

# **Assessing human impact on Rostherne Mere, UK, using the geochemistry of organic matter**

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## **Abstract**

Recent changes in the geochemistry of organic material from a hyper-eutrophic lake (Rostherne Mere, UK) were investigated using the stable isotope, geochemical, and molecular composition of radiometrically-dated sediment cores. Modern samples suggest that recent sedimentation is dominated by algal production, however the presence of higher thermal stability components suggests there is also a minor component of allochthonous organic material. Sediment cores reveal that absolute proxy values and the magnitude of observed changes are broadly homogenous across the lake basin, in both deep and shallow

water environments. A transition to environmental conditions favouring enhanced algal productivity in more recent sediments is suggested by a progression to higher TOC and lower C/N,  $\delta^{13}\text{C}_{\text{org}}$ , and average *n*-alkane chain length through the cores. A strong covariance between TOC and Rock-Eval Hydrogen Index implies this transition is driven by an increasing algal contribution rather than being a response to variations in the source of organic matter. Decadal trends and abrupt shifts in organic geochemical proxies, such as TOC,  $\delta^{13}\text{C}_{\text{org}}$ , and C/N, are suggested to be directly related to changes in external anthropogenic nutrient loading following the construction and decommissioning of point-source sewage treatment plants, set against a background of intensive agriculture. The development of hyper-eutrophic conditions likely occurred in stages, where rapid transitions are associated with the commencement of sewage effluent input in the 1930s, population increases in the 1980s, and a dramatic reduction in external nutrient loads in the 1990s. Recovery of the lake ecosystem is limited by internal nutrient recycling, and organic proxies indicate that the geochemistry of sediments has remained relatively constant since effluent diversion. This study highlights the utility of organic geochemical parameters, such as  $\delta^{13}\text{C}_{\text{org}}$ , Rock-Eval, and biomarkers, in tracing recent eutrophication processes in lakes to provide evidence for the timing and scale of anthropogenic environmental change.

## **1. Introduction**

Organic matter is a key biogeochemical component of lake sediment, and comprises a complex mixture of material that originates from organisms living within the lake and in its surrounding catchment. In many productive lakes the autochthonous detritus of phytoplankton is the dominant source of organic material to sediments (Cohen, 2003), although the contribution from macrophytes and allochthonous organic matter pools, such as terrestrial plant materials and soils, can also be important (Meyers, 1997). The geochemical composition of lacustrine organic matter can be used to distinguish the source of material

preserved in lake sediments and identify alteration pathways, and therefore is able to provide information on environmental change over the history of a lake (Meyers & Ishiwatari, 1993). Established bulk geochemical proxies, including the carbon isotope composition of organic matter ( $\delta^{13}\text{C}_{\text{org}}$ ) and carbon to nitrogen ratio (C/N), have been widely used for palaeoenvironmental reconstructions (Meyers & Teranes, 2001), but cannot fully discriminate between different organic pools (Holtvoeth et al., 2016), and therefore a combination of other techniques are needed.

Rock-Eval pyrolysis is a screening technique typically associated with determining the hydrocarbon potential of petroleum source rocks (Espitalie et al., 1977), however the technique has more recently been applied in palaeoenvironmental investigations to examine the organic composition of sediments from soils, wetlands, and lakes (Zocatelli et al., 2012; Khan et al., 2013; Sebag et al., 2013; Lacey et al., 2015; Newell et al., 2016). These studies highlight how Rock-Eval parameters can be used to assess the relative contributions from different sources of organic matter in sediments, and also provide evidence on the geochemical quality and degradation history of organic material. In addition, specific compounds can be traced to an individual organism, group, or process using biomarkers (Peters et al., 2005), which enables the recovery of detailed palaeoenvironmental information (Castañeda & Schouten, 2011). While the geochemical analysis of bulk sediment is useful for detailing general trends, biomarkers provide specific information on production, delivery, and preservation of organic sediment components (Meyers, 1997); for example, homologues of saturated long straight-chained hydrocarbons, the normal alkanes (*n*-alkanes), can be used to differentiate sources of organic matter. Short chain *n*-alkanes are typical of aquatic algae ( $n\text{C}_{15-21}$ ), a large component of which are dominated by the  $n\text{C}_{17}$  alkane (Clark & Blumer, 1971; Cranwell et al., 1987), mid chains ( $n\text{C}_{23-25}$ ) are associated with submerged aquatic

macrophytes, and longer chains ( $nC_{27-35}$ ) are a dominant component of higher plants and emergent macrophytes (Eglinton & Hamilton, 1967; Ficken et al., 2000).

Organic matter from both aquatic and terrestrial sources may also be subject to mineralisation and degradation, and consequently it is important for palaeoenvironmental reconstructions to identify the extent of alteration (Meyers & Teranes, 2001). The concentration of hydrogen in hydrocarbons is influenced by the redox conditions of the depositional environment, as hydrocarbons are commonly volatile when in the presence of oxygen (Cohen, 2003), therefore Rock-Eval proxies can provide evidence on the geochemical quality and degradation history of organic matter.

Recent eutrophication of lakes is common due to the increased input of phosphorus (P) and nitrogen (N), from effluent, agricultural run-off, and other anthropogenic nutrient sources. Eutrophication can lead to a loss of biodiversity and also directly affect the health of humans and animals through declining water quality (Krivtsov et al., 2001; Moss et al., 2005). One such eutrophic lake is Rostherne Mere in the UK. Over much of the 20<sup>th</sup> century (1935-1991), the lake received effluent from a local sewage treatment works in addition to nutrient-rich run-off from surrounding farmland (Carvalho et al., 1995). The site is of international importance (National Nature Reserve, Ramsar site, Site of Special Scientific Interest) and has a long-standing history of scientific monitoring since 1912 (Pearsall, 1923; Reynolds & Bellinger, 1992). In addition, Rostherne has a well-documented account of anthropogenic catchment disturbance making it a unique site to investigate human impact on the lacustrine environment through the organic geochemistry of lake sediments. Previous geochemical work on surface sediments and a 30 cm core from the deepest part of the lake suggests there is a high contribution of algal material to the most recent sediments (Brooks et al., 1976; Gaskell & Eglinton, 1976; Cardoso et al., 1983), although no chronological work was carried out on the core. A further 35-cm sediment core taken in 1993 was analysed for molecular and

isotope composition (Prartono & Wolff, 1998), which also indicated a strong allochthonous contribution in surficial sediments, but similar to previous work no dating was carried out on the core and end-member materials were not defined.

Here, we present a new multi-proxy organic geochemical record from Rostherne Mere that combines total organic carbon (TOC) and total nitrogen (TN) concentrations (from which we calculate C/N),  $\delta^{13}\text{C}_{\text{org}}$ , Rock-Eval parameters, and biomarkers (*n*-alkanes) on radiometrically dated ( $^{210}\text{Pb}$ ,  $^{137}\text{Cs}$ , and  $^{241}\text{Am}$ ) sediment cores. This study also uses trap sediments, fresh vegetation, and soil samples to identify the main constituents of modern sedimentary organic matter and improve end-member characterisation, as well as evaluate the role of degradation in both anoxic and oxic environments through litter bag experiments. Our main aim was to assess quantitatively the geochemistry of recent and sedimentary organic matter to provide a calibration of modern materials and to identify the primary drivers and timing of environmental change at Rostherne Mere. This supports characterisation of the transition to hyper-eutrophic conditions and the effectiveness of remedial actions at the lake. We assume that major shifts in the geochemical composition of sediments will occur contemporaneously with the introduction and cessation of sewage effluent input, as a result of organic matter productivity changes due to variations in external nutrient loading. Overall, this work seeks to demonstrate the efficacy of using organic geochemistry to investigate the impact of human activity on lacustrine environments.

## **2. Study site**

Rostherne Mere (53°20'N, 2°24'W) is the most northerly of a sequence of three meres, together with Mere Mere and Little Mere, located 17 km south-west of Manchester in the Shropshire-Cheshire plain, UK (Figure 1).. The elongate lake basin most likely formed as a

kettle hole, which deepened due to gradual subsidence caused by the dissolution of underlying salt-bearing strata following ice sheet retreat at the end of the Last Glacial period (Banks, 1970; Reynolds, 1979). The lake has a maximum depth of 31 m, an average depth of 14 m, and covers an area of approximately 0.5 km<sup>2</sup>, making it one of the largest and deepest of the Shropshire-Cheshire meres (Reynolds, 1979). The large lake volume of 6.6×10<sup>6</sup> m<sup>3</sup> (Woof & Wall, 1984) produces a comparatively long water residence time of 1.5-2.7 years (Carvalho, 1993; Moss et al., 2005), although recent high resolution monitoring suggests a residence time of closer to ca. 0.8 years (A. Radbourne & D. Ryves, unpublished data). The majority of water inflow to Rostherne Mere is derived from Rostherne Brook (~75%) entering at the southwest margin of the lake, which drains 7 km<sup>2</sup> of the 9 km<sup>2</sup> catchment (Carvalho et al., 1995). The remaining water input is divided between contributions from small springs and transitory drainage ditches, groundwater and direct rainfall (Carvalho, 1993; Carvalho et al., 1995). Water output comprises evaporation from the lake surface during the summer months and river outflow to the southeast margin through Blackburn's Brook. The lake is monomictic and thermal stratification typically arises between April and November causing anoxia in the hypolimnion. Dissolved oxygen stratification takes place from early Summer and peak deoxygenation occurs during late September to early October and can extend to the base of the thermocline at around 12 m water depth (Davison & Woof, 1984; Scott, 2014). The stratification regime is enhanced by eutrophication, which has increased over the 20<sup>th</sup> century due to anthropogenic nutrient enrichment sourced from greater agricultural runoff and upstream sewage effluent input (Carvalho et al., 1995; Krivtsov et al., 2001). Effluent has been discharged into the lake from two sewage treatment plants since their construction in 1935, dominating the external nutrient load sourced from the Rostherne Brook. The sewage flow was diverted away from the catchment in 1991, following which total external nutrient loads were measured to have decreased by up to 75% (Moss et

al., 1997). Conversely, internal nutrient variation was relatively consistent between years, steadily decreasing until ~2002 (Moss et al., 2005), and has since stabilised showing only minor oscillations (A. Radbourne & D. Ryves, unpublished data). High concentrations of P and N accumulate in the hypolimnion during the spring-autumn period of stratification and anoxia, which are then released and transferred to the epilimnion following overturn in late-November (Carvalho et al., 1995; Scott, 2014). The substantial degree of internal nutrient loading within the basin results in high levels of primary productivity and the development of algal blooms. Summer months are dominated by cyanobacteria and cryptophytes, principally consisting of *Anabaena* spp., *Aphanizomenon flos-aquae*, and *Cryptomonas*, and planktonic diatom blooms, mainly comprising *Asterionella formosa*, *Aulacoseira granulata*, and smaller centrics (e.g. *Stephanodiscus* and *Cyclostephanos* spp.), are typical of spring and autumn months (Carvalho, 1993; Moss et al., 2005). Submerged aquatic plant species diversity today is very low and in littoral areas < 1m deep dominated by *Callitriche hermaphroditica*. Lake margins are dominated by a narrow band of *Phragmites australis* with some *Typha angustifolia*, behind which is *Salix* scrub and alder carr (*Alnus glutinosa*) with areas of relatively species-rich wetland vegetation (Natural England, 2012).

### **3. Materials and methods**

#### *3.1 Core recovery and sampling*

A 27 cm-long sediment core (RM-S1) was recovered from the northwestern part of the lake in June 2014 at a water depth of 4.9 m (oxic zone; Figure 1) using a HON-Kajak gravity corer (Renberg, 1991). The RM-S1 core was extruded in the field at 1-cm intervals. A further 51 cm-long core (RM-D4) was retrieved from the central pelagic area in September 2014 at a water depth of 30.8 m (anoxic zone; Figure 1). The coring tube was packed with a foam block, wrapped in polythene film, and transported to the British Geological Survey (BGS).

The tube was split, the core halves were frozen, and the working half was sectioned partially thawed at 1 cm intervals and then freeze-dried.

To enable the examination of the geochemical composition of autochthonous and floating allochthonous organic matter, open tube sediment traps were installed in the central pelagic area of the lake in April 2014 (~30 m water depth). A shallow trap was set at 8 m below the lake surface, and a deeper trap at 22.5 m below the lake surface. The open traps consist of four plastic tubes set in a cross shape that have a trapping ratio of 6.25 (450 mm length, 72 mm internal diameter). The traps were recovered and emptied in June 2014, reset at the same location and depths, and a further collection was conducted in September 2014. Following each collection, the trap samples from respective sampling depths were amalgamated into polypropylene bottles.

To investigate the major allochthonous sources of organic matter to Rostherne Mere, surface soil samples were taken from a grassland meadow, woodland, and from the channel of the Rostherne Brook using a spade. In addition, submerged and emergent macrophytes were collected from the shoreline of the mere, including *Phragmites*, *Typha*, and the submerged macrophyte *Callitriche*. The *Phragmites* and *Typha* samples were divided into two groups; the first included the stem and leaves, and the second comprised the roots. The five plant samples (*Phragmites* root/stem, *Typha* root/stem, and *Callitriche*) were then each sub-divided into three aliquots. To examine the possible effects degradation may have on the geochemistry of the fresh vegetation, two of the three aliquots of each sample were sealed within individual 125 µm mesh bags using cable ties. One group of five aliquots were submerged in the central pelagic area at a water depth of 26.5 m within the deoxygenated zone, and the second group of five aliquots were submerged in the shallow oxic zone in the northwestern part of the lake at a water depth of 5.2 m. Each of the groups were left in the water column at their respective depths for three months between April and September 2014,



and then recovered and transferred to polythene bags. The trap sediment, vegetation, and soil samples were freeze-dried and freezer milled prior to analysis.

### *3.2 Chronology*

Sediment samples from cores RM-D4 and RM-S1 were analysed at the Environmental Radiometric Facility, University College London, for  $^{210}\text{Pb}$ ,  $^{226}\text{Ra}$ ,  $^{137}\text{Cs}$ , and  $^{241}\text{Am}$  by direct gamma assay using ORTEC HPGe GWL series well-type coaxial low background intrinsic germanium detectors (as described by Yang & Turner, 2013).  $^{210}\text{Pb}$  was determined by means of its gamma emissions at 46.5 keV. Samples were stored for three weeks in sealed containers to allow for radioactive equilibration, and  $^{226}\text{Ra}$  was then measured by the 295 keV and 352 keV gamma rays emitted by its daughter isotope  $^{214}\text{Pb}$ .  $^{137}\text{Cs}$  and  $^{241}\text{Am}$  were determined by their emissions at 662 keV and 59.5 keV. Corrections were applied for the effect of self-absorption of low energy gamma rays within the sample (Appleby et al., 1992).

### *3.3 Isotope analysis of organic matter and C/N*

Samples were analysed for TOC, TN, and  $\delta^{13}\text{C}_{\text{org}}$  at the BGS. For the soils and lake sediments, any carbonate present was removed prior to analysis using 5% hydrochloric acid for 24 hours and then rinsed in deionised water to neutral pH, dried at 40 °C, and homogenised in agate. The prepared samples were analysed in tin capsules using an on-line system comprising a Costech ECS4010 elemental analyser (TOC and TN) and a VG Triple-Trap and Optima dual-inlet mass spectrometer ( $\delta^{13}\text{C}_{\text{org}}$ ).  $\delta^{13}\text{C}_{\text{org}}$  were calibrated to the Vienna Pee Dee Belemnite (V-PDB) scale using within-run laboratory standards (BROC2 and SOILB) calibrated against NBS-19 and NBS-22 international standards. Analytical reproducibility ( $1\sigma$ ) for the within-run laboratory standards was  $<0.05\text{‰}$  for  $\delta^{13}\text{C}_{\text{org}}$ , and  $<0.5\%$  for TOC and TN. C/N is given as the weight ratio.

### 3.4 Rock-Eval pyrolysis

A Rock-Eval 6 analyser at the BGS configured in standard mode (pyrolysis and oxidation as a sequential process) was used to analyse samples using a technique previously employed for lake sediments and peats (Lacey et al., 2015; Newell et al., 2016). The performance of the instrument was assessed by comparison to the accepted values of Institute Français du Pétrole standards (IFP 160000, S/N1 5-081840). Rock-Eval parameters were determined as the amount of hydrocarbons previously generated and distilled upon heating to 300 °C (S1), those generated through thermal cracking of bound organic matter (S2), the CO<sub>2</sub> released during pyrolysis (S3), TOC (sum of pyrolysed and residual carbon), and mineral carbon (MinC; e.g. calcium carbonate). The hydrocarbons generated as a proportion of the total organic matter, Hydrogen Index (HI), is calculated from  $S2 \times 100 / \text{TOC}$ , and the amount of oxygen with respect to total organic matter, Oxygen Index (OI), is calculated from  $S3 \times 100 / \text{TOC}$ .

### 3.5 *n*-Alkane analysis

Freeze-dried samples from cores RM-S1 and RM-D4, trap sediment, vegetation and soils were analysed for *n*-alkanes at BGS using the technique described by Newell et al. (2016). The estimated ratio of terrigenous to aquatic *n*-alkanes (TAR; Bourbonniere & Meyers, 1996) was calculated using the quantified peak areas  $\text{TAR} = (nC_{27} + nC_{29} + nC_{31}) / (nC_{15} + nC_{17} + nC_{19})$ . To further characterise the distribution of *n*-alkanes, the weighted average *n*-alkane carbon chain length (ACL; Bush & McInerney, 2013), where  $C_n$  is the *n*-alkane concentration and  $n$  the number of carbon atoms, was determined using  $\text{ACL} = \Sigma(C_n \times n) / \Sigma(C_n)$ . The relative contribution to lake sediments from submerged or floating macrophytes in comparison to emergent and terrestrial plants ( $P_{\text{aq}}$ ; Ficken et al., 2000) was calculated using  $P_{\text{aq}} = (nC_{23} + nC_{25}) / (nC_{23} + nC_{25} + nC_{29} + nC_{31})$ .

## 4 Results

### 4.1 Chronology

For core RM-S1, total  $^{210}\text{Pb}$  activity reached equilibrium with supported  $^{210}\text{Pb}$  activity at 23 cm depth (Figure 2). The maximum level of unsupported  $^{210}\text{Pb}$  is found below the surface at 3.5 cm, suggesting an increase in sedimentation rates in recent years. A transition to lower levels of unsupported  $^{210}\text{Pb}$  between 15 and 18 cm implies high sedimentation rates and implies that sediment slumping may have occurred, which is likely to have been instantaneous. A well-resolved  $^{137}\text{Cs}$  peak at 12.5 cm (Figure 2) most probably reflects the fallout from atmospheric testing of nuclear weapons in 1963. A simple constant rate of  $^{210}\text{Pb}$  supply (CRS) model was used for dating, and the  $^{137}\text{Cs}$  peak at 12.5 cm, along with a concomitant peak in  $^{241}\text{Am}$  (Figure 2), was used to correct the final  $^{210}\text{Pb}$  chronology for core RM-S1 such that 12.5 cm was deposited in 1963.

In core RM-D4, total  $^{210}\text{Pb}$  activity reached equilibrium with supported  $^{210}\text{Pb}$  activity at 49 cm depth (Figure 2). A relatively uniform sedimentation rate across the upper 35 cm of the core is suggested by a balanced exponential decline in unsupported  $^{210}\text{Pb}$  activity, with increased sedimentation rates at around 38 cm. The well-resolved  $^{137}\text{Cs}$  peak at 37.5 cm most likely is contemporaneous with the peak observed in core RM-S1 at 12.5 cm and reflects the maximum fallout from atmospheric nuclear weapons testing in 1963, which is supported by a concomitant excursion in  $^{241}\text{Am}$  (Figure 2). The final  $^{210}\text{Pb}$  chronology for core RM-D4 was corrected by using the  $^{137}\text{Cs}$  and  $^{241}\text{Am}$  tie point of 37.5 cm equating to 1963.

### 4.2 Trap sediments, soils, and vegetation samples

Results for trap sediments, soils, and vegetation samples are given in Table 1 and Figure 3. The trap collection from June shows high MinC (6%) and TOC (10%), and low C/N (7) and  $\delta^{13}\text{C}_{\text{org}}$  (-31‰). TOC, C/N, and  $\delta^{13}\text{C}_{\text{org}}$  increase in the September trap (19%, 8, and -28‰,

respectively), and MinC decreases to 3%. Rock-Eval parameter HI is broadly consistent between the two collection periods (390-400), although decreases to 370 in the deep water June trap coincident with higher OI (219), and OI is lower (~164) in September. ACL and TAR remain consistent between June and September collections. All of the trap sediments show a narrow *n*-alkane composition dominated by a strong maximum at *n*C<sub>17</sub>, with only a minor contribution from longer-chained *n*-alkanes (Figure 3). Total alkane concentrations are higher for the September trap sediments (average = ~ 104 μg g<sup>-1</sup>) compared to the trap sediments collected in June (average ~ 60 μg g<sup>-1</sup>).

All three soil samples show very low MinC (<1%) and relatively low TOC (~3-6%), however C/N is much higher for woodland soil (C/N ~35) than for inflow and grassland soils (C/N ~12; Table 1). Woodland and inflow soils have similar HI and OI (~204 and 237, respectively), with grassland soil showing marginally higher values for both HI and OI (266 and 260, respectively). Soil δ<sup>13</sup>C<sub>org</sub> ranges between -27.8 and -29.5 ‰ (Table 1). Ratios of *n*-alkanes are relatively consistent between the different soil samples, with high ACL (~27.6) and high TAR (~32.2). Total *n*-alkane concentrations for soils are comparatively low (between ~ 9 and 20 μg/g), with varying carbon chain length maxima at *n*C<sub>27</sub>, *n*C<sub>29</sub>, and *n*C<sub>31</sub> for woodland, inflow, and grassland soils, respectively (Figure 3).

Fresh samples of the dominant vegetation from around and within the lake (*Phragmites*, *Typha*, and *Callitriche*) show low MinC and high TOC, HI, and OI (Table 1). The submerged aquatic macrophyte *Callitriche* has lower C/N (12.3) and higher δ<sup>13</sup>C<sub>org</sub> (-24.5 ‰) compared to *Phragmites* and *Typha* root and stem samples (C/N >30, δ<sup>13</sup>C<sub>org</sub> ~ -28.0 ‰; C/N could not be calculated for *Phragmites* and *Typha* roots due to very low N). Fresh *Phragmites* and *Typha* root samples and *Callitriche* have an ACL of ~ 25, whereas the stem samples of *Phragmites* and *Typha* show moderately higher ACL values of between 26 and 28. The fresh vegetation samples show a broad *n*-alkane composition (Figure 3), typically having maxima

above  $nC_{25}$ , where total  $n$ -alkane concentrations range from  $\sim 10 \mu\text{g/g}$  (*Phragmites* root) up to  $\sim 486 \mu\text{g/g}$  (*Typha* stem). The  $n$ -alkane composition of *Callitriche* shows a bimodal distribution with carbon chain length maxima at both  $nC_{23}$  and  $nC_{29}$  (Figure 3).

Vegetation samples that were submerged in the lake and exposed to anoxic and oxic conditions typically have lower TOC when compared to their fresh vegetation equivalent, although *Phragmites* and *Typha* stems only display a minor change in TOC (Table 1). A  $-4 \text{‰}$  shift in  $\delta^{13}\text{C}_{\text{org}}$  of *Callitriche* is observed between fresh and degraded samples, whereas both root and stems of *Phragmites* and *Typha* have more consistent values. Conversely, HI remains unchanged in *Callitriche* between fresh and degraded samples, in contrast to *Typha* and *Phragmites* which show a major increase in HI for those samples exposed to oxic conditions and a lesser increase for anoxic conditions. OI decreases substantially in all degraded vegetation samples. Total  $n$ -alkane concentrations for degraded *Typha* and *Phragmites* samples are generally higher than fresh counterparts, and lower for *Callitriche* (Figure 3). The overall  $n$ -alkane distribution of degraded *Typha* and *Phragmites* is consistent with the observed pattern in fresh samples, however *Callitriche* shows a shift to an  $nC_{27}$ -dominated assemblage under oxic conditions and is  $nC_{27}$ - $nC_{29}$ -dominated under anoxic conditions, also losing a component of long- and short-chained homologues.

#### 4.3 Sediment cores

Bulk organic geochemical data and  $n$ -alkane-based ratios for core RM-S1 are shown in Figure 4 and averaged data are provided in Table 1. From the Rock-Eval data, MinC varies between 1 and 3% and is lowest at the base of the core ( $< 1\%$ ), showing a trend towards higher values up to the top of the core, while TOC is stable around 11% up to 17 cm, after which values increase between 17 and 7 cm, and then show fluctuations around a mean of 16% in the uppermost sediments. There is a higher degree of variability in C/N in the lower

core with values of between 14 and 16. Above 17 cm, C/N is more stable showing a trend to lower values and decreasing to a C/N of 10 at the very top of the core. As with TOC,  $\delta^{13}\text{C}_{\text{org}}$  has consistent values of around  $-27\text{‰}$  between the base of the core and 19 cm, and then decreases from 17 cm to the top of the core, with an increase in the rate of decline between 10 and 5 cm, where values are approximately  $-29\text{‰}$ . A progressive increase in HI from a value of 232 up to 296 is observed between the base of the core and 10 cm, which is followed by a transition to lower HI around 5 cm and then a subsequent increase to values around 300 at the top of the core. OI shows an opposing trend to HI in the lowermost core where values decrease from 166 to 145 between 27 and 16 cm, and then OI shows a consistent increase through the rest of the core to a value of 204 in the most recent sediments. The *n*-alkane-based ratios ACL and TAR are both high in the lower core around 28 and 12, respectively, and show a shift to lower values of around 25 and 3, respectively, in the upper core. The  $\text{P}_{\text{aq}}$  ratio has an opposing trend, where lower values of 0.2 at the bottom of the core transition to higher values around 0.5 in the upper core.

Bulk geochemical data and *n*-alkane based ratios for core RM-D4 are given in Figure 5 and averaged data are provided in Table 1. MinC progresses from lower to higher values throughout the core, ranging between approximately 1 and 4%. TOC has variable but increasing values of between 5 and 13% at the bottom of the sequence up to around 30 cm, after which values are more consistent between 11 and 13% until the upper few cm of sediment where an increase to ~15% TOC is observed. C/N shows a broad decline to lower values throughout the core, between values of 14 at the very base of the core and 9 at the top of the core. As with C/N,  $\delta^{13}\text{C}_{\text{org}}$  decreases through the core from  $-27\text{‰}$  in basal sediments to  $-30\text{‰}$  at the top of the core, with a pronounced shift to lower  $\delta^{13}\text{C}_{\text{org}}$  around 13 cm core depth. An increase in HI from roughly 180 to 280 is observed through the bottom 20 cm of the core, which is followed by relatively stable values of around 275 between 31 and 3 cm,

followed by an increase to 333 in surface sediments. From the base of the core to 24 cm, OI shows a decreasing trend from a maximum value of 212 down to a minimum value of 156. The decrease is followed by a shift to higher and more stable values of around 187 through to 5 cm, after which OI decreases to around 172 in the uppermost sediments. In the lower core (51 to 17 cm), the *n*-alkane ratios ACL and TAR are both high (~27 and 7, respectively) and show a change to lower values in the upper core (~25 and 2, respectively). The ratio  $P_{aq}$  shows little overall variation through core RM-D4, with values ranging between 0.18 and 0.24.

## 5 Discussion

### 5.1 Characterisation of modern sediments and post-depositional processes

Substantial, long-term shifts across bulk and molecular organic geochemical proxies are evident in sediment cores RM-D4 and RM-S1 from Rostherne Mere (Figure 4 and 5). To better understand the primary driving influences and historical context of the observed changes, it is important to be able to determine the mechanisms underlying proxy variations by examining the organic matter that contributes to the lake sediments through end-member characterisation and evaluating the potential for post-depositional changes.

Previous studies on the composition of organic matter in Rostherne Mere suggest the predominance of organic material transferred to recent sediments comprises autochthonous components (Gaskell & Eglinton, 1976; Cardoso et al., 1983). This is confirmed by the uniform *n*-alkane maxima at  $nC_{17}$  observed across sediment trap data (Figure 3), suggesting a principal abundance of algal material (Clark & Blumer, 1971; Cranwell et al., 1987). Diatom blooms between March and June can account for over 90% of annual diatom growth (Livingstone & Reynolds, 1981), and have been dominated in recent years by the species *Asterionella formosa* and *Aulacoseira granulata* (Reynolds, 1978; Moss et al., 2005). In

summer-autumn months, the most prevalent taxa of phytoplankton are the cyanobacteria *Aphanizomenon flos-aquae* and *Anabaena* spp., as well as the cryptophyte *Cryptomonas* (Moss et al., 2005). Trap sediments collected in June and September therefore represent end-member values for the major phytoplankton blooms in Rostherne, where the diatom-dominated assemblage (April-June) has  $\delta^{13}\text{C}_{\text{org}}$  of  $\sim -31$  ‰ and the cyanobacteria-cryptophyte assemblage (June-September) has  $\delta^{13}\text{C}_{\text{org}}$  of  $\sim -28$  ‰, and both have C/N typical of aquatic organic matter of between  $\sim 7$  and  $8$  (Table 1). The difference in  $\delta^{13}\text{C}_{\text{org}}$  between the two collection periods of  $+3$  ‰ is most likely driven by the removal of  $^{12}\text{C}$  from the dissolved inorganic carbon pool by productivity-driven export to sediments during summer months. High HI values of  $\sim 400$  for the shallow traps are also indicative of a kerogen-rich algal source of organic material, and are consistent with the deeper trap sediments collected through June-September. However, lower HI values ( $\sim 370$ ) from the deeper trap during the collection between April and June, which, alongside a higher OI, suggests that greater degradation occurs within the water column upon sinking and may be related to the onset of stratification and development of anoxia. In the months before anoxia takes hold throughout the hypolimnion and extends to the thermocline, reaching a maximum in late September (Davison & Woof, 1984), sinking organic matter derived from algal blooms will be exposed to a more oxygenated water column and hence there is a greater potential for oxidation. This implies that degradation in the water column may lead to lower HI and higher OI values of sinking organic matter when compared to fresh in situ material, as described by Meyers & Lallier-Verges (1999), however overall interpretations of material provenance (i.e. terrestrial vs. aquatic sources) are still valid.

A comparison of trap sediments with the uppermost samples of core RM-D4 (rather than core RM-S1 due to coring location and month of recovery) illustrates the transition and incorporation of fresh organic material from the water column into the sedimentary record.



As mean annual sedimentation rates are on average  $0.7 \text{ cm year}^{-1}$  (based on the  $^{210}\text{Pb}$  chronology of core RM-D4), the uppermost core sediments from the deep water site should represent approximately one annual cycle of sediment deposition. The low  $\delta^{13}\text{C}_{\text{org}}$  of  $-29.4$  ‰ in core top sediments is indicative of a predominantly algal source of organic matter, which is likely dominated by diatom and cyanobacteria material. However, although  $\delta^{13}\text{C}_{\text{org}}$  is consistent between trap and sediment cores, C/N and HI show a change to higher (7.6 to 9.6) and lower (390 to 316) values, respectively, in core top samples. C/N may potentially vary due to a differential loss rates between C and N in the sediment as part of early diagenetic processes, where N is lost at a higher rate than C (Gälman et al., 2008). While this may be a factor in producing the difference in C/N observed at Rostherne, the distribution of *n*-alkanes in the uppermost sediments indicates a source of higher C/N organic material is more likely the cause for divergence between the water column and sediment record. The alkane distribution for core top samples shows a distinct peak at *nC*<sub>17</sub>, relating to a predominant component of algal material. Yet there is a clear secondary maxima centred on *nC*<sub>29</sub> suggesting the presence of longer-chained hydrocarbons and indicating the delivery of non-algal material to the sediments, which would also contribute to lower HI values in core tops (Talbot & Livingstone, 1989).

Although the distinctive alkane maxima at *nC*<sub>17</sub> in trap sediments confirms a dominant algal signal, subsequent smaller peaks are also observed between *nC*<sub>25-33</sub>, which is concordant with the presence of longer-chained hydrocarbons in core top samples. Submerged aquatic macrophytes, mainly comprising *Callitriche* at Rostherne (Carvalho, 1993), show *n*-alkane peaks between *nC*<sub>27-31</sub> corresponding to the pattern found in core tops and traps, but also have major peaks at *nC*<sub>21</sub> and *nC*<sub>23</sub>. Maxima at these chain lengths are not found in the trap or core top sediments and therefore it is unlikely that *Callitriche* is one of the major contributors to the Rostherne organic pool. This is still found to be the case even when post-depositional

processes are considered as, although the samples subjected to degradation in both oxic and anoxic environments show a preferential loss of the shorter-chained fraction (carbon maximum shifts from  $nC_{23}$  to  $nC_{27}$ ), clear peaks are still found at  $nC_{21}$  and  $nC_{23}$ . This is also the case for *Typha* root samples, which show dominant peaks at  $nC_{23}$  and  $nC_{25}$ . The leaves and stems of *Phragmites* and *Typha* have an *n*-alkane distribution more consistent with that of trap and core top samples, however, any contribution would have to be minor given their high TOC, C/N and S<sub>2</sub>, which are significantly higher than all trap and core samples (Table 1 and Figure 6). Following degradation, the distribution of *n*-alkanes from plant samples remains fairly constant (Figure 3), and therefore no overall signal change would be expected due to early diagenetic processes. Although diagenetic changes do not appear to influence the *n*-alkane distribution in samples, total *n*-alkane concentrations are observed to change between fresh and degraded samples indicating that a component of the plant material was lost. The process of selective degradation appears to have a more pronounced influence on bulk parameters, such as C/N and S<sub>2</sub> (Figure 6), though even after degradation the geochemical separation between autochthonous and allochthonous organic constituents remains distinct and traditional interpretations of material source differentiation are still valid (Meyers & Teranes, 2001). One exception are the values for *Callitriche*, which indicate degraded samples assimilate  $\delta^{13}C_{org}$ , C/N, and S<sub>2</sub> values that are congruous with cores and traps (Figure 6). Nonetheless *Callitriche* most likely provides a negligible contribution to the sediment architecture given its *n*-alkane distribution.

A further source of organic material to the lake and its sediments is from soil. The transfer of soil organic matter from catchment to the lake is most likely dependent on inflow from Rostherne Brook, accounting for ~75% of water input, with secondary input via surface runoff from surrounding land through small streams and drainage ditches (Carvalho, 1993; Carvalho et al., 1995). The *n*-alkane composition of soils from the inflow river channel shows

peak concentrations at  $nC_{29}$  with lesser peaks at  $nC_{27}$ ,  $nC_{31}$  and  $nC_{25}$ , which is broadly similar to the distribution associated with woodland and grassland soils, and is representative of the distribution observed across longer-chained homologues in trap and core top sediments (Figure 4 and 5). Although woodland soil has high C/N and  $\delta^{13}C_{org}$ , soils from inflow and grassland catchment localities have a relatively low organic matter content (low TOC and S2; Figure 6) and have  $\delta^{13}C_{org}$  and C/N similar to core tops (Table 1). This suggests that these soils may supply the main component of the lake's allochthonous sediment. Further validation of organic matter provenance can be ascertained using the S2 pyrograms (Figure 3). Fresh plant samples have a unimodal distribution characterised by peaks at low pyrolysis temperatures (330-375 °C), whereas trap and core top samples show a bimodal distribution with maximum hydrocarbon abundance around 360 °C and a secondary, broader peak of lower hydrocarbon abundance extending between ~400 and 500 °C. The only samples to be characterised by a strong bimodal distribution are those from soils, which are most likely due to the incorporation of high thermal stability organic matter such as lignin and tannin (Carrie et al., 2012; Newell et al., 2016). This suggests that Rostherne surface sediments are characterised by both low and high thermal stability components, indicating that input of organic materials is dominated by the autochthonous production of algal matter with a contribution from soils and that marginal vegetation belts most likely have a restricted influence on sediments. Given that the trap collections between April and September are significantly dominated by algal-derived organic matter (Figure 3), the higher thermal stability components found in core top sediments are most likely transferred during winter months, which may account for a proportion of geochemical offset between the trap and core sediments.

## 5.2 Organic geochemistry of the sediment record

The two cores from Rostherne were retrieved in shallow (RM-S1; oxic) and deep (RM-D4; anoxic) water areas of the lake, and so can provide information on spatial differences in sedimentation across the basin and the influence that redox conditions have on sediment proxies. Considering the different sedimentation rates and, therefore, sampling resolution for each core, the overall profile of all proxies is largely consistent both in terms of the magnitude of observed changes and in absolute values between the two coring locations (Figure 4 and 5). Core RM-D4, in comparison to core RM-S1, shows lower average TOC, C/N, and HI between 1955 and present day (given the common chronological  $^{210}\text{Pb}$  tie point; Table 1), which may suggest that a greater proportion of the deeper water sediments comprise a higher concentration of lithogenic material with a detrital origin. This is corroborated by similar average  $\delta^{13}\text{C}_{\text{org}}$  and OI for RM-S1 and RM-D4 (around  $-28.6\text{‰}$  and 178, respectively). Rostherne sediments therefore indicate that proxy values are generally consistent across both shallow and deep water, and oxic and anoxic depositional environments, and that sediment focussing does not overly influence the interpretation of the records.

The surface sediments of Rostherne Mere are shown to comprise predominantly of organic matter dominated by autochthonous components, however shifts in proxy values are observed in deeper horizons (Figure 4 and 5) which is also indicated by previous investigations (Cardoso et al., 1983; Prartono & Wolff, 1998). Lower  $\delta^{13}\text{C}_{\text{org}}$  and C/N, and higher TOC and HI in sediments from the upper parts of the core are consistent with a productive, eutrophic lake system and an algal-dominated source of organic matter. However, towards the lower core, higher  $\delta^{13}\text{C}_{\text{org}}$  and C/N, and lower TOC and HI may reflect either a shift in the balance of organic material being supplied to the lake from different sources, where allochthonous material was more dominant in past years, or potentially could be the result of diagenetic

modification to the organic matter over time. The change from higher to lower  $\delta^{13}\text{C}_{\text{org}}$  through both cores may also be partly attributed to the lowering of  $\delta^{13}\text{C}$  atmospheric  $\text{CO}_2$  due to anthropogenic burning of fossil fuels (the Suess effect; Keeling, 1979).

Post-depositional diagenesis has been shown to have no significant impact on the  $\delta^{13}\text{C}$  of algal organic matter (Hodell & Schelske, 1998; Meyers & Lallier-Verges, 1999), and although changes in the molecular composition of vascular plant material are observed,  $\delta^{13}\text{C}$  is also consistent between fresh and buried allochthonous materials (Meyers et al., 1995). As with  $\delta^{13}\text{C}$ , C/N values are considered to be typically representative of the original material, where changes are not of a sufficient magnitude to override larger-scale source determinations (Meyers & Ishiwatari, 1993). A relative change in C/N has been shown to be driven by the selective loss of N over C at a rate of  $\sim 1:1.5$ , leading to increasing C/N through time (Gälman et al., 2008). However, the rate of diagenetic loss of both C and N is highest within 5 years after deposition (20% and 30%, respectively), and reduces considerably over time with a maximal C/N difference of +2 between recent and degraded sediments (Gälman et al., 2008). The observed  $\delta^{13}\text{C}_{\text{org}}$  and C/N change of approximately +1.6 ‰ and +4, respectively, from the upper to lower sediments of cores RM-S1 and RM-D4 (1955-present) therefore suggests that the balance between autochthonous and allochthonous components is shifting downcore and recording palaeoenvironmental change, rather than primarily reflecting the influence of diagenetic processes.

To assess further the change in organic constituents, S2 pyrograms (the main constituent of HI) and *n*-alkane distributions are plotted in Figure 4 and 5. Pyrograms from the uppermost sediments of cores RM-S1 and RM-D4 show a bimodal distribution across the S2 temperature range of 300 to 600 °C, where shallower sediments, as previously discussed, are characterised by a distinct peak at a pyrolysis temperature of around 360 °C (S2a) and a second less pronounced peak (S2b) at the higher pyrolysis temperature of 460 °C. A strong

S2a peak form indicates the presence of organics with a low-moderate thermal stability, such as those derived from lipid cell wall materials, and therefore implies a signal driven by algal-derived organic matter (Carrie et al., 2012). This is confirmed by the presence of short-chained hydrocarbons in both cores (Figure 4 and 5), in particular the peak observed at  $nC_{17}$ , which is commonly associated with algal-dominated sediments (Clark & Blumer, 1971). In both RM-S1 and RM-D4 there is a clear change from the upper to mid-core depths, where both cores show a marginally reduced hydrocarbon abundance in contrast to upper sediments and a bimodal distribution characterised by a stronger S2b peak at 460 °C. In the lowermost sediments, there is a considerable reduction in hydrocarbon abundance observed for both cores when compared to upper- and mid-levels, and the S2 peak structure becomes broader and less defined between S2a and S2b, in particular for core RM-D4. A shift to an S2b-dominated pyrogram form suggests a lower relative abundance of algal matter in sediments and a greater contribution of higher thermal stability elements (Carrie et al., 2012). This implies that these sediments contain a component of terrestrial material, which is validated by the reduction in short-chained hydrocarbons in *n*-alkane profiles and a shift to a long-chained-dominated assemblage between  $nC_{27}$  and  $nC_{33}$ .

This change is also illustrated in the *n*-alkane ratios ACL and TAR (Figure 4 and 5). Higher ACL and TAR in sediments from the lower core indicate a terrestrial dominance of organic matter content (Bourbonniere & Meyers, 1996; Bush & McInerney, 2013) and confirm a greater proportion of long-chained hydrocarbons are present. Contrastingly, the upper core is characterised by lower ACL and TAR indicating an up-core change from a relatively low abundance to a higher content of lake-derived organic matter. This, together with a parallel rise and strong covariance between TOC and HI ( $R > 0.8$  for both cores), suggests that the signal is driven by an increasing contribution of autochthonous organic matter rather than terrestrial inputs diminishing (Meyers, 2003; Steinmann et al., 2003). A change to

environmental conditions favouring increased aquatic productivity in the upper core is further supported by a transition to lower C/N indicating a greater contribution from non-vascular algal organic matter (Meyers & Teranes, 2001). In addition to lower C/N, increasing MinC through the cores implies greater endogenic carbonate precipitation most probably in response to increased phytoplankton activity and photosynthetic uptake of dissolved CO<sub>2</sub>, thereby driving calcium carbonate super saturation in the water column (Ito, 2001; Leng & Marshall, 2004; Scott, 2014).

### *5.3 Impact of anthropogenic environmental change*

At the base of core RM-S1, relatively high TOC (>10 %) and moderate HI (>230) suggests that Rostherne was a productive lake and had likely developed enriched conditions before the end of the 19<sup>th</sup> century (Figure 7). This is supported by an extended history of human activity and catchment modification in the local area, including agriculture, habitation of surrounding villages, and infrastructure development. Cultural eutrophication was most likely strengthened following a period of forest clearance and agricultural development after ca. 600 BP (Nelms, 1984), with long-term monitoring and sedimentary diatom analysis confirming that enriched conditions have existed since at least 1912 (Pearsall, 1923). TOC is relatively consistent at the base of core RM-S1 up to ca. 1953, after which there is a substantial shift to higher TOC and HI suggesting greater organic matter accumulation (Figure 7). This shift is also observed in  $\delta^{13}\text{C}_{\text{org}}$  and C/N which are stable until ca. 1953, followed by a marked trend towards the lower values of present day. Lower, but consistent, TOC and higher C/N and  $\delta^{13}\text{C}_{\text{org}}$  at the base of the core correspond to lower accumulation rates of planktonic diatoms and organic carbon burial before the 19<sup>th</sup> century until the mid-20<sup>th</sup> century (Nelms, 1984; Radbourne et al., 2017), which is implied to reflect reduced productivity compared to more recent conditions in Rostherne Mere.

The comparatively rapid shift of the mid-1950s observed in core RM-S1 is not seen in the corresponding interval of core RM-D4, which has a lowermost  $^{210}\text{Pb}$  date of  $1955 \pm 8$  years (Figure 2). A reduction in sedimentation rate between ca. 1955 and 1964 is observed in both cores, as are the relatively stable sedimentation rates between ca. 1964 and present day, suggesting the dating on the two cores is comparable and well-resolved subsequent to 1955. The rapid TOC increase in core RM-S1 between 1953 and 1955 may correspond to a similar increase between 1940 and 1947 in core RM-D4, which is substantiated by similar trends in C/N,  $\delta^{13}\text{C}_{\text{org}}$ , and HI through the same interval (Figure 7). The base of core RM-S1 has larger chronological errors associated with the  $^{210}\text{Pb}$  chronology (Figure 2), and a high sedimentation rate in core RM-S1 through this zone implies the potential for sediment slumping and/or an erosional event, which may explain the discrepancy between the cores. However, there is an overall distinct shift in TOC from the lower part of RM-S1 dated to the early 20<sup>th</sup> century (ca. 1919-1933) characterised by low TOC (~11%) to the mid-20<sup>th</sup> century (~14%), in addition a large perturbation in TOC is also observed in RM-D4 around ca. 1935 (Figure 7). This pronounced change in organic carbon content is most likely related to the construction of a local sewage treatment works in 1935 that discharged effluent upstream of Rostherne until its diversion in 1991 (Carvalho et al., 1995). Nutrient budgeting has shown that the effluent comprised around 70% of the external nutrient load from the Rostherne Brook inflow (Moss et al., 1997), which suggests a considerable increase in nutrient (P) loading following the construction of the treatment plants.

In RM-S1 and RM-D4,  $\delta^{13}\text{C}_{\text{org}}$  remains moderately consistent at  $-28\text{‰}$  until ca. 1960 and then starts to progressively decrease, which is also observed as a trend to higher TOC and HI, and lower C/N (Figure 7). These changes are broadly concurrent with an abrupt transformation of the lake's cyanobacteria assemblage which took place around 1958 (Livingstone, 1979), as well as diatom replacement by species characteristic of very eutrophic



waters and a reduction in the accumulation of non-planktonic diatoms (Nelms, 1984).

Although a primary cause of this change may be related to a transition from phytoplankton being N-limited to a state where populations were instead controlled by light availability (Reynolds & Bellinger, 1992), it is suggested that this ultimately was driven by an increase in the use of synthetic agricultural fertilisers in turn leading to further nutrient enrichment in the lake (Reynolds, 1979). Many lakes around the UK and Europe similarly provide evidence for eutrophication starting in the mid-19<sup>th</sup> century, often with a more distinct phase since c. 1950 (Battarbee et al., 2011; Bennion et al., 2012; Anderson et al., 2014; Heathcote et al., 2015). This change in TOC and  $\delta^{13}\text{C}_{\text{org}}$  is gradual and of a lesser magnitude compared to the effects of sewage effluent, which is most likely associated with a lower contribution to external nutrient loading from agricultural runoff (up to 27 %; Moss et al., 1997). This period is followed by overall higher TOC, and decreasing C/N and  $\delta^{13}\text{C}_{\text{org}}$ , which is associated with an increasing sedimentation rate through until ca. 1980 (Figure 7).

In the early 1980s, an increase in TOC and an overall rapid decrease in  $\delta^{13}\text{C}_{\text{org}}$  and C/N is observed in both cores (Figure 7). A higher rate of decreasing  $\delta^{13}\text{C}_{\text{org}}$  and C/N, and higher TOC, suggests sediments contain an increasing proportion of autochthonous organic matter, which is consistent with lower TAR and ACL (Figure 4) and is simultaneous with an abrupt increase in the rate of organic carbon burial after ca. 1980 (approximately +15 %, Figure 7; Radbourne et al., 2017). These proxy changes likely reflect a period of peak eutrophication perhaps due to a further increase in nutrient loading from the sewage treatment works, given the similarity with the intensification of eutrophication following their construction in the 1930s. Originally, the treatment works served a population of 550 people, however by the 1980s they had become overloaded due to local housing development (Moss et al., 2005), and by 1990 served a population of 3350 people (Carvalho et al., 1995).

Following the transition to lower  $\delta^{13}\text{C}_{\text{org}}$  from ca. 1980, a change to relatively stable  $\delta^{13}\text{C}_{\text{org}}$  is observed between ca. 1994 and present day in parallel with a continued C/N decrease in both cores (Figure 7). This broad-scale change follows the closure of the sewage treatment works in 1991, however the shift from decreasing to low and stable  $\delta^{13}\text{C}_{\text{org}}$  in both cores post-dates the diversion of effluent by several years. Although external nutrient loading fell dramatically following diversion, where the phosphorus load in Rostherne Brook reduced by ~78 %, total outputs from the lake remained constant (Carvalho et al., 1995). In contrast to upstream Little Mere, where there was a distinct and immediate decline in total P following sewage diversion, substantial internal nutrient loading in Rostherne resulted in increased lake concentrations of total P between 1992 and 1993 (Carvalho et al., 1995), and only a limited decline in concentration over the next decade (Moss et al., 2005). Since ca. 1994, stable  $\delta^{13}\text{C}_{\text{org}}$  around  $-29.5 \text{ ‰}$  are consistent with modern trap sediments, which implies that although external nutrient loading has been greatly reduced, internal nutrient loading has likely resulted in a slow remediation of hyper-eutrophic conditions and restricted ecological recovery.

## 6 Conclusions

- The dominant source of organic matter to the sediments of Rostherne Mere is derived from autochthonous material, which is based on maxima in n-alkanes at  $n\text{C}_{17}$  and overall low  $\delta^{13}\text{C}_{\text{org}}$ , C/N, and high HI values in trap and core top data. Catchment soils also contribute to the sediments, most likely transferred during winter months, as suggested by differences in HI and C/N between traps and core tops, and the presence of longer-chained hydrocarbons in core tops.
- Conventional interpretations of organic matter provenance remain valid after sinking and burial, as changes in the values of geochemical proxy data are minor. Early diagenetic processes in core sediments do not overprint the signal of changing

palaeoenvironmental conditions, which is important for setting recent anthropogenic-induced changes in a historical context. Sediment focussing and differential redox conditions across the lake basin also do not exert considerable influence over the general profile of excursions observed in multi-proxy records.

- Autochthonous organic matter production in the lake increased over the 20<sup>th</sup> century and the transition to hyper-eutrophic conditions took place at a variable rate. This was principally dependent on the delivery of point-source sewage effluent and enhanced phosphorus loading, with key increases in productivity following the construction of a sewage treatment works in mid-1930's, the introduction of synthetic fertilisers in the late 1950's, and population growth through to the 1980's. In recent years, remediation has reduced the influence of external nutrient loading, however internal nutrient loading has slowed ecological recovery.
- Overall, this study has shown how sedimentary organic matter can sensitively record recent eutrophication processes in lakes. It highlights the efficacy of using a combination of geochemical techniques, in particular  $\delta^{13}\text{C}_{\text{org}}$ , Rock-Eval, and *n*-alkane biomarkers, and the need integrate modern end-member calibrations and lacustrine sediment cores to better understand the timing and scale of environmental change through the Anthropocene.

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## Figure captions

Figure 1: A) Location of Rostherne Mere, Little Mere and Mere Mere, and the sites of Rostherne and Mere sewage treatment works (STW), and B) Map of Rostherne Mere with bathymetric data (in meters; Scott, 2014), showing the coring locations of RM-S1 (N53°21'28.7", W2°23'22.1") and RM-D4 (N53°21'12.6", W2°23'19.4"; Map data: Google, Bluesky).

Figure 2: Radiometric chronology versus depth for cores RM-S1 and RM-D4, showing the CRS model  $^{210}\text{Pb}$  dates (black squares) with respective error bars and sedimentation rate (grey circles). Fallout radionuclide concentrations versus depth are also shown for total  $^{210}\text{Pb}$ , supported  $^{210}\text{Pb}$ , unsupported  $^{210}\text{Pb}$ ,  $^{137}\text{Cs}$ , and  $^{241}\text{Am}$ . A zone of high sedimentation rate in RM-S1 implies the potential for sediment slumping, which is highlighted by the horizontal grey bar.

Figure 3: Chain-length distributions of *n*-alkane biomarkers (black bars) and Rock-Eval S2 pyrograms (grey lines) of modern sediment traps (water depth of shallow = 8 m, deep = 22.5 m), catchment soils (inflow/woodland/grassland), and fresh and degraded (oxic/anoxic) vegetation samples (*Phragmites/Typha/Callitriche*) samples. S2 pyrogram y-axis values (abundance of hydrocarbons in mg HC/g TOC) are given as integrated peak area (Hydrogen Index) in Table 1.

Figure 4: Organic geochemistry of sediment core RM-S1, showing MinC (equivalent to TIC), TOC, C/N,  $\delta^{13}\text{C}_{\text{org}}$ , HI, OI, ACL,  $\text{P}_{\text{aq}}$ , and TAR.  $^{210}\text{Pb}$  ages are shown to the right of the diagram. Grey bar marks area of high sedimentation rate. Rock-Eval S2 pyrograms are shown for three horizons (each pyrogram represent the given core depth) with *n*-alkane biomarker distributions (biomarker samples were combined from the sample depths given in the pyrograms).

Figure 5: Organic geochemistry of sediment core RM-D4, showing MinC (equivalent to TIC), TOC, C/N,  $\delta^{13}\text{C}_{\text{org}}$ , HI, OI, ACL,  $\text{P}_{\text{aq}}$ , and TAR.  $^{210}\text{Pb}$  ages are shown to the right of the diagram. Rock-Eval S2 pyrograms are shown for three horizons (each pyrogram represent the given core depth) with *n*-alkane biomarker distributions (biomarker samples were combined from the sample depths given in the pyrograms).

Figure 6: Plot of  $\delta^{13}\text{C}_{\text{org}}$  versus C/N (top) and Rock-Eval modified Van Krevelen diagram of S2 versus TOC (bottom) for sediment trap, vegetation, soil, and core samples. Trap sediments for June and September are split between shallow (S) and deep (D) water samples. Blue dashed boxes show typical  $\delta^{13}\text{C}$  and C/N values of major sources of lacustrine algae and  $\text{C}_3$  land plant organic matter to lake sediments (Meyers & Teranes, 2001). The boundaries between organic matter types are defined as HI = 700 and HI = 200 after Langford & Blanc-Valleron (1990).

Figure 7: Comparison of TOC, C/N,  $\delta^{13}\text{C}_{\text{org}}$ , HI, and sedimentation rate from cores RM-S1 and RM-D4 plotted against calendar age, showing organic carbon (OC) burial rate (Radbourne et al., 2017) and the onset and cessation of sewage effluent delivery to the lake (vertical grey lines). The lowermost sedimentation rate for each core was extrapolated to the base of each sequence to provide a chronological framework for the full record.

Table 1: Geochemical, isotope, and Rock-Eval composition of modern vegetation (fresh and oxic/anoxic degraded), sediment trap, and catchment soil samples. Averages between 1955 and present day for sediment cores RM-S1 and RM-D4 are provided. The *n*-alkane ratios ACL,  $\text{P}_{\text{aq}}$ , and TAR are also given for each sample.