Mobilisation of iron and manganese from sediments of a Scottish Upland reservoir
Corinna ABESSER* and Ruth ROBINSON

School of Geography and Geosciences, University of St Andrews, St Andrews, KY16 9AL, UK
*e-mail corresponding author: c.abesser@bgs.ac.uk

ABSTRACT

High concentrations of manganese (>50 μg L⁻¹) and iron (>226-467 μg L⁻¹) have caused water quality problems at the Megget Reservoir during an excessive draw down of the reservoir water level in 1997/98. Sediment column studies were carried out to investigate how internal processes, namely sediment resuspension and diffusion from the sediment, contribute to the mobilisation of these elements in the water column. It was found that sediment resuspension is by far the more effective mechanism in mobilising iron and manganese and also causes persistent discoloration of the water. Concentrations of up to 80 mg L⁻¹ iron and 16 mg L⁻¹ manganese were observed in the experimental core water columns which are equivalent to 490 μg L⁻¹ and 97 μg L⁻¹ in a 27m water column (= maximum water depth during reservoir draw down in 1997/98). From sediment particle size data it was calculated that wind speeds over 13 m s⁻¹ can trigger sediment resuspension at water depths of up to 20 m and have the potential to disturb sediments in large parts (>25% at full capacity) of the reservoir. Under current climatic conditions, such wind events are infrequent but increasing wind speeds and rainfall intensities, as predicted by climate change scenarios, may cause resuspension events to become more frequent, reducing the reservoir’s water quality over prolonged periods and limiting its use as a drinking water resource. Such resuspension events are also expected to occur in other water bodies in the study region, adversely affecting their water quality and increasing the ecosystem productivity. Diffusion from the sediment mostly affects the cycling of manganese. Maximum concentrations in the experimental columns were equivalent to 70-130 μg L⁻¹ in a 3-5 m water column and were comparable to in-situ concentrations measured in the bottom waters of the reservoir (3-5 m above the sediment) during summer stratification. Sediment release is unlikely to cause serious water quality problems as only a limited amount of available manganese is found within the sediment. However, the accumulation of manganese and iron in the bottom waters may increase with changing climatic condition, in particular when the external inputs of these elements increase as a result of higher catchment loading associated with the predicted rise in rainstorm intensities.

Key words: iron, manganese, sediment resuspension, diffusion, drinking-water quality, reservoir, laboratory study

1. INTRODUCTION

Manganese (Mn) and iron (Fe) are present in water supplies as a result of natural processes involving both catchment erosion and redox-related dissolution of Fe- and Mn-containing minerals at or near the sediment-water interface. Increased concentrations of these elements in water supply bodies concerns many water authorities in the UK (Heal 2001) and has also caused problems in Australia (Zaw & Chiswell 1999), the Ukraine (Linnik & Zubenko 2000) and the USA (Hsiung & Tisue 1994) where approximately 40% of the public water supply systems are affected (AWWA 1987). Temporary deterioration in water quality due to increased Mn has occurred in raw waters from the Ellan valley in Wales (Schofield et al. 1991) and in Loch Bradan, south-west Scotland (Little & Mcfadzean 1991). Increased Mn concentrations were also found in the Megget Reservoir during a forced draw down of the reservoir water level in 1997/98 from 55 m (full capacity) to 27 m (28% capacity) in order to conduct repairs to the dam. As a result, concentrations at the water treatment works increased to 50 μg L⁻¹ for Mn and to 350 μg L⁻¹ for Fe, approaching and exceeding the maximum admissible concentrations in UK drinking water of 50 μg L⁻¹ and 200 μg L⁻¹, respectively (DWI 2000). Problems with high Mn concentrations in the drinking water have forced the introduction of filtration systems and Mn removal stages at various water treatment works in Scotland and have led to regular, controlled flushing of the distribution system being carried out to reduce Mn deposits in the pipes.

While catchment inputs from streams provide the primary, external loading of Fe and Mn to reservoirs and lakes, internal processes, both biochemical and physical, can be responsible for increased concentrations in the water column as a result of remineralisation and diagenetic (redox-controlled) release from the sediments. High concentrations of Mn in Loch Bradan (Gavin et al. 2001), for example, have been attributed to the dissolution of Mn-containing minerals at or near the sediment-water interface, and sediment core studies have shown that release of Mn from sediments is a common phenomenon in Scottish freshwater lochs (Bryant et al. 1997). Mobilisation of Fe and Mn during early diagenesis is coupled to suboxic conditions in the sediment where Fe(III) and Mn(IV), present in form of Mn and Fe (hydr)oxides, provide an electron acceptor for the microbial degradation of reactive organic matter (Froehlich et al. 1979; Lovley & Phillips 1986). In the
process, Fe(III) and Mn(IV) are reduced to the more soluble Fe(II) and Mn(II) and released into the porewaters from where they diffuse upwards towards the sediment-water interface. Where redox conditions are favourable, this can provide a constant input of dissolved species into the overlying water column (Davison et al. 1982; Warnken et al. 2001).

Advective transport, including sediment resuspension, resulting from wind/wave-induced turbulences has also been linked to increased metal (and nutrient) concentrations in the water column of aquatic systems (ten Hulscher et al. 1992; Brassard et al. 1994). Resuspension events can be generated by exposure of deep-water regions to higher energy regimes (e.g., wind or water movement) and often occur in reservoirs where large water level fluctuation and regular draw down events are frequent (Effler et al. 1998). Excessive reservoir draw down was held responsible for the increased metal release in the Dnieper reservoirs (Linnik & Zubenko 2000).

In this study, laboratory-based experiments were carried out to investigate (1) how a reduced oxygen supply to the water column (as experienced during summer stratification) affects the sediment release of Fe and Mn, and (2) what impact sediment resuspension (as experienced during autumn turnover/storm events), and related metal mobilisation from sediments, has on the water quality of the overlying water column. Undisturbed sediment cores were installed in the laboratory and subjected to various treatments simulating different internal and external forcing conditions (e.g., oxygen depletion and shear stress). This paper discusses the laboratory-based experiments and their implications for reservoir management.

2. STUDY AREA

The Megget Reservoir (Fig. 1) is located in the Southern Uplands in the UK, about 50 km south of Edinburgh (NGR 319 622). It is an important source of drinking water and one of Scottish Water's largest supply reservoirs. It has a surface area of 2.59 km² and its yield to supply (at the time of construction in 1983) was estimated as 102.3 MI d⁻¹ (Jowitt & Hay-Smith 2002).

The reservoir is a monomictic, oligotrophic system. Its catchment area is underlain by Fe and Mn-rich Ordovician and Silurian greywackes and shales which are locally covered by glacial and post-glacial deposits (e.g., peat). The soils are predominated by organic-rich peaty podzols, peaty gleys and blanket peat and to a lesser degree by mineral soils including brown forest soils, rankers and podzols. The reservoir is fed by six major tributaries which, during periods of increased runoff, deliver high loads of organically-complexed Fe and Mn to the reservoir (Abesser et al. 2006). The deepest part of the reservoir is situated towards the east with a maximum depth of 55 m at the dam (at full capacity). The western part is relatively shallow with alluvial fans developed around areas of stream inflows. During periods of thermal stratification, bottom waters remain oxygenated but dissolved oxygen (DO) is depleted progressively from the lower layers of the reservoir (minimum DO 7.6 mg L⁻¹ at 1 m above reservoir bottom). Distinct vertical gradients in total Fe and Mn concentrations develop during late summer/early autumn and differences between surface and bottom waters of up to 150 µg L⁻¹ (Fe) and 100 µg L⁻¹ (Mn) have been observed prior to mixing and overturn of the reservoir (Abesser 2003).
The reservoir sediments consist of grey-brown silty muds which dominate the central parts of the reservoir and are intermixed with fine sands closer to the shores. The organic carbon content (as determined by Loss on Ignition) of the surficial sediments (top 10-15 cm) is relatively high and can make up a quarter (24%) of the total dry mass (Abesser 2003). The sediment chemical composition varies between the shallow end of the reservoir and the deeper parts (in the middle and dam area) indicating variations in the proportions of feldspar, clay minerals, quartz and biogenic components in these sediments.

The Southern Uplands have a predominantly cool and temperate climate that is dominated by frontal systems from a general Atlantic source (mainly westerly direction). Wind speeds average around 3.9 m s⁻¹ (median value calculated from hourly data between 01 September 1999 and 30 September 2001 measured at the Environmental Change Network site at Sourhope (NGR 387 622)) but strong winds from the north prevail during the winter and can exceed speeds of 20 m s⁻¹.

3. METHODS

3.1. Core collection

Six undisturbed sediment cores were collected from the Megget Reservoir on 4th October 2001 at two different sites (A and C) within the reservoir (see Fig. 1). At the time of core collection, the reservoir was well-mixed with no temperature or chemical stratification.

The sediment cores were collected with a Jenkins core sampler into 50 cm-long perspex tubes with an inner diameter of 5 cm. The sampler captures the upper 10–30 cm of the sediment and a corresponding column of overlying water without disturbing the core or the sediment interface (Mortimer 1971). Prior to core collection, 10 litres of reservoir water were collected in a sample-rinsed polyethylene container for use during the laboratory experiments.

On collection, the core quality was inspected and cores were arranged according to their future use. Two cores were used for (immediate) chemical sediment analysis and were prepared for dissection by removing the water overlying the sediment with a siphon. The final 5 cm of the water column (above the sediment) were collected into clean sample-rinsed polyethylene bottles for chemical analysis. The cores were then dissected into 1-cm segments. The individual segments were transferred into pre-labelled plastic bags and stored in a cool box until they were transferred to the cold store. The cores collected for laboratory experiments remained sealed and were transported to the laboratory and stored in the cold store at 9 °C until the next day when they were required for the experimental setup. The time between collection of the cores and the first experimental sampling/measurements was <3 days.

3.2. Experimental settings

The four undisturbed sediment cores were installed in the laboratory and subjected to various treatments simulating different internal and external forcing conditions as summarised in Table 1. Cores collected for the resuspension experiment, C1 and C5, were fitted with a wall-mounted motor to which a 20-cm-long stainless steel stirrer with 1-cm mixing blades was attached. The stirrer was adjusted to sit 12-14 cm above the sediment surface. The depth of resuspension was controlled by adjusting the speed of the stirrer and uniformity was achieved by constant revolutions. The cores were left uncovered and exposed to light in order to reproduce the conditions in the shallow areas of the reservoir which are exposed to wind-induced turbulence and prone to resuspension. Cores collected for the sediment diffusion study, C3 and C6, were wrapped in aluminium foil to reduce photosynthetic processes in the water column and then sealed with plastic film to minimise oxygen exchange at the water-air interface during core incubation. Core C3 was maintained at 9 °C ±1 °C in a temperature-controlled room to simulate conditions at the water-air interface during core incubation. Core C6 was kept at a temperature of 21 °C ±2 °C. The cores remained isolated from atmospheric oxygen throughout the incubation period, except for the short intervals of sample collection when the seal was removed for approximately 5 minutes.

The four cores were incubated over a 21-day period. At regular intervals (3-4 days), *in situ* measurements were made of temperature and redox potential. They were taken in the upper 10 cm of the water column in order to minimise column mixing and avoid disturbance of the sediment-water interface. The redox potential (Eh) was measured using a gel-filled Orion electrode (Thermo Scientific Inc.). All measurements were corrected to the mV values that would be obtained by a Standard Hydrogen Electrode (SHE) and converted to pE. Water column samples were collected for the analy-

<table>
<thead>
<tr>
<th>Core ID</th>
<th>C1</th>
<th>C3</th>
<th>C5</th>
<th>C6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Core length</td>
<td>13.5 cm</td>
<td>20.0 cm</td>
<td>11.5 cm</td>
<td>18.0 cm</td>
</tr>
<tr>
<td>Treatment</td>
<td>Resuspension</td>
<td>Diffusion</td>
<td>Resuspension</td>
<td>Diffusion</td>
</tr>
<tr>
<td>(top 4–5 cm)</td>
<td>(9 °C)</td>
<td>(top 1–2 cm)</td>
<td>(21 °C)</td>
<td></td>
</tr>
<tr>
<td>Duration of treatment</td>
<td>21 days</td>
<td>21 days</td>
<td>21 days</td>
<td>21 days</td>
</tr>
<tr>
<td>Sampling frequency</td>
<td>3–4 days</td>
<td>3–4 days</td>
<td>3–4 days</td>
<td>3–4 days</td>
</tr>
</tbody>
</table>
sis of total and dissolved metals as well as pH by slowly lowering a glass tube to a position just above the water-sediment interface. The glass tube was then capped and an integrated sample of the water column was withdrawn. Samples collected for determining the water column pH were emptied into a clean glass beaker and the pH was measured immediately. All other samples were collected into clean polyethylene bottles. For the analysis of the dissolved (soluble-colloidal) metal fraction, samples were filtered through a 0.45 µm pore size filter (Whatman) and acidified to 1%-vv with concentrated Arista® grade nitric acid. The core water column was then replenished with reservoir water that was collected on the day of core collection and stored at a temperature of 9 °C ±1 °C. Prior to replenishing, the water temperature was adjusted to the core water temperatures of 9 °C and 21 °C, respectively. After each measurement and sampling event (i.e. every 3-4 days), cores C1 and C5 were resuspended as follows: Core C1 was resuspended for 1 hour to a depth of 4-5 cm and core C5 was resuspended for 4 hours to a depth of 1-2 cm. On completion of the experiments, the cores were dissected (as previously described 3.1) and the weakly-bound metal (Fe, Mn, Al) fraction was determined as detailed below.

3.3. Analytical methods

3.3.1. Sediment analysis

A single extraction technique was applied to the sediment segments following the operationally-defined procedure by Loring and Rantala (1992) which is detailed in Abesser (2003). This method is based on the use of 25%-vv acetic acid (HOAc) to release the proportion of weakly-bound metals from such sites as ion-exchange positions, amorphous compounds of Fe and Mn, carbonates and those metals weakly held in organic matter. Hence, the extracted fractions should approximate the proportion of reactive Fe and Mn that is available for diagenetic remobilisation, bearing in mind that some differences may exist between such chemical dissolution of metal oxides and the actual process of microbial utilisation of mineral-bound metals (Hyacinthe et al. 2006). By definition, the extracted fraction also includes freshly-formed Fe and Mn hydroxides that precipitated from the pore waters during core segmentation and drying. Consequently, changes in the extracted-metal profiles prior to and after the experiment also reflect changes in metal distribution in the pore waters during the incubation period.

For the (kinetic) extraction, 2 g of oven-dried and crushed (not grinded) sediment were extracted with 25 ml of 25%-vv acetic acid (HOAc) by agitating the sample in an orbital shaker (at 200 rpm) for 6 hours. The supernatant was then decanted and the sediment was washed with 10 ml deionised water to remove the remaining dissolved components. The supernatant as well as wash water were separated from the sediment by centrifugation and decanted into a 50-ml polyethylene container. The extracted sample was then made up to 50 ml with deionised water. All samples were stored at 4 °C in a temperature-controlled room until analysis by ICP-OES. Acidification of the samples was not necessary as the sample extract had an acidity of 10-12%-vv HOAc and the pH was sufficiently low (pH ~2.5) to prevent re-adsorption of the metals to the walls of the storage vessels.

3.3.2. Water sample and supernatant analysis

Analysis of column water samples and sediment extraction (supernatant) samples was carried out using an Inductively Coupled Plasma-Optical Emission Spectrometer (ICP-OES). Water samples with metal concentrations significantly higher than the calibration range were diluted 20-fold using Arista® grade nitric acid as dilutant. Sediment extraction samples were also diluted (1) 10-fold with de-ionised water to reduce the acidity of the sample to 1%-vv (as required by the ICP OES analysis method); and (2) a further 10-fold with nitric acid to reduce metal concentrations to lie within the calibration range of the method.

3.4. Calculating potential for wind-driven sediment resuspension

In order to estimate how wind conditions at the Megget Reservoir might affect sediment resuspension, bottom shear stress values (fluid shear stress acting on the bottom sediment due to wind-induced wave action) and critical entrainment threshold values (critical shear stress value at which sediment of median particle size becomes entrained/resuspended) were calculated from the available wind speed, water depth and particle size data.

Particle size data were available from a sediment survey carried out in August 2000 (Abesser 2003) which included 11 sampling stations across the Megget Reservoir. Water depths at these sampling sites ranged from 9.3 m to 43.9 m (Fig. 1). The wind speed data were recorded at the ECN monitoring site at Sourhope (NGR 387 622) and the fetch for the western end of the reservoir was approximated as 1000 m.

Calculations of bottom shear stress values were based on the method outlined in Lawson et al. (2007):

$$\tau_{h_m} = \rho(f / 2)U_b^2$$

(1)

where \(\tau\) is bottom shear stress (Pa) due solely to waves, \(U_b\) is the orbital velocity, \(\rho\) is water density (kg m\(^{-3}\)), and \(f\) is a friction factor based on wave amplitude and roughness length \(k_s\) (which is assumed to be \(3 \times D_{50}\) where \(D_{50}\) is the grain diameter for which 84% of the sediment sample by mass is finer). Bottom wave orbital velocity and the friction factor were calculated following Lawson et al. (2007):

$$U_b = \frac{H_w \pi}{T \sinh(\frac{2Hz}{\lambda})}$$

(2)
where $H_{\text{sig}}$ is the significant wave height (m), $T$ is the wave period (s), $\lambda$ is the wavelength (m) and $A$ is wave amplitude (m).

Wave height ($H_{\text{sig}}$) and wave periods ($T$) were calculated using Airy Wave theory in the form presented by Prakash (2004):

\begin{align}
H_{\text{sig}} &= 5.112 \times 10^{-4} w F^{3/2} \\
T &= 6.238 \times 10^{-2} (w F)^{1/3}
\end{align}

where $w$ is the wind speed (m s$^{-1}$), $g$ is the acceleration due to gravity (m s$^{-2}$) and $F$ is the maximum effective fetch (m). The over-water wind speed $w$ (m s$^{-1}$) was estimated to be 20% higher than the recorded over-land wind speed for fetches of less than 16 km (Prakash 2004). The shallow water wave equation was then used to calculate wavelength $\lambda$ (m):

$$\lambda = T \sqrt{gh}$$

Critical entrainment of sediment with a grain-size less than fine sand can be calculated following Akahori et al. (2008):

\begin{equation}
\tau = \left( 1 - \frac{D_s}{D_{50}} \right) \frac{\rho_s}{\rho w} \sqrt{\frac{D_s}{\rho w}}
\end{equation}

where $D_s$ is the non-dimensional grain-size, $\rho_s$ and $\rho$ are sediment and water density, respectively, and $D_{50}$ is the median grain size ($D_{50}$). The Shields curve, which is commonly used to estimate critical entrainment threshold values, is known to fit poorly to silts and clays and Cao et al. (2006) have proposed that a non-dimensional critical entrainment threshold $\tau$ can be formulated as:

$$\tau = 0.1414 \left( \frac{D_s}{D_{50}} \right)^{0.2306}$$

when $D_s < 6.61$, which is the case for the median grain sizes in the Megget Reservoir.

4. RESULTS

4.1. Core Resuspension

Iron and Mn concentrations in the water column of cores C1 and C5 increase dramatically during initial resuspension (Figs 2 and 3). They remain fairly constant during subsequent resuspension events and appear to be homogeneously distributed throughout the core water.
Iron and manganese mobilisation

47

column (Figs 2 and 3). Concentrations of total Fe (suspended plus dissolved) in the water column are up to 5 times higher than those of total Mn with maximum Fe concentrations of 79.5 mg L⁻¹ in C1 and 29.4 mg L⁻¹ in C5, compared to 16.3 mg L⁻¹ and 12.6 mg L⁻¹ of Mn. Most Fe is present in particulate form and soluble-colloidal Fe only accounts for 3% to 4% (1.6-3.0 mg L⁻¹) in C1 and 4% to 7% (0.5-1.6 mg L⁻¹) in C5 of the total water column concentrations. The proportion of soluble-colloidal Mn is much higher ranging between 39% and 55% (5.2-6.8 mg L⁻¹) in C1 and between 67% and 76% (5.1-8.8 mg L⁻¹) in C5. The total concentrations of Fe and Mn are higher in core C1, where a higher amount of sediment is suspended in the water column. The proportion of soluble-colloidal species, however, is lower even during reducing conditions where Fe²⁺ and Mn²⁺ are expected to predominate (Figs 4a and 4b).

The observed Fe and Mn concentrations in the resuspended core water columns are very high compared to concentrations in the reservoir due to the high sediment-to-water ratio (0.6-0.8) in these experimental cores. Assuming the dilution of these concentrations through homogeneous mixing of the resuspended sediment into a 27 m water column (= water depths at deepest point during maximum reservoir draw down) the concentrations of total Fe and Mn are higher in core C1, where a higher amount of sediment is suspended in the water column. The proportion of soluble-colloidal species, however, is lower even during reducing conditions where Fe²⁺ and Mn²⁺ are expected to predominate (Figs 4a and 4b).

The observed Fe and Mn concentrations in the resuspended core water columns are very high compared to concentrations in the reservoir due to the high sediment-to-water ratio (0.6-0.8) in these experimental cores. Assuming the dilution of these concentrations through homogeneous mixing of the resuspended sediment into a 27 m water column (= water depths at deepest point during maximum reservoir draw down) the concentrations of total Fe and Mn reduce to 210 μg L⁻¹ and 89 μg L⁻¹ (if the top 2 cm of sediment are resuspended) and to 490 μg L⁻¹ and 97 μg L⁻¹ (if the top 5 cm of sediment are resuspended).

In the diffusion cores, Fe concentrations (total + filterable) increase constantly during the course of the experiment (Fig. 5a). Concentrations increase from 0.8 to 4.1 mg L⁻¹ in core C6 (maintained at 21 °C ±2 °C) and from 0.5 to 2.0 mg L⁻¹ in core C3 (maintained at 9 °C ±1 °C). They are higher at the sediment-water interface (0-5 cm) (triangles in Figs 4 and 5) relative to the overlying water column (>5 cm) (rectangles in Figs 4 and 5). The overall increase in Fe during the incubation period is statistically significant (at the 95% confidence level) in both cores with slopes of 0.16 (R² = 0.85) in core C6 and 0.05 (R² = 0.68) in core C3. The total concentrations of Fe and Mn are higher in core C1, where a higher amount of sediment is suspended in the water column. The proportion of soluble-colloidal species, however, is lower even during reducing conditions where Fe²⁺ and Mn²⁺ are expected to predominate (Figs 4a and 4b).

Water column redox (pE) and pH of the resuspension cores are plotted in figure 4. The plot shows that circumneutral pH prevails in the water column of both cores throughout the experiments with median pH of 6.57 and 6.86 in C1 and C5, respectively. Redox potentials remain high (pE >4) in core C5 throughout the experiment. In core C1, the pE drops dramatically after the initial resuspension event (pE <1) but then rises after 12 days of incubation until initial pE conditions are re-established.

Core resuspension is associated with a significant increase in water colour, which changes from colourless to orange-brown during initial resuspension. The discolouration of the water persists throughout the duration of the experiments. Filtration of the samples using 0.45 μm Whatman filter disks removes colouration of the water completely together with more than 90% of the Fe and 24% to 60% of the Mn.

4.2. Core Diffusion

In the diffusion cores, Fe concentrations (total + filterable) increase constantly during the course of the experiment (Fig. 5a). Concentrations increase from 0.8 to 4.1 mg L⁻¹ in core C6 (maintained at 21 °C ±2 °C) and from 0.5 to 2.0 mg L⁻¹ in core C3 (maintained at 9 °C ±1 °C). They are higher at the sediment-water interface (0-5 cm) (triangles in Figs 4 and 5) relative to the overlying water column (>5 cm) (rectangles in Figs 4 and 5). The overall increase in Fe during the incubation period is statistically significant (at the 95% confidence level) in both cores with slopes of 0.16 (R² = 0.85) in core C6 and 0.05 (R² = 0.68) in core C3. Most Fe is present in particulate form, although colloidal-soluble Fe accounts for 14% to 36% (0.27-1.37 mg L⁻¹) of the total concentration in core C6 and 13% to 43% (0.16-0.64 mg L⁻¹) in core C3. Manganese is present almost exclusively (94-100%) in colloidal-soluble form. Concentrations initially increase from 2.2 mg L⁻¹ to 3.9 mg L⁻¹ in core C3 (Fig. 6), but remain constant after 15 days. In core C6, Mn concentrations increase from 1.3 mg L⁻¹ to 2.7 mg L⁻¹ during the first 12 days of incubation but then decrease to 2.2 mg L⁻¹ (Fig. 6). The increase is statistically significant (at the 95% confidence level) in core C3 (slope = 0.08, R² = 0.90) but not in core C6.

As for the resuspension cores, the observed core water concentrations are very high compared to concentrations in the reservoir. However, accounting for dilution of these concentrations by diffusion into an overlying water column of 3-5 m yields concentrations

Fig. 4. pE-pH diagram for (a) Fe and (b) Mn species in (resuspension) cores C1 (open triangles) and C5 (closed triangles) (The diagram is drawn for conditions in the water column of core C5 at the end of the experiment: Fe = 2.4×10⁻⁷ mol L⁻¹ and Mn = 1.2×10⁻⁴ mol L⁻¹ at 19 °C.) Complexation with dissolved organic matter is not considered in the diagram.
in the range of 100-160 μg Fe L⁻¹ and 70-110 μg Mn L⁻¹ (at 21 °C) as well as 40-70 μg Fe L⁻¹ and 80-130 μg Mn L⁻¹ (at 9 °C).

The pH and pE ranges measured in the core water column are plotted in figure 7. The pH is similar in both cores (pHmedian = 6.70 and 6.54, respectively) and shows little variation (<0.25 pH units) throughout the experiments. Redox potentials (pE) (measured in the upper water column) remain high throughout the experiment with pEmedian = 7.4 in core C3 and pEmedian = 6 in core C6.

### 4.3. Core metal distribution

Figure 8 illustrates the distribution of reactive Fe and Mn in the sediment cores analysed (a) prior to and (b, c) after the experiments. Concentrations of aluminium (Al) are included in the graphs to provide a "conservative" reference for evaluating the relative changes in Fe and Mn in the cores. There are clear differences in the amount of Fe and Mn present within the sediment of the different cores. Concentrations are generally higher in the pre-experimental cores compared to the diffusion cores. The largest change is observed for extractable (reactive) Mn which declines from mean concentrations of 142 mg L⁻¹ (A) and 144 mg L⁻¹ (C4) in the pre-experimental cores to 19 mg L⁻¹ (C3) and 15 mg L⁻¹ (C6) in the diffusion cores. Comparison of means (based on Student’s t-test) shows that this reduction is statistically significant (at the 95% confidence level). Similarly, mean concentrations of extractable (reactive) Fe in the diffusion cores C3 (184 mg L⁻¹) and C6 (133 mg L⁻¹) differ significantly (at the 95% confidence level) from the pre-experimental core C4 (234 mg L⁻¹) but not from core A (134 mg L⁻¹). In the resuspended cores C1 and C5, concentrations are similar to pre-experimental conditions, only core C1 shows a (statistically) significant reduction in Mn (mean 65 mg L⁻¹).

There are distinct differences in the vertical distribution of reactive Fe and Mn within the sediment cores. In the pre-experimental cores (Fig. 8a), Fe and Mn are enriched in the top 0-2 cm of the sediment. The Mn peak is better established than that of Fe and peaks at higher concentrations. Below 3 cm, concentrations of extractable Fe constantly increase with depth with a maximum at around 7 cm in core C4. Sediment profiles of the diffusion cores (Fig. 8b) show similar enrichment of weakly-bound Fe in the upper 3-4 cm of the core, in
particular in C6. Below this zone, concentrations gradually decline and concentrations in the deeper parts of the sediment column are low. Core C3, incubated at 9 °C, shows little Fe depletion at depth. However, the sediment of both cores is almost completely depleted of Mn with low concentrations (Mn < Al) at all depths throughout the profile. In the resuspension cores (Fig. 8c), the distribution of reactive Fe and Mn is similar to that of the pre-experimental cores in core C5 (top 1-2 cm resuspended) but resembles that of the diffusion cores in core C1 (top 4-5 cm resuspended).

4.4. Wind-driven sediment resuspension

The grain-sizes available for transport have \( D_{50} \) values that range from 7.95 µm (MS11) to 15.03 µm (MS6) (Tab. 2). Bottom shear stress estimates are calculated for the shallow part of the reservoir with water depths of <20 m using equations 1-6. The results for site MS11 (water depths 13.8 m) are presented in figure 9a for hourly wind speed data observed during the study period (September 1999-October 2001). The value of the critical entrainment threshold (\( \tau_c \)) at this site is

---

**Fig. 7.** pE-pH diagram for (a) Fe and (b) Mn species in (diffusion) cores C3 (closed circles) and C6 (open circles) (The diagram is drawn for conditions in the water column of core C6 at the end of the experiment: Fe = 2.5×10⁻² mol L⁻¹ and Mn = 6.5×10⁻⁵ mol L⁻¹ at 19.4 °C).

**Fig. 8.** Distribution of reactive Mn and Fe in the pre-experimental and experimental cores (plot shows concentrations in mg L⁻¹ (x-axis) versus depths below sediment-water interface in cm (y-axis)).
equal to 0.27 (based on equations 7 and 8) and this value is exceeded for all bottom shear stresses greater than 0.035. Bottom shear stresses of this magnitude can be generated by wind speeds of 13 m s\(^{-1}\) and above.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Depth (m)</th>
<th>(D_{50}) ((\mu)m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MS1</td>
<td>20.4</td>
<td>8.75</td>
</tr>
<tr>
<td>MS2</td>
<td>39.0</td>
<td>9.54</td>
</tr>
<tr>
<td>MS3</td>
<td>43.9</td>
<td>12.96</td>
</tr>
<tr>
<td>MS5</td>
<td>40.8</td>
<td>9.82</td>
</tr>
<tr>
<td>MS6</td>
<td>29.6</td>
<td>15.03</td>
</tr>
<tr>
<td>MS7</td>
<td>36.6</td>
<td>8.48</td>
</tr>
<tr>
<td>MS8</td>
<td>19.2</td>
<td>8.74</td>
</tr>
<tr>
<td>MS9</td>
<td>30.9</td>
<td>10.08</td>
</tr>
<tr>
<td>MS11</td>
<td>13.8</td>
<td>7.95</td>
</tr>
<tr>
<td>MS13</td>
<td>22.5</td>
<td>9.76</td>
</tr>
<tr>
<td>MS14</td>
<td>33.2</td>
<td>11.02</td>
</tr>
</tbody>
</table>

Such strong winds occur in the study region predominantly during the autumn and winter period. They are relatively infrequent (less than 2\% of the time) (Fig. 9b), but have the potential to disturb sediments in the shallow areas of the reservoir with water depths below 20 m. At full capacity, these areas make up about one third of the total reservoir area (Fig. 1).

5. DISCUSSION

Resuspension introduces high quantities of Fe and Mn into the core water columns. The five fold enrichment of (total) Fe relative to (total) Mn reflects the higher retention of Fe within the sediment (Abesser 2003). Almost all Fe in the resuspended water column is present in particulate form and Fe\(^{3+}\) appears to be the stable Fe species in the water column of the two cores during most of the experiment (Fig. 4a). Most Fe is probably present as non-reactive, refractory Fe oxides and/or incorporated in the crystal structure of clay minerals and silicates. Only one third is likely to be present as reactive oxyhydroxides (Tessier et al. 1979; Davison & De Vitre 1992), which can be reduced under low redox conditions and probably provide the main source of Fe within the sediments. Manganese is mostly present as dissolved Mn\(^{2+}\) which is the stable species at the prevailing pHe-pE conditions in the experimental columns (Fig. 4b). The high proportion of dissolved species reflects the low stability of particulate Mn in the aquatic environment where it is mostly present as reactive oxyhydroxides (Davison 1993). These are rapidly reduced in the suboxic water column and/or when encountering the sediment-water interface. As a result, Mn often accumulates near the sediment-water interface (Davison 1993) while the deeper sediment layers are depleted as seen in the profiles of the pre-experimental cores (Fig. 8). Extrapolating the observed core column concentrations to the conditions in the reservoir during the draw down period gives Fe concentrations in the range of 210-490 \(\mu\)g L\(^{-1}\). These values are comparable to concentrations measured in the reservoir (226-467 \(\mu\)g L\(^{-1}\)) and at the treatment works (350 \(\mu\)g L\(^{-1}\)) during maximum draw down, suggesting that resuspension was a major contributor to the observed water quality deterioration in the reservoir. The extrapolated concentrations of Mn are somewhat higher (89-97 \(\mu\)g L\(^{-1}\)) than those measured at the treatment works (50 \(\mu\)g L\(^{-1}\)). Since much of the Mn is present in dissolved form, it is likely that oxidation has removed some of the Mn from the water column during the 60 km passage of the water from the reservoir to the treatment works. Resuspension causes serious discolouration of the water which is probably caused by the presence of particulate matter (silt/clay particles, particulate organic matter, Fe particles) and not by dissolved organic substances which often are the cause for discoloration in upland streams and lakes (Freeman et al. 2001; Pace & Cole 2002). These particles have low settling velocities and also provide good sorption sites for dissolved metals. The sorption potential of these particles can increase during resuspension as increased inter-particle collisions and enhanced particle disaggregation create additional, new sorption sites on newly-formed particle surfaces (Brass-
Iron and manganese mobilisation

Sorption of Fe (and Mn) onto these new sites removes dissolved species from the water, as indicated in C1, but it also enhances the residence time of Fe (and Mn) in the water column as the sorbed species are more stable under the prevailing pH-pE conditions. Combined with low settling velocities, this implies that after resuspension events, prolonged time is required for the recovery of the system’s original water quality.

Wind-induced resuspension in the Megget Reservoir is estimated to occur at wind speeds of 13 m s⁻¹ and above and can affects sediments at water depth <20 m. At full capacity, these shallow areas constitute about one third of the total reservoir area (Fig. 1). Hence, water quality deteriorations resulting from such resuspension events may be substantial, particularly during autumn when the reservoir volume can be as low as 60% of the full reservoir capacity even in years of normal reservoir operation (i.e., no artificial draw down) (Abesser 2003). Wind speeds in excess of 13 m s⁻¹ occur predominantly during the autumn and winter period and, under current climatic conditions, are relatively rare (<2%). However, climate change scenarios predict a 5% change in wind speeds for Scotland as well as an increase in rainfall intensities during the winter month, particularly in eastern Scotland (Hulme et al. 2002; Barnett et al. 2006). This implies that natural, wind-induced resuspension events may become more frequent and associated water quality problems (e.g., colouration) may prevail throughout the autumn/winter period until conditions are sufficiently calm to allow particle settling. Resuspension events can also be expected to occur in other supply reservoirs and/or freshwater lochs in the study region, most of which are located in similar geological and geomorphological settings. Besides the adverse effects on water quality, such events are likely to increase nutrient availability and entrainment in the water column, thereby impacting on the lake's ecosystem productivity (Schallenberg & Burns 2004). The effect of increasing rainfall intensities on resuspension has not been investigated in this study. However, bottom currents generated by high discharges to the reservoir during intensive rainfall events can contribute to sediment resuspension and can even impact on deep lake/reservoir regions that are otherwise not affected by wind-induced stresses (Hawley & Lee 1999). In addition to sediment disturbance, such high-rainfall events also supply large quantities of dissolved Fe and Mn via inputs from the feeder streams with concentrations of up to 1200 μg L⁻¹ and 150 μg L⁻¹, respectively, during peak runoff (Abesser et al. 2006).

Under undisturbed conditions, mobilisation within and release from the sediment becomes an important source of dissolved Fe and Mn in the water column. Concentrations of Fe and Mn are almost an order of magnitude lower, compared to those resulting from resuspension, and are mostly controlled by chemical factors (i.e., temperature, pE, mineral solubility and saturation) rather than by physical processes. Sediment profiles suggest that Fe²⁺ and Mn²⁺ are mobilised within the sediment and released into the overlying water column. This release occurs even though oxic conditions prevail in the upper water column throughout the experiments (probably because oxygen enters the core water column during sample collection and core replenishment). However, soluble Fe²⁺ is not stable in the water column under these pE (and pH) conditions (Fig. 7a) and is rapidly oxidised to particulate Fe³⁺ probably in the form of amorphous ferric hydroxides. Assuming a half life of 4 hours in oxygenated environments (pH 7, 10 °C) (Davison & Seed 1983), Fe²⁺ can be expected to travel about 0.5 cm (if only molecular diffusion operates) in a plentiful supply of oxidants before being re-oxidised (Hamilton-Taylor & Davison 1995). This is consistent with observations in the core tubes where Fe precipitates form about 0.5-3 cm above the sediment on the inside of the core tubes. Dissolved Mn²⁺ is oxidised about 106 times slower than Fe²⁺ (at circumneutral pH) (Martin 2005) and is more stable in the core water columns of the diffusion cores (Fig. 7b). Its high mobility and the constant release of dissolved Mn²⁺ from the sediment has led to Mn depletion in the core sediments of C3 and C6 (Fig. 8) so that concentrations in the water column could no longer increase after 12-15 days (Fig. 3). Extrapolating from the concentrations in the diffusion cores to an overlying water column of 3-5 m gives concentrations similar to those measured in the reservoir (~3-5 m above the sediment) during summer stratification which averaged around 125 μg L⁻¹ for Fe and 105 μg L⁻¹ for Mn. This confirms that sediment release, as simulated in this study, occurs during summer stratification. However, it is unlikely to cause water quality problems as the release of Fe is low (and Fe is also rapidly oxidised) and Mn within the reservoir sediments is probably depleted during summer stratification, thereby limiting the amount of Mn that can accumulate in the bottom waters. In addition, some of the Mn may actually be removed from the water column as indicated by the decline in Mn concentrations in core C6. A major pathway for such Mn removal in natural systems is bacterial Mn oxidation (Tebo et al. 1997) which results in the formation of biogenic Mn oxides in the form of vernadite (Wehrli et al. 1995), birnessite (Tebo et al. 2004) or reduced Mn oxides, such as hausmannite or mangane. These processes commonly occur at elevated temperatures, i.e., between ~15 °C to ~30 °C (Tipping 1984) and could explain the formation of brown spots, interpreted as Mn precipitates, on the inside of core tube C6. The absence of such precipitates in the colder waters of core C3 (~9 °C) supports this assumption of biologically-mediated oxidation. The form in which the Mn oxide phase is present was not determined in this study, but should be investigated in future research efforts to establish the role of biota in Mn cycling within the reservoir.
An obvious limitation of this study is the small number of cores and replicate treatments that were employed during the experiments. This means that there is only limited information on the spatial variability in metal concentrations and distribution within the reservoir sediments and implies that some of the observed differences in metal release may be due to heterogeneity in the sediment distribution and chemistry between the cores. Furthermore, the results from this incubation experiment only provide a "snapshot" of the fluxes between the sediment and the water. They are specific to the part of the reservoir where the cores were collected and the environmental conditions (temperature, pH, redox, etc.) that prevailed in the core columns during incubation. Considering these limitations, sediment release rates were not calculated as part of this study as more spatially and temporally varied data are required to determine representative numbers.

6. CONCLUSIONS

Resuspension can introduce large quantities of Fe and Mn into the water column of the Megget Reservoir and, in this oligotrophic system, is far more effective than release from sediment diffusion. It is more important for the mobilisation of Fe which, in comparison to Mn, is present in large quantities within the sediments. Under the current climate, natural resuspension events at the Megget Reservoir are rare, but are expected to become more frequent as climatic conditions change towards increased wind speeds and higher rainfall intensities. Such an increase is likely to affect other reservoirs and lochs in the study region leading to increased sediment resuspension with adverse effects on water quality and ecosystem productivity.

In the absence of physical disturbance, Mn and Fe are delivered to the water column by the rapid reduction of oxyhydroxide particles at the sediment water interface and/or within the sediment. Reductive dissolution and sediment release are unlikely to cause water quality problems in the Megget Reservoir as the release of Fe is low and Mn within the sediments will be quickly depleted. However, accumulation of these elements in the bottom waters may increase with changing climatic condition, in particular when the external Mn and Fe inputs increase as a result of higher catchment loading associated with the predicted rise in rainstorm intensities.

Although extrapolation of experimental results to in situ conditions is often difficult and despite the limited number of cores, this laboratory study has successfully separated the internal processes related to resuspension and sediment diffusion and has helped to evaluate their potential impact on the water quality of the Megget Reservoir in the past and in the future.

ACKNOWLEDGEMENTS

The authors would like to thank Alexander Kirika (CEH Bush) for his help with collecting the sediment cores, Margaret and Colin Neal (CEH Wallingford) for providing access to and help with the ICP-OES analysis as well as for their continued support during the project and their great hospitality. Further thanks to Richard Bates (University of St Andrews) for providing the bathymetric data of the Megget Reservoir as well as to Heather Wickham (CEH Wallingford) for her help with running the ICP-OES. David Kinniburgh (BGS Wallingford) is thanked for the provision of the PhreePlot programme which was used for calculating and plotting pH-pE diagrams as well as for his invaluable comments on this manuscript. We further thank the two anonymous reviewers for their comments which greatly helped to improve this manuscript. The Environmental Change Network (ECN) is acknowledged for providing all wind speed data used in this study. Funding and facilities for this study have been provided by East of Scotland Water (now part of Scottish Water) and by the University of St Andrews.

REFERENCES


