC67; 1883 in SC57) than in those from the North Basin (1916 in SC64; 1920 in SC68).

Major and trace element variability are presented in Figs. 3, 4 and supplementary Figs. 1 - 4, and total carbon and nitrogen and stable isotopes δ\(^{13}\)C and δ\(^{15}\)N in Fig. 5. These results are discussed in section 4 below in the context of the overall pollution and eutrophication history of Windermere. Details of the geochemical microfabrics and barium mineralogy encountered are given below.

3.3. Sediment fabrics and preservation of lamination

3.3.1. Sediment microfabrics and lamina types
Microfabrics vary from laminated through homogeneous and unstructured sediment to pervasively pelleted, with pellets ranging in size between 50-350µm, (Fig. 6). Terrigenous silty clays (Fig. 6c) form the dominant sediment type and are interspersed with a more porous sediment (PS) type (paler in optical microscopy/ darker in BSEI) (Fig. 6b), comprising silty clay and organic material. Fe and/or Fe and Mn minerals typically occur within porous sediment laminae, for example in Unit I of SC68 and in near surface sediment (Fig. 6b). The silty clays of Unit II are distinctly dark-coloured under optical microscopy (Fig. 6d).

Obvious lamination comparable to regular varves (Zolitschka et al., 2015) was rarely observable directly on the split core surfaces. However, a combination of optical microscopy and BSEI of embedded sediment reveals a range of lamina types, the most common being 0.8 – 5.5 mm thick and comprising alternations of detrital terrigenous and higher porosity laminae with the latter often containing autochthonous organic material laminae and reduced detrital content (Figs 5c). The high porosity laminae also often contain enrichments of Fe or both Fe and Mn minerals (see 3.4.1, below).

3.3.2. Diatom ooze laminae

Microlithostratigraphic analysis shows that Units II and IV are abundant in diatom remains. In SC68, diatoms are concentrated in distinct laminae ranging in thickness from 0.2 to 1 mm (Fig. 7). Twelve diatom ooze laminae occur in SC68 and a single lamina is present in SC64. The greater preservation of laminae in SC68 reflects its deep basin location that would be more prone to anoxia (see 4.3.3, below). Imaging of stub samples taken from the laminae reveals that they are composed of near-monospecific concentrations of Asterionella formosa in Unit II, and Aulacoseira subarctica, in Unit IV (Fig. 7).

3.4. Geochemical fabrics
3.4.1. Fe- and Mn-rich laminae

Fe and/or Mn minerals typically occur within porous sediment laminae, for example in Unit I of SC68 and in near surface sediment (Fig. 6b). Enrichments in Fe and Mn occur intermittently in all the cores but are most abundant in Unit I of the deep North Basin (SC68) (Fig. 3). Peaks in Mn in the lower laminated sediments were associated with concentrations of a rhombohedral mineral which was identified as rhodochrosite (MnCO$_3$) by EDS elemental mapping and line scans (Fig. 8). The rhodochrosite occurs as both individual crystals with a typical size range of 2–10 µm, but also as dense clusters of crystals, up to 60 µm across (Fig. 8A). Mn peaks in the upper core interval are not associated with rhodochrosite, but rather with amorphous Mn oxyhydroxides. Fe peaks throughout the sediments were shown to be associated with authigenic amorphous and crystalline iron (oxy)hydroxide (Fe(O)OH), most likely lepidocrocite, occurring as subhedral platy agglomerates ranging in size from between 50 µm to 700 µm (Fig. 8B).

Selected intervals were investigated with EDS elemental mapping and line scans (Fig 9). To indicate variation in detrital sediment, Si is also shown. Decreases in Si coincide with increases in Mn and or Fe in the more porous laminae, they are also indicated by paler zones in the thin section photomicrographs (Fig. 9). In some cases, the laminae appear in pairs separated by terrigenous-rich laminae, with the Mn lamina located above or overlapping with the Fe-rich lamina (Fig. 9a, c). In this instance the laminae are distinct on the EDS maps and line scans despite pelletisation. In other cases, the order is less clear (Fig. 9) although the Mn enrichment (0.05 to 2.2 mm thick) generally occur towards the upper part of the Fe-enrichment (0.2 to 3 mm thick) while in some cases only Fe enrichments (0.1 to 1 mm thick) occur (Fig. 9). The mineral rich-detrital rich laminae couplets range between 0.14 and 0.8 cm in thickness with an average of 0.24 cm. Given the LSR of 0.29 cm/ yr$^{-1}$ for this Pb-dated section of the core, a quasi-annual occurrence is plausible for the upper intervals.
3.4.2. Barium mineralogy

Barium (Ba) content is marked in all cores, except SC68, by peak values near the sediment water interface (SWI), with maximum values of up to 0.2 wt% in core SC57. Below this there are smaller increases above the clay horizon in the North Basin cores, SC64 and SC68 and there is elevated Ba content within the upper part of Unit II in SC68 and SC67. In SC68 Ba also shows high variability at the base of the core. SEM examination of sediment reveals the common presence of small, typically 2–12 μm-sized crystals of barite that may occur in clusters or individually. SEM EDS analysis indicates that these have a normal barite BaSO$_4$ composition near the core top, but within Unit IV they contain Pb and so are on the Barite (BaSO$_4$)-Anglesite (PbSO$_4$) solid solution series (Fig. 10). The barite-anglesite crystals are most abundant in SC57 and SC64 but occur in all cores and increase in abundance in the mid and upper part of Unit IV. Here microlithostratigraphic analysis shows the presence of BaSO$_4$ in the SWI appearing as individual euhedral minerals up to 5 μm and in clusters up to 35 μm.

4. Discussion

4.1. Causes and significance of pelleting

In lake waters deeper than 20 m, pelletal structures (50–300 μm) are produced mainly by conveyer-belt feeding tubificid oligochaetes or by chironomid larvae (McCall and Tevesz, 1982). Both mechanisms can cause vertical redistribution of sediment but usually over very small distances (2–9 cm). In Windermere it was found that vertical redistribution by bioturbation was limited to 1.5 cm (Fielding et al., 2018) and that while this disrupted the original sediment structure in some places it did not completely remove it. The presence of pellets does not, however, necessarily imply sustained oxic conditions since oligochaetes such as *Tubifex* spp. are known to survive in anoxia (Famme and Knudsen, 1985) and have
been observed in abundance in the profundal sediments of Windermere even with seasonally anoxic bottom water conditions (Reynoldson, 1987). Furthermore, some chironomids contain haemoglobin, and as such can survive in anoxic conditions through a range of respiration strategies (Brodersen, 2008). In this instance, therefore, it is the extent of pelletisation that is likely related to the relative degree and persistence of sediment anoxia, since the overall benthic activity would be inhibited during prolonged anoxia (Reynoldson, 1987).

4.2. Use of Mn and Fe content and mineralogy as redox indicators

Mn and Fe are sensitive redox indicators since both metals have soluble reduced ions but on oxidation form solid oxyhydroxides (Davison, 1993). There are two circumstances in which Mn and Fe minerals may be used to reconstruct ventilation histories in lakes. In lakes with anoxic bottom waters, the dissolved ions of Fe and Mn are present. In such lakes, for example, Lake Zurich, seasonal ventilation of bottom waters and the introduction of oxygen leads to the formation and subsequent sedimentation of solid phase Fe- and Mn- oxyhydroxides that may preserve a record of ventilation events in seasonally laminated sediments (Naeher et al., 2013). In these cases, laminae of Fe/Mn minerals record lake ventilation events. On the other hand, in lakes without persistent anoxia, consumption of oxygen in the hypolimnion during the summer-autumn stratification period promotes sediment anoxia and leads to mobility of Fe and Mn ions. In this case Fe and Mn will precipitate at a redox boundary, either within the sediment (as is the case in Lake Baikal; Och et al., 2012) or at the sediment water interface (SWI) as pore waters interact with oxygenated bottom waters (Schaller et al., 1997). From monitoring of Windermere, hypolimnetic anoxia has only been recorded in the deepest part of the South Basin late in the stratified summer period occasionally between 1979 and 1991 during the period of peak eutrophication (Hamilton-Taylor and Willis, 1990; Pickering, 2001). The presence of Fe or Mn layers in Windermere sediments (Figs. 3, 4, 9), therefore, records periods where reduced bottom water oxygenation led to reduced oxygen in the upper sediments, and a shallowing of the
oxycline in the sediment, facilitating the mobility of Fe and Mn and their subsequent oxidation at
or near the SWI (Fig. 3). The occurrence of Fe laminae without overlying Mn concentrations (Fig.
9) indicates that some overturning episodes were not of sufficient duration or intensity to allow
the slower-oxidising Mn to precipitate suggesting release of Mn to the water column. In
scenarios where reducing conditions in the sediment result in the dissolution of Mn
oxyhydroxides there may still be the potential to preserve a record if rhodochrosite is formed (Yu
et al., 2016) as was evidently the case in Unit IV of SC68. The distribution and variability in Fe
and Mn may therefore be used to reconstruct the Windermere sediment redox history.

4.3. The sedimentary record of the pollution and eutrophication of Windermere

4.3.1. Asynchrony between the South and North Basin

The increase in metals (Pb, Zn, Cu) and As in the North Basin occurs around 1910–1920,
lagging the appearance in the South Basin by around 70-100 years for Pb and 30 years for
Zn, Cu and As (Figs. 3, 4; Supplementary Figs. 1-4). This lag in delivery to the North Basin is
unlikely to be due to differences in the sources of metals (sewage and atmospheric), as this
would have occurred almost synchronously on a catchment to regional scale. Instead, it may
be that the efficient delivery of the metals to the sediment was enhanced by increased algal
productivity and scavenging during flux. For example, sediment trap studies from
Windermere show take up of Zn by diatoms (Reynolds and Hamilton-Taylor, 1992) and
enrichments of both Zn and Pb were found associated with the eutrophic-indicating diatom
Asterionella formosa (Lund, 1957). Furthermore, the extracellular polymeric substances (EPS)
exuded by both diatoms and associated bacteria in lake “snow” aggregates (Schweitzer et
al., 2001) are also efficient scavengers of trace metals (Bhaskar and Bhosle, 2006; Comte et al.,
2008). The smaller water volume of the South Basin makes it more susceptible to enrichment
and eutrophication than the North Basin and it also receives a higher nutrient load
(Pickering, 2001), so that the earlier increase in metals may be related to the earlier increases
in algal productivity and associated scavenging from the water column. There is also a
similar lag in the $\delta^{15}$N of organic matter that is consistent with the earlier eutrophication in
the South Basin. Such earlier productivity increases in the South Basin are also recorded in
algal pigment records with South Basin production increasing markedly from around 1860
and that of the North Basin increasing more gradually from 1890 to highs from around 1940
(McGowan et al., 2012).

4.3.2. 1800–1940: The onset and increase in pollution

Increases in Pb content in the early to mid-19th century in the South Basin (SC67–1810s;
SC57–1850s) and from around 1910–20 in the North Basin represent the first significant signs
of anthropogenic contaminants in the sediment. Although there was considerable mining
and quarrying for metals in the North of the Windermere catchment in the 19th century, the
periods of metal extraction do not coincide with increases in metal content in the lake
sediment (Miller et al., 2014). Double-spike Pb isotope analysis of Windermere sediments
dating from the 1840s to the 1920s reveals a likely Pb source to be from Carboniferous coal,
related to the launch of coal-fired steam ships from 1845 and the further expansion of their
use following the opening of the Kendall-Windermere railway in 1847 (Fig. 11) (Miller et al.,
2014). The increase in Pb prior to other metals is also recorded in the sediments of the
adjacent Blelham Tarn (Ochsenbein et al., 1983). The increased use of petroleum in road
vehicles is also a significant contributor of Pb from 1920 onward (Miller et al., 2014).

The increase in the other metals (Zn, Cu, Hg; Supplementary Figs. 1-4) and As from 1880–
1990 and concomitant darkening of the sediment in the South Basin (Fig. 4), coincides with
increased sewage discharge from the rapidly increasing population centres around the lake
(McGowan et al., 2012) and this is consistent with previous metal studies in Windermere
(Hamilton-Taylor and Willis, 1990). High levels of As and Hg persisted in the South Basin
from the 1920s through the following six decades. Increasing values of the δ^{15}N of organic matter in the South Basin sediment also occur from around 1890 (SC57), in step with the metals (other than Pb) from this time. In the North Basin, by contrast, a prominent increase in δ^{15}N of organic matter together with metals does not occur until the 1930s (Figs 3, 13, Supplementary Figs 1, 2). Elevation in δ^{15}N of organic matter can be attributed to a number of causes including increased algal productivity (Hodell and Schelske, 1998) or the input of isotopically heavy nitrate from primary sewage or farm runoff (Meyers, 1994; Moorhouse et al., 2018). Although, in Windermere all of these may be applicable, recent analysis shows that lake primary producer variability tracks modifications to waste water treatment works in both the North (~65% variance explained) and South Basins (~62% variance explained) (Moorhouse et al. 2018). The darkening of the sediment also coincided with the first rise to prominence of the eutrophic-indicating diatom Asterionella formosa (Sabater and Haworth, 1995) (Figs 7, 12). Elevated productivity in both basins is also evidenced by increasing δ^{13}C values of organic matter through the 19th and early 20th centuries (Fig. 5). Phytoplankton preferentially take up the lighter ^{12}C isotope during photosynthesis, but in periods of sustained high productivity they will increasingly utilise the heavier isotope resulting in an increase in δ^{13}C values (Schelske and Hodell, 1995). The increase in δ^{13}C values in Windermere is consistent with steadily increasing algal productivity during this period. The δ^{13}C values are around 1‰ heavier in the South Basin, possibly reflecting its greater susceptibility to eutrophication due to its smaller volume, although phytoplantonic species composition or organic matter source changes could also affect this (Gu et al. 1999). While there is broad agreement with δ^{13}C values from an earlier Windermere core study (McGowan et al., 2012) other organic proxies (TOC, TN, C:N) show a somewhat different trend. This could be due to differences in analytical methods since McGowan et al. measured carbon by loss on ignition whereas we used an elemental analyser.
Regarding sediment redox conditions, the regular occurrence of Fe and Mn laminae in the deep North Basin (SC68 – 54 m water depth) prior to the 19th century indicate intermittent sediment anoxia, but with regular deep-water ventilation in the period prior to eutrophication. When compared to the LSR of the core (<27.5 cm depth: 0.29 cm yrs⁻¹, >27.5 cm depth: 0.04 cm yrs⁻¹) the regular spacing of these mineral laminae (Fig. 3) (0.2–0.7 cm) suggests these features represent redox changes, and resulting deposition of Mn and or Fe, occurring on annual to multi-annual scale (Fig. 9a). The pattern is consistent with summer stratification and a reduction in bottom water oxygenation that promoted anoxia in the sediment and allowing the release and upwards diffusion of dissolved Fe and Mn ions. This would have been followed by lake turnover in the autumn/ winter that led to re-oxygenation of bottom waters and formation of the Fe and Mn oxides that precipitated at a redox boundary within the surficial sediment. The occurrence of rhodochrosite rather than Mn oxyhydroxides suggests that reducing conditions in the sediment may have driven dissolution of the oxyhydroxides, but that the Mn subsequently combined with carbonate (Yu et al., 2016). The less frequent occurrence of Fe laminae and presence of only a single Mn lamina in SC64 is consistent with its location in shallower waters (26m) which remain oxygenated for longer than deeper waters in the hypolimnion. Here, the occasional Fe and Mn laminae represent periods where sufficient dysoxia occurred in even the shallower waters, followed by reoxygenation resulted in the concentrating of redox sensitive elements in the sediments.

As the colour changes from pale brown to grey (Unit I to II) in the 1920s these Fe and Mn laminae become progressively less abundant until they cease, with the slower oxidising Mn disappearing first. The concomitant decrease in the extent of pelletisation of the sediment is also consistent with progressively increasing bottom water hypoxia and sediment anoxia (Fig. 3). Combined with the cessation of Mn oxyhydroxide preservation and the lack of
pelletisation in the shallower SC64, this suggests increasing incidence and persistence of sediment anoxia in the North Basin (Fig. 3).

4.3.3. 1940–1980: Peak pollution and eutrophication, sediment anoxia and bottom water hypoxia

Sustained high concentrations of metals were present in both basins from around 1940 coinciding with further darkening of the sediment, high levels of sediment organic carbon and persistent high values of the δ¹⁵N of organic matter. This was likely driven by an increase in resident population during World War II that eventually overwhelmed the existing sewage treatment facilities at Tower Wood (McGowan et al., 2012) leading to increased sewage discharge to the lake. In addition, following WW II the application of N rich artificial fertiliser significantly increased UK wide (McGowan et al., 2012) making agricultural run off another likely driver of persistent high values of the δ¹⁵N.

In both basins, peak organic carbon contents occurred from 1950–1980 and this also coincides with some of the highest δ¹³C values marking peak productivity (Fig. 5). Lake monitoring during this period recorded blooms dominated by the diatom Asterionella formosa, a characteristic indicator of eutrophication (Lund, 1972; Lund et al., 1963). These recurrent annual blooms occurred in May/June with massive flux in late June. Close monitoring of the 1947 bloom observed a flocculent mass of cells accumulating on the surface sediment (Lund et al., 1963). From the 1940s, increasing sediment anoxia in the deep North Basin (SC68), as evidenced by a decrease in pelletisation and cessation of Fe and Mn laminae, led to the preservation of several of these layers as sedimentary laminae recording the near-monospecific nature of the Asterionella formosa blooms (Figs 3, 4, 7, 12), with the last preserved around 1983 (Fig. 12). Increasing diatom productivity and eutrophication was further driven by increases in soluble reactive phosphorus, particularly in the South Basin,
together with increases in nitrate in both basins from the mid-1960s (Reynolds and Irish, 2000). Sustained, elevated concentrations of metals and As occurred throughout this period with peaks occurring mainly between 1950–1960.

During the time interval 1940–1980, peak sediment anoxia is indicated in all cores with the most persistent in the deep North Basin (SC68). The presence of Fe and Mn laminae and decrease in pelletisation in SC67 (1960–1980) indicate that enhanced sediment anoxia also extended to shallower zones of the lake at this time. Intermittent Fe laminations in the deep South Basin (SC57) are consistent with sediment anoxia developing reducing conditions sufficient to mobilise both Fe and Mn but with insufficient bottom water re-ventilation to re-oxidise the Mn. This sediment evidence is consistent with observations of seasonally declining oxygen concentrations in the summer-autumn hypolimnion (Talling and Heaney, 1988) and episodic bottom water anoxia recorded in the deep South Basin from 1979 (Pickering, 2001). In fact, elevated Mn concentrations were observed in bottom waters during late season stratification in the 1970s (Hamilton-Taylor et al., 1984; Pickering, 2001).

The peak sedimentary metal concentrations at this time may have been boosted by the scavenging of metals by Mn- and Fe-oxides forming in the bottom waters (Hamilton-Taylor and Davison, 1995; Xue et al., 1997).

There is no evidence of authigenic pyrite from SEM observations, so although the bottom sediments were anoxic, they were rarely sulphidic and there is no evidence for sulphate reduction and pyrite formation consistent with sulphur speciation studies in Windermere (Davison et al., 1985). However, increases in sulphur occur in the South Basin cores in Unit II (>0.5%) and all sediment cores contain frequent 2–20 µm-sized barite crystals (Fig 9). The occurrence of the barite as euhedral crystals suggests a different mechanism of formation from that of the adjacent seasonally anoxic Esthwaite Water, where small (3–4 µm) spherical granules of barite are biologically mediated within protozoa (Finlay et al., 1983; Smith et al.,
The origins of the Ba concentrations that led to the formation of the euhedral barite crystals in Windermere may be similar to those responsible for the transfer of Ba from bottom waters to the topmost sediment in Lake Biwa, Japan. Here, the hypolimnion reaches minimum dissolved oxygen concentrations at the end of the stratification period in December but is never totally depleted (Sugiyama et al., 1992). In this case, during the period of stratification, enhanced anoxia within bottom sediments leads to the release of Fe and Mn into the oxic bottom waters where they are re-oxygenated to form hydrous oxides. The hydrous Mn oxides have a strong affinity for Ba which they readily scavenge and this results in draw down of Ba to the sediment as the hydrous Mn oxides settle (Sugiyama et al., 1992). Although water column sulphate is low in freshwater lakes, bacterial oxidation of organic sulphur (Fakhraee et al., 2017) may have been important in producing the sulphate for barite formation. Unlike the pure barite crystals in the near surface sediment, the crystals in Unit III consistently contain Pb. The presence of barite-anglesite solid solution has not been reported before from lake sediments and has primarily been associated with mining waste and contaminated soils (Courtin-Nomade et al., 2008; Fernandez-Gonzalez et al., 2013).

4.3.4. 1980–2010: Partial Recovery

From around 1980 (Units II to IV) the sediment colour changes back to pale brown in SC68 and SC67 and becomes less dark in SC64 and SC57. In the South Basin, decreased sediment anoxia is also indicated by the resumption of pelletisation (enhanced benthic activity), cessation of the Fe and Mn laminae, and decreases in sedimentary S. The decline in Pb after peaks in the 1960s follows the marked decline in atmospheric Pb since peaks in the 1970s (Rippey and Douglas, 2004). A post 1960s decline in metals and As in the South Basin may be ascribed to the effective operation of the then newly expanded (new sedimentation tanks
and additional filters) Tower Wood Sewage treatment works from 1967 (Fig. 10) (McGowan et al. 2012).

Phosphate stripping was introduced 1991–1992 at the sewage treatment works at Tower Wood (South Basin) and Ambleside (North Basin) (Fig. 11) and measured deep water oxygen concentrations showed a marked improvement in the 1990s (Pickering, 2001). The North Basin (SC68) shows a progressive decline in the δ¹⁵N of organic matter indicating a diminishing contribution from isotopically heavy sewage, which may in part be due to the building of extra STW (sewage treatment works) at Gasmere and Elterwater (Fig. 11), but the persistence of diatom ooze laminae resulting from diatom mass flux record sustained eutrophication. On the other hand the δ¹⁵N of organic matter in the South Basin (SC57) shows no such decline, highlighting continued influence of isotopically heavy N in sewage input. There is also a significant increase in organic carbon content in the North Basin. Although phosphate stripping was introduced in 1991/1992, phosphate contents in the sediment actually increase synchronously in all cores at this time and reach concentrations of 1–2% in the surface sediment. Sustained concentrations of soluble reactive phosphorus are also present in lake waters (Winfield et al., 2008). Microlithostratigraphic and EDS line scan analysis show that P is associated with Fe oxyhydroxides in the top few cm of sediment, suggesting redox mobilisation and potential P release from the sediment to lake waters (see discussion on redox-driven processes in 4.2, above). This highlights the issue that although external P loading may be reduced in mitigation programmes, internal P loading originating by release from a pool earlier accumulated in the sediment may jeopardize improvement in water quality (Søndergaard et al., 2003).

Since 1980, water column surveys identify both Asterionella formosa and Aulacoseira spp. to be abundant in the spring (Canter and Haworth, 2010; Feuchtmayr et al., 2012; Haworth, 1988). Other key spring bloom species present, include Tabellaria flocculosa and Fragilaria crotonensis
with *T. flocculosa* identified as more abundant than *Aulacoseira* spp. (Feuchtmayr et al., 2012). These species do not form sediment laminae suggesting that they are less important for controlling flux than either *Asterionella formosa* or *Aulacoseira subarctica*. The lamina records show that *Aulacoseira subarctica* has replaced *Asterionella formosa* as the dominant lamina-former since the early 1990s, matching with broader changes observed in the FBA plankton records from the south end of the North Basin (Thackeray et al., 2015) (Fig. 12). Clearly, only a few of the surface blooms recorded by the FBA correspond to laminae in SC68. This may reflect some combination of the different locations (north end of the North Basin for the core and south end for the FBA sampling), flux attenuation (so that only the most intense blooms sedimented to depth) and intermittent bioturbation as evidenced by pelletisation (Fig. 3). The installation of phosphate stripping to the Tower Wood STW in 1991/1992 has led to a decrease in eutrophication. This is therefore consistent with the view that *Asterionella formosa* favours fully eutrophic conditions while *Aulacoseira subarctica* thrives with moderate increases in nutrients but is disadvantaged by further enrichment (Gibson et al., 2003).

Throughout Unit IV (1980 to the 2000’s) the geochemistry and micropalaeontology of the sediments show a partial recovery from eutrophication, but further indicate that the recovery has not been sustained, consistent with evidence for a renewed deterioration in oxygen levels in lake waters (Jones et al., 2008).

4.4. The modern lake - 2010–2014: Topmost sediment and recent redox-driven processes

4.4.1. *Mn, Fe and Ba*

Microfabric and geochemical analysis shows that with the exception of SC67, the topmost sediment of Unit I (1.5–3 cm) in all cores is enriched in Mn and Fe. This is consistent with upwards diffusion of reduced and mobile Mn and Fe from the anoxic sediment and their
subsequent precipitation at the redox boundary near the SWI as oxyhydroxides (Davison, 1993). The exceptional concentrations of 12.5 wt% MnO in SC57 (Deep South Basin) requires further explanation and suggests advection of Mn to the deep South Basin. The mechanism for this may be analogous to that identified in the Swiss lake Baldaggersee where Mn mobilisation leads to “geochemical focusing”. This process involves the release of Mn from anoxic sediments to overlying oxygenated waters where it starts to reprecipitate, but because this takes days to weeks there is down slope movement resulting in concentrations in the deeper part of the lake (Schaller and Wehrli, 1997). A prerequisite for this to occur is for anoxic sediment to be in contact with an oxic water column, a situation which is apparently increasingly occurring during the stratified period in Windermere (Jones et al., 2008). A similar redox-driven “geochemical focusing” process is also invoked for Fe, although this is less mobile with a tendency to precipitate more rapidly as oxyhydroxides (Schaller and Wehrli, 1997). The two cores that show the greatest Mn enrichment (SC 64 and SC57) also show significant Ba enrichment to values exceeding 0.2 wt% in the deep South Basin. This association is consistent with the sequestration of Ba by Mn oxyhydroxides (Section 4.3.3, above) as also documented from Lake Biwa (Sugiyama et al., 1992).

4.4.2. Arsenic

There is relative enrichment of As in the surface sediments of SC68, SC64 and SC57 to values approaching 70 ppm (Figs 3, 4). Arsenic is readily adsorbed by Fe oxyhydroxides in sediment, and the cycles of Fe and As are tightly coupled (Belzile and Tessier, 1990; Couture et al., 2010; Dixit and Hering, 2003; Pierce and Moore, 1982). Field and experimental evidence indicates that As is also adsorbed by hydrous Mn oxide (Takamatsu et al., 1985) and arsenite becomes adsorbed to more freshly formed Fe and Mn oxyhydroxides (Dixit and Hering, 2003; Farmer and Lovell, 1986). This can lead to concentrations in surfaces sediments far in excess of that delivered from the water column (Farmer and Lovell, 1986)
and has likely led to the enrichment of As in surface sediments of Windermere. Furthermore, where a legacy of anthropogenic As pollution remains in deeper sediments, it can remain elevated in the surface sediments long after exposure to anthropogenic sources of As have been reduced (Fabian et al., 2003). There can be further enrichment of As in surface sediments as a result of sediment anoxia where Fe and Mn oxyhydroxides and adsorbed arsenate are reduced to labile species and diffuse upward to the redox boundary. As is the case for Mn, As may be released to the water column during periods of lake stratification, especially where hypoxia develops in the hypolimnion and remains for long periods (Keimowitz et al., 2017). Progressive warming driven by climate change has increased stratification in Windermere (Maberly and Elliott, 2012) so that this is likely to be an ongoing process that may adversely affect the ecosystems (Thackeray et al., 2013) including fisheries (Jones et al., 2008).

4.5. Implications for future water quality and potential hazards

Many countries, including the UK, have failed to introduce the formal Sediment Quality Standards (SQS) such as those set by the Australian, New Zealand (ANZECC/ARMCANZ, 2000), Canadian (CCME EPC-98E), and Netherlands governments (Dutch Water Act) (Fig. 14) (Burton Jr, 2002). These are designed to be adopted to control sediment contamination considered dangerous to benthic life and human health (Burton Jr, 2002). The enrichment of Pb and As in Unit III (especially in the 20th century) too often exceed these standards in both of the South Basin cores for Pb. Concentrations of As in all cores, (except SC64) also exceed the SQS (Fig. 14). This is concerning if these sediments were to be exposed to water column hypoxic or disoxia conditions where both Pb and As can become mobile (Hamilton-Taylor and Davison, 1995). Evidence of mass transport deposits and slope failure scarps found in the distal lake slopes suggest that this is increasingly possible (Fielding et al., 2018; Miller et al., 2013). What is more concerning is that in both North Basin cores, As
values at the SWI exceed all three SQS. As discussed, Mn and As dissolution and migration under low oxygen conditions to the above water column is common in warm monomictic lakes during the Summer/Autumn bottom water oxygen low (Keimowitz et al., 2017). This would suggest that significant quantities of As are mobile and being released into the water column during Summer and Autumn months (El Bilali et al., 2002). As discussed above the Water Framework Directive legally obliges stakeholders to return water bodies to “good ecological and chemical status” with reference to pre-anthropogenic chemical, ecological and environmental conditions. Our combined geochemical and sediment microfabric study demonstrates that, despite mitigation measures being put in place, pollution issues still remain in Windermere. With climate change driving current trends of longer and more intense summer lake stratification, the degree of mobilisation of toxic elements may increase in the future, requiring a reappraisal of mitigation strategies.

5. Conclusions

A multi-proxy investigation of sediment cores from Windermere, England’s largest natural lake, has yielded a detailed history of changing lake and catchment conditions over the past 300 years. Prior to the 19th century in the lake’s South Basin and the 20th century in the North Basin, Fe and Mn rich laminae indicate regular, seasonal-scale ventilation of bottom waters. Following local population increases and associated increasing fossil fuel use through the 19th century, Pb content in the sediment increases first in the South Basin (SC67–1810s; SC57–1850s) and from 1910–20 in the North Basin. This is followed later by increases in other metals (Zn, Cu, Hg), and As. At the same time increases in sedimentary δ13C, and the appearances of monospecific diatom ooze lamina, together with decreasing Fe and Mn lamina show a move to decreasing bottom water ventilation caused by eutrophication.
Greater values of sedimentary $\delta^{15}\text{N}$ through the same period are also consistent with enhanced productivity coupled with increases in sewage discharge and farm runoff in to the lake, with the latter two inputs being likely sources of metals and nutrient enrichment. The synchronous increase in metals with indicators of lake productivity also point toward the enhanced incorporation and adsorption of metals to settling diatom aggregates bringing elevated flux to the sediments. Through the middle of the 20th century benthic activity intermittently ceased in the deeper North Basin due to persistent strongly reducing conditions in the sediment and bottom waters as indicated by increased $S$ content, low $\text{Fe-Mn}$ values and the formation of unusual $\text{Pb}$-bearing barite mineralisation, hitherto only described from toxic mine wastes and contaminated soils. From 1980 there was a partial recovery, with bioturbated sediment reflecting increases in oxygenation of deep waters. In the South Basin however, elevated $\delta^{15}\text{N}$ of organic matter indicates continued impacts of sewage discharge. Imaging and X-ray microanalysis using scanning electron microscopy has enabled the identification of seasonal-scale redox mineralisation of $\text{Mn, Fe, Ba}$ related to intermittent sediment anoxia. Recent, persistent sediment anoxia, strengthened by hypoxic bottom waters during seasonal stratification, has resulted in trace element mobilisation. Subsequent oxidation at the SWI or within the bottom waters has caused significant enrichment of $\text{Mn, Fe, As, P and Ba}$ in the surficial sediment, and in the North Basin, As enrichment at the surface exceeds Sediment Quality Standards set by the Australian, New Zealand, Canadian and Netherlands governments. It would thus appear that despite mitigation measures being put in place pollution issues still remain in Windermere, and with climate change driving current trends of increased duration of summer stratification in the lake, the mobilisation of toxic elements may increase in the future.

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Figure and table captions

Fig. 1. Location of study area and cores. Inset shows position of Windermere in relation to the Lake District topography. The extent of the Windermere catchment is given by the
dashed line. Key rivers and water bodies feeding the lake are shown, together with major population centres and sewage treatment works (Triangles). (1.5 column fitting image)

Fig. 2. Age depth model measurements of artificial radionuclides for cores SC68 (a), SC64 (b), SC67 (c), SC57 (d) and calculated linear sedimentation rates (LSR). R² shows the linear trend for the Ln Pb data. In the age panel black circles show the Pb CF:CS LSR age depth model at 2 cm intervals. Vertical error indicates bulk sample interval and horizontal error represents the extent of the maximum and minimum age depth based on machine measurement error. White circles show the Cs based age depth at 1986 and 1963 with vertical error also indicating the bulk sample interval. Red circles show data points that were removed due to being partly within the mass transport deposit (MTD). Grey triangles show the age depth at the base of the core derived from linear interpolation between Pb and C derived ages (Table 2)(2 column fitting image).

Fig. 3. Stratigraphy and Geochemistry for North Basin Cores SC68 and SC64. Core depth in cm, core photograph, core x-radiograph, lithological units, lithostratigraphy, sediment fabric types, Pb CF:CS LSR age depth model for the North Basin gravity cores. For geochemistry black lines show Itrax ED-XRF Fe, Mn, Pb, As, S and P (lower scale). Red dots show discreet WD-XRF data for K₂O, Fe₂O₃, MnO, Pb, As, S and P₂O₅ (titles in brackets, upper scales). Vertical errors on WD-XRF show the sampling interval. Water depths of each coring site are shown above the corresponding core. (2 column fitting image).

Fig. 4. Stratigraphy and Geochemistry for South Basin Cores SC67 and SC57. Core depth in cm, core photograph, core x-radiograph, lithological units, lithostratigraphy, sediment fabric types, Pb CF:CS LSR age depth model for the South Basin gravity cores. For geochemistry black lines show Itrax ED-XRF Fe, Mn, Pb, As, S and P (lower scale). Red dots show discreet WD-XRF data for K₂O, Fe₂O₃, MnO, Pb, As, S and P₂O₅ (titles in brackets, upper scales).
Vertical errors on WD-XRF show the sampling interval. Water depths of each coring site are shown above the corresponding core. (2 column fitting image).

**Fig. 5.** Organic chemistry for SC68 and SC57. Core depth in cm, core photograph, core x-radiograph, lithostratigraphy, sediment fabric types, $^{210}\text{Pb}$ CF:CS LSR age depth model for the South Basin gravity cores. This is followed by $\delta^{13}\text{C}$, $\delta^{15}\text{N}$, TOC, TN, C/N (grey dots). Vertical error shows the sampling interval for each data point (2 column fitting image).

**Fig. 6.** Detailed sediment and microfabric types as identified using optical thin section microscopy (left) and backscatter electron imagery (middle). Fabric types: [a] pelleted [b] homogenous, [c] laminated (alternating porous and terrigenous-rich laminae), [d] dark muds. Key sediment fabric features are featured (left). (1.5 column fitting image).

**Fig. 7.** Position of *Aulacoseira subarctica* bloom lamina (A) within SC11 optical thin section and back scatter electron image (BSEI) and *Asterionella formosa* bloom lamina (B) within SC14 optical thin section and BSEI, and detailed optical thin section and BSEI form both A and B. (2 column fitting image).

**Fig. 8.** Core photograph of SC68 with depth and age showing the position of rhodochrosite minerals shown in panel A, with corresponding EDS elemental map showing Mn highlighted in purple. Lower panel shows the EDS line scan of Mn and Ca in mineral across x-y. Panel B shows amorphous Fe oxyhydroxide, as determined by EDS from location indicated in panel A. (2 column fitting image).

**Fig. 9.** Optical thin section, back scatter electron image, mineral lamina log, EDS map (Si, Mn and Fe) and EDS line scan (Si, Mn and Fe) of alternating laminae for thin sections (A) SC68_19, (B) SC68_20 and (C) SC68_21 from SC68. (2 column fitting image). (2 column fitting image).
Fig. 10. Core SC68 photograph along with depth and age showing the position of samples containing examples of (A) multiple barite-anglesite minerals, and (B, top) and individual barite-anglesite mineral. (B, bottom) Also shown is an EDS line scan of Pb, Ba and S in mineral B. (2 column fitting image).

Fig. 11. A time line of the anthropogenic environmental variables. Modern artificial fertiliser manufacturing began in the UK in the early 20th century (dashed line), exceeded 200 Tonnes \(N \times 10^3\) use nationwide in \(\sim 1945\) (solid line) and peaked in \(\sim 1990\) (diamond) (McGowan et al. 2012). \(\Delta 15N\) and aspects of significant sediment chemistry results from this paper are shown for the North (SC68) and South Basin (SC57). For chemistry, solid lines represent onset and increase in concentration; dashed lines represent decreasing values. Full geochemistry is given in Figs 3 and 4 and discussed in the text. (1.5 column fitting image).

Fig. 12. Comparison of sediment diatom record with FBA diatom record through the period 1977–2014. Asta - Asterionella Formosa; Aula - Aulacoseira subarctica. (1 column fitting image)

Fig. 13. SC68 (a), SC64 (b), SC67 (c) and SC57 (d) Pb (black dots, left y-axis), As (red triangles, right y-axis) concentrations over time along with Australian-New Zealand, Canadian (short dashes) and Netherlands (long dashes) sediment quality standards for Pb (black) and As (red). (2 column fitting image)

Table 1. Gravity core location description and coordinates, depth and length.

Table 2. The depth, age and 2σ calibrated ages of the top most \(^{14}\)C from the piston cores at each site. The depth scale is composite constructed using overlap between the gravity and piston cores. Mean ages and radiocarbon 2σ age ranges are expressed years before present (1950 A.D.).
Figure 1
Figure 2

(a) SC 68 LSR = 0.28 cm yr\(^{-1}\)
Depth (cm) \(^{137}\)Cs Unsupported \(^{210}\)Pb (Bq/g) \(^{210}\)Pb\(_{ss}\) Age

(b) SC 64 LSR = 0.14 cm yr\(^{-1}\)
Depth (cm) \(^{137}\)Cs Unsupported \(^{230}\)Th (Bq/g) \(^{230}\)Th\(_{ss}\) Age

(c) SC 67 LSR (0 - 10.5 cm) = 0.29 cm yr\(^{-1}\)
LSR (10.5 - 30.5 cm) = 0.12 cm yr\(^{-1}\)
Depth (cm) \(^{137}\)Cs Unsupported \(^{230}\)Th (Bq/g) \(^{230}\)Th\(_{ss}\) Age

(d) SC 57 LSR (0 - 11.5 cm) = 0.27 cm yr\(^{-1}\)
LSR (11.5 - 35.5 cm) = 0.12 cm yr\(^{-1}\)
Depth (cm) \(^{137}\)Cs Unsupported \(^{230}\)Th (Bq/g) \(^{230}\)Th\(_{ss}\) Age
Figure 3
Figure 4
Figure 6

**Sediment and Fabric type**

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Figure 7
### Table 1

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### Table 2

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<td>bulk</td>
<td>1020</td>
<td>1109 - 1125</td>
</tr>
<tr>
<td>64</td>
<td>92</td>
<td>leaf</td>
<td>2544</td>
<td>2363-2620</td>
</tr>
<tr>
<td>67</td>
<td>48</td>
<td>twig</td>
<td>1214</td>
<td>1083-1160</td>
</tr>
<tr>
<td>57</td>
<td>78</td>
<td>wood</td>
<td>1651</td>
<td>1554-1732</td>
</tr>
</tbody>
</table>
6. Supplementary material

6.1. Detailed methods

6.1.1. Coring and location

Following the bathymetric surveys outlined in Miller et al. (2013 cores were taken with a Uwitec 86 mm diameter gravity corer over one day using two small vessels. On recovery sodium polyacrylate was used to solidify the water and preserve the water-sediment interface (WSI), a simple log was then taken before transport to the British Ocean Sediment Core Facility (BOSCORF).

6.1.2. Initial sampling and core logging

Digitally imaging was done using a Geotek™ Multi Sensor Core Logging-Core Imaging System (MSCL-CIS) and X-rayed. Slab samples were taken continuously with overlapping
sections for microfabric studies. The 1 cm-thick slabs were further X-rayed using a Hewlett Packard Faxitron X-radiography cabinet at 35 kVe for 10 seconds.

6.1.3. Microlithostratigraphy

To complement geochemical analyses, changes in the sediment microfabric were studied. Following sub sampling, slabs underwent fluid-displacive resin embedding after which covered thin sections (CTS) for optical microscopy and polished thin sections (PTS) for backscatter electron imagery were prepared. PTS were imaged and analysed using a Carl Zeiss LEO 1450VP Scanning Electron Microscope (SEM).

6.1.4. Geochemical analysis

6.1.5. Itrax XRF Core scanning

The core archive-half surface was analysed using an itrax XRF core scanner that incorporates and Energy Dispersive X-Ray Fluorescence analysis (ED-XRF) system (Croudace et al., 2006) with a step size of 200 μm. Data points were excluded from the analysis where surface discontinuities (e.g. cracks etc) led to reduced count rates. Data points with zero validity or a mean standard error of >5 were also excluded from analysis following recommended procedures. Elemental data (counts) was normalised by dividing it by total kilocounts per second (kcps) for each interval to account for changes in the core density and surface high between sample points and cores (Croudace et al. 2006).

6.1.6. Wavelength Dispersive XRF

In order to calibrate and complement the ED-XRF semi-quantitative itrax analysis, continuous 1 cm-thick homogenised bulk samples were analysed to produce fully quantitative results using a Philips Magix-Pro Sequential Wavelength Dispersive XRF Spectrometer WD-XRF (Almelo – Holland), with a 4 kW rhodium anode X-ray tube.
6.1.7. SEM-based energy dispersive X-ray microanalysis

The EDS instrumentation for elemental microanalysis was an Oxford Instruments X-Act 10mm$^2$ area Silicon Drift Detector, coupled with the AZtec Energy software system (v.3.1). Line scans were run at 20 kV, a WD of 19 mm, and a dwell time per analysis site of 60 seconds, with EDS data collected at ~5,000 cps. The calculated data have been acquired using standardless analysis, hence all results are normalized to 100%. Minimum detection limits are 0.195% for Na K$\alpha$ and decreasing to 0.085% for Ca K$\alpha$ (Goldstein et al., 2003).

6.1.8. TOC, TN, $\delta^{13}$C

SC68 from the North Basin and SC57 from the South Basin were analysed for TOC, TN, $\delta^{13}$C. 1 g samples were taken continuously at 2 cm intervals and freeze dried for 48 hours. Between 13 to 15 mg were taken from 6 samples representing end member sediment types in both cores were analysed for calcium carbonate (CaCO$_3$) content at the University of Southampton on AutoMate + CM5015 coulometer. All samples were analysed for TOC, TN, $\delta^{13}$C using a Elementar Vario Isotope Cube Elemental Analyser equipped with a TCD (Thermal conductivity detector) which is interfaced with an Isoprime 100 continuous flow isotope ratio mass spectrometer (IRMS). Acetanilide was used as an elemental standard for C and N and USGS40 and USGS41 as international reference materials for the normalisation of the isotope ratios.
Supplementary Fig. 1.

Stratigraphy and geochemistry for SC68. Core depth in cm, core photograph, core x-radiograph, lithostratigraphy, sediment fabric types, $^{209}$Pb CF:CS LSAR age depth model for the North Basin gravity cores. For geochemistry black lines show Itrax ED-XRF Zn, Ba, Cu, Ni and Co (lower scale). Red dots show discreet WD-XRF data for Zn, Ba, Cu, Ni and Co (titles in brackets, upper scales). Vertical errors on WD-XRF show the sampling interval. Water depths of each coring site are shown above the corresponding core.
Supplementary Fig. 2.

Stratigraphy and geochemistry for SC64. Core depth in cm, core photograph, core x-radiograph, lithostratigraphy, sediment fabric types, $^{209}$Pb CF:CS LSAR age depth model for the North Basin gravity cores. For geochemistry black lines show Itrax ED-XRF Zn, Ba, Cu, Ni and Co (lower scale). Red dots show discreet WD-XRF data for Zn, Ba, Cu, Ni and Co (titles in brackets, upper scales). Vertical errors on WD-XRF show the sampling interval. Water depths of each coring site are shown above the corresponding core.
Supplementary Fig. 3.

Stratigraphy and geochemistry for SC67. Core depth in cm, core photograph, core x-radiograph, lithostratigraphy, sediment fabric types, $^{209}$Pb CF:CS LSAR age depth model for the North Basin gravity cores. For geochemistry black lines show Itrax ED-XRF Zn, Ba, Cu, Ni and Co (lower scale). Red dots show discreet WD-XRF data for Zn, Ba, Cu, Ni and Co (titles in brackets, upper scales). Vertical errors on WD-XRF show the sampling interval. Water depths of each coring site are shown above the corresponding core.
Supplementary Fig. 4.

Stratigraphy and geochemistry for SC57. Core depth in cm, core photograph, core x-radiograph, lithostratigraphy, sediment fabric types, $^{209}$Pb CF:CS LSAR age depth model for the North Basin gravity cores. For geochemistry black lines show Itrax ED-XRF Zn, Ba, Cu, Ni and Co (lower scale). Red dots show discreet WD-XRF data for Zn, Ba, Cu, Ni and Co (titles in brackets, upper scales). Vertical errors on WD-XRF show the sampling interval. Water depths of each coring site are shown above the corresponding core.