Historical anthropogenic heavy metal input to the south-eastern North Sea

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Abstract
The Helgoland Mud Area (HMA) in the German Bight, covering an area of approximately 500 km², is one of a few depocentres for finer sediments in the North Sea. Radiocarbon and ²¹⁰Pb analyses revealed continuous sedimentation over the last several centuries. Zinc (Zn) and lead (Pb) contents in the sediments show a distinct increase towards the youngest most sediments with the thickness of the heavy metal enriched sediments ranging from 15 to 103 cm. Stratigraphic data indicate that the onset of heavy metal enrichment is diachronous progressing north-westward over the depocentre, paralleled by a decrease in the thickness of the enriched layer. Beginning already during medieval times, the enhanced input of Zn and Pb seemingly is related to silver and zinc mining in the Harz Mountains and the Erzgebirge, well-known mining areas since the Bronze Age. Both regions are directly connected to the HMA by the Elbe and Weser rivers. Zn and Pb enrichment began in the south-eastern HMA and progressed subsequently with an average of 10 m per year north-westward, most likely triggered by variations in river discharge and by the hydrodynamic setting. Quantitative assessments of the Zn and Pb content in the sediments suggest that since the onset of enhanced Zn and Pb deposition, the anthropic Zn and Pb input in the HMA amounts to ~ 12,000 t and ~ 4000 t, respectively.

Introduction

The southern North Sea with its mesotidal and high wave-energy levels is generally characterised by sediment redistribution as the dominating sedimentological process. Only a few depocentres for the deposition of finer sediments exist in the North Sea (Lohse et al. 1995). One is the Helgoland Mud Area (HMA) in the German Bight in the SE North Sea (Fig. 1). In its centre, up to 30 m of sediments of Holocene age have accumulated (von Haugwitz et al. 1988). Despite generally low water depths of ~ 30 m in the west and ~ 15 m in the east (Fig. 1), a spatially coherent sedimentation pattern (Irion et al. 1987) as well as the presence of undisturbed palaeo-records (e.g. Scheurle et al. 2005) demonstrate the long-term depositional nature of the HMA. This also holds true for the most recent times, when human activities contributed substantially to sediment redistribution in the German Bight (Rijnsdorp et al. 1998; OSPAR Commission 2000), as a high-resolution paleo-record from the HMA can be well correlated to the 120-year-long instrumental Helgoland Roads time series (Scheurle and Hebbeln 2003).

Fürstner and Reineck (1974) were the first to describe an increase in the content of heavy metals in the uppermost decimetres of the sediments of the HMA. Based on ²¹⁰Pb and ¹³⁷Cs dating, Dominik et al. (1978) estimated that the increased input of these heavy metals (including Zn and Pb) started in the second half of the nineteenth century, with a second marked increase between AD 1930 and 1950. Analysing a set of 21 sediment cores, Irion et al. (1987) showed the presence of a distinct spatial pattern within this depocentre, marked by increasing thicknesses of Zn-enriched layers from a few centimetres in the west to > 300 cm in the east. Without having any detailed stratigraphic data, Irion et al. (1987) assumed that the increased Zn input only began in post-war times due to the dumping of contaminated harbour and estuarine muds. However, their interpretation yielded sedimentation rates for the uppermost sediments of the HMA of
> 30 mm year\(^{-1}\), which are much higher than estimates of 2 mm year\(^{-1}\) based on sedimentological observations (Reineck 1963), 7.7 mm year\(^{-1}\) based on \(^{210}\)Pb and \(^{137}\)Cs dating (Dominik et al. 1978), and 1.6 to 3 mm year\(^{-1}\) based on radiocarbon dating (Hebbeln et al. 2003; Serna et al. 2010).

Based on the analysis of sediment core GeoB 4801-1 from the HMA, Hebbeln et al. (2003) showed an onset of the Zn increase as early as AD 1750 (corrected here to AD 1850, see below). This early onset was linked to silver and zinc mining activities in the Harz Mountains and the Erzgebirge. Both are well-known mining areas located along the tributaries of the Elbe and Weser rivers that both drain into the German Bight. Near these mining areas and along the rivers, silver and zinc mining left traces in the environment since medieval times (e.g. Bartels 1996; Matschullat et al. 1997; Ortlam 1989).

There is agreement that the heavy metal enrichment in the uppermost sediments of the HMA is based on human activities (Förstner and Reineck 1974; Dominik et al. 1978; Irion et al. 1987; Hebbeln et al. 2003). Such pollution can cause severe problems. A mobilisation of heavy metals, such as Zn and Pb, from the sediments in marine environments could endanger marine organisms and consequently negatively influence the marine food chain (Förstner and Müller 1973). Indeed, Rotstigen (2009) showed that both the absolute abundance and the diversity of benthic foraminifera in the HMA has decreased dramatically since the beginning ~ AD 1800, coinciding with an increase in abnormal foraminifera morphologies.

Although the available data draw a coherent picture of (south-)eastward increasing thicknesses of the Zn- and Pb-enriched layer in the HMA, the temporal development of anthropogenic induced Zn and Pb inputs is uncertain. As the onset of enhanced heavy metal input to the HMA in the mid-twentieth century (Irion et al. 1987) has been challenged by radiocarbon-based data documenting a much earlier onset (Hebbeln et al. 2003) and as the so far favoured interpretation of a temporally coherent onset over the entire HMA (Irion et al. 1987) would require drastic variations in sedimentation rates, the temporal development of enhanced Zn and Pb inputs to the HMA requires a critical evaluation. Therefore, based on
a set of five new dated sediment cores, this study aims to analyse the spatio-temporal development of anthropogenic Zn and Pb input to the HMA, to identify possible sources and to understand the transport and deposition of heavy metals (and sediments).

Regional setting

The sediment transport mechanisms in the southern North Sea are dominated by the bedload transport of mobile sands and by the suspended load of fine particles supplied by several rivers, however, with the latter being largely exported from the region due to relatively high hydrodynamic energy levels (e.g. Puls et al. 1997; Zeiler et al. 2000). The HMA is the only significant depocentre for fine sediments in the southern North Sea, except from some intertidal flats and some estuaries (Figge 1981). However, its evolution is still under discussion. The following theory for the evolution of the HMA is based on the studies of Hertweck (1983), von Haugwitz et al. (1988) and Mayer (1995). When sea level rose and erosive wave action resulted in the disconnection between Eiderstedt Peninsula and the island of Helgoland between 1500 years (von Haugwitz et al. 1988) and 2000–3000 years ago (Irion et al. 1987), the modern-day circulation in the SE North Sea with longshore currents from W to E and from S to N was established. This has been assumed as the main trigger for the onset of the deposition of suspended particulate matter (SPM) in the HMA (Reineck et al. 1967).

The continuous deposition of finer sediments in the HMA is attributed to a small-scale, anti-clockwise eddy that is driven by the interactions of longshore coastal currents, the discharge from the Elbe and Weser rivers, and tidal dynamics (Hertweck 1983). This eddy can be subdivided into five small-scale eddies, one of them being the main driver for the deposition of fine sediments in the HMA (Mayer 1995). A three-dimensional, numerical model for the transport of SPM revealed the existence of vertical eddies with the rising side situated above the HMA that appear to be consistent at nearly every wind direction (Mayer 1995). This eddy can be subdivided into five small-scale eddies, one of them being the main driver for the deposition of fine sediments in the HMA (Mayer 1995). A three-dimensional, numerical model for the transport of SPM revealed the existence of vertical eddies, one of them being the main driver for the deposition of fine sediments in the HMA (Mayer 1995). A three-dimensional, numerical model for the transport of SPM revealed the existence of vertical eddies, one of them being the main driver for the deposition of fine sediments in the HMA (Mayer 1995).

Methods

A set of six gravity cores with lengths between 6.87 m and 11.41 m was collected from the HMA during RV Meteor cruise M 40/0 in 1997 (Table 1) in close proximity to each other (Fig. 1). The sediment cores were split lengthwise into two halves. One half was analysed for the chemical composition of the sediments using an XRF Core Scanner III (AVAATECH Serial No. 12) at MARUM, University of Bremen. This instrument allowed for a non-destructive, semi-quantitative determination of major and minor elements (e.g. heavy metals). In this study, only Zn and Pb contents will be discussed. The other half was used to take discrete samples for the quantitative analysis of Zn and Pb contents and to determine the age of the sediment at selected depth levels.

For XRF scanning, the split core surface was covered with a 4-μm thin SPEXcerti Prep Ultralene 1 foil to avoid contamination of the XRF measurement unit and desiccation of the sediment. Net intensities of Zn and Pb were measured at 1-cm intervals for the upper two core sections for each core. Further downcore, the scanning interval was increased to 2 cm. Measurements were conducted over a 1-cm² area with a down-core split size of 10 mm using generator settings of (1) 30 kV, a current of 1.0 mA, and (2) 10 kV, a current of 0.3 mA, both with a sampling time of 20 s directly at the split core surface. Data were acquired using a Canberra Digital Spectrum Analyser DAS 1000 and an Oxford Instruments 100W Neptune X-ray tube with Rhodium (Rh) target material. The raw data spectra were processed by the analysis of X-ray spectra by applying the Iterative Least Square software (WIN AXIL) package from Canberra Eurusys. The resulting data were provided for every measured element in counts per 20 s and the Zn and Pb data being focused on here are provided as normalised net intensities, i.e. as percentages of the total counts for each individual measurement, to minimise errors due to variable water content, grain size variations and others (Tjallingii et al. 2007). For analysing marine sediment cores, XRF scanning has become increasingly popular in the past decades because it allows non-destructive extraction of near-continuous records of element intensities from sediment cores with a minimum of analytical effort with the results being seen as valuable indicators for true element concentrations (e.g. Weltje and Tjallingii 2008).

To calibrate the XRF scanner data for Zn and Pb, 66 discrete samples of 10 cm³, taken at positions corresponding to the core-scanner measurements, were taken, freeze dried and grounded using a mortar and pestle. Then 700 ± 0.6 mg of each sample was weighed into a ceramic crucible and mixed with 4200 ± 1 mg of a di-lithium tetraborate fusion flux (Spectromelt® A10; Merck) and about 1000 mg of ammonium nitrate. After 1 h of pre-annealing at 500 °C, the samples were fused and transformed into glass beads and measured for the Zn and Pb contents via wavelength dispersive XRF spectrometry (WD-XRF) at ICBM, University of Oldenburg. To determine the trueness and precision of the method, an international reference material (GSR_6) and in-house standards (Peru-1, Loess 1.6) were used. The results for the three
Radiocarbon (AMS14C) dating. Samples were wet-sieved over different depths were taken for accelerator mass spectrometer in a stratigraphic framework, discrete samples from the HMA. All AMS 14C ages were corrected for 4806-1. For core GeoB 4801-1, the age model of Scheurle performed on cores GeoB 4802-1, 4803-1, 4805-1 and 4806-1 was used. All AMS 14C ages were corrected for isotopic fractionation using δ13C values. Resulting conventional radiocarbon ages were converted into calendar years using the Calib 7.1 software (Stuiver et al. 2017) with the Marine 13 calibration curve (Reimer et al. 2013) considering the global mean reservoir age of 400 years (Bard 1998), which has been confirmed for nearby Danish marine waters (Heier-Nielsen et al. 1995). Results are given as the median probability calendar ages accompanied by the 1σ (68.3% probability) and 2σ ranges (95.4% probability).

To put the historic development of Zn and Pb contents in the HMA in a stratigraphic framework, discrete samples from different depths were taken for accelerator mass spectrometer radiocarbon (AMS14C) dating. Samples were wet-sieved over 500, 150 and 63 μm mesh size sieves and oven dried at 50 °C for 24 h for further analysis. For 17 samples, approximately 10 mg of mixed benthic foraminifera was picked from the fraction 500–150 μm and sent to the Beta Analytics Limited Radiocarbon Laboratory in London or to the Poznan Radiocarbon Laboratory. Radiocarbon measurements were performed on cores GeoB 4802-1, 4803-1, 4805-1 and 4806-1. For core GeoB 4801-1, the age model of Scheurle et al. (2005) was used. All AMS 14C ages were corrected for isotopic fractionation using δ13C values. Resulting conventional radiocarbon ages were converted into calendar years using the Calib 7.1 software (Stuiver et al. 2017) with the Marine 13 calibration curve (Reimer et al. 2013) considering the global mean reservoir age of 400 years (Bard 1998), which has been confirmed for nearby Danish marine waters (Heier-Nielsen et al. 1995). Results are given as the median probability calendar ages accompanied by the 1σ (68.3% probability) and 2σ ranges (95.4% probability).

Particle-size measurements of sediments of cores GeoB 4801-1 and 4806-1 were performed at the Senckenberg Institute, Department of Marine Science, Wilhelmshaven. Samples taken every 5 cm were desalted in semi-permeable hoses over 24 h. The desalted samples were washed through a 63 μm mesh size sieve to separate the mud fraction (< 63 μm) from the sand fraction (> 63 μm). In addition, at the MARUM, University of Bremen, 17 samples of core GeoB 4803-1 were washed through a 63 μm mesh size sieve to separate the mud fraction (< 63 μm) from the remaining sediment. The weight percent of the mud (< 63 μm) and the sand fractions (> 63 μm) were calculated by weighing the dry sediment before and after sieving.

Table 1 Site information for the sediment cores collected from the Helgoland Mud Area taken by the German RV Meteor during cruise M40/0 in 1997

| Core number | Latitude | Longitude | Water depth (m) | Core length (m) |
|-------------|----------|-----------|----------------|----------------|
| GeoB 4801-1 | 54°06.7’ N | 8°02.2’ E | 25 | 11.41 |
| GeoB 4802-1 | 54°05.7’ N | 8°04.4’ E | 25 | 8.01 |
| GeoB 4803-1 | 54°01.7’ N | 8°09.0’ E | 22 | 7.25 |
| GeoB 4804-1 | 54°07.8’ N | 8°10.8’ E | 20 | 6.87 |
| GeoB 4805-1 | 54°06.8’ N | 8°07.6’ E | 21 | 7.34 |
| GeoB 4806-1 | 54°03.1’ N | 8°00.6’ E | 29 | 7.40 |

Results

XRF scanner data

The relative Zn and Pb contents reveal similar background values in the lower parts of all cores, with Zn and Pb values ranging between 0.7% and 1.9% and between 0.3% and 1.1%, respectively (Fig. 2). Towards the top of the cores, a significant increase of the relative Zn and Pb contents is observed reaching Zn and Pb values ranging between 1.2% and 2.9% and between 0.4% and 1.6%, respectively (Fig. 2), except for GeoB 4804-1. The uppermost 8 cm of core GeoB 4804-1 was crushed and squeezed during retrieval, and, therefore, could not be analysed via XRF Core Scanner. These uppermost sediments enriched in Zn and Pb differed in both contents and thickness. Core GeoB 4803-1 shows both the thickest enriched sediment layer (103 cm) as well as very high relative Zn (mostly > 2%) and Pb (mostly > 1%) contents, with these high contents largely mirrored in core GeoB 4802-1, however, only in its upper 43 cm. Both cores are characterised by a decreasing trend in the Zn and Pb relative contents in the uppermost sediments. In cores GeoB 4801-1, 4805-1 and 4806-1, the pattern of the Zn and Pb relative contents is very similar with the significant increase occurring at similar core depths of 15–25 cm, with maximum relative Zn and Pb contents ranging between 1.9% and 2.3% and between 1.3% and 1.4%, respectively.

Quantitative XRF data

The quantitative XRF measurements of the 66 samples used for the calibration of the semi-quantitative XRF core scanner data yielded Zn and Pb contents up to 300 and 100 ppm, respectively, whereas the background contents for Zn and Pb are ~ 45 and ~ 20 ppm, respectively (Fig. 3).

Calibration of Zn and Pb scanner data

Cross-plots between the quantitatively measured Zn and Pb contents and the corresponding relative contents based on the semi-quantitative scanner data show coefficients of determination (R²) of 0.65 for Zn and 0.72 for Pb (Fig. 3). For the Zn data, the cross-plot reveals one outlier (Fig. 3a) that, however,
does not strongly influence the calibration. Thus, the equations used to calibrate the Zn and Pb XRF scanner data (see Fig. 3) include all 66 quantitatively measured samples.

**Calculated quantitative Zn and Pb contents**

For the lowermost core sections, the calibrated XRF Core Scanner data reveal mean absolute background contents of ~41–46 ppm for Zn in the individual cores (Table 2, Fig. 4). After the increase in the upper core sections, the maximum zinc contents vary between ~100 ppm (GeoB 4805-1) and ~200 ppm (GeoB 4802-1 and GeoB 4803-1; Table 2, Fig. 4). The maximum contents of ~200 ppm in cores GeoB 4802-1 and GeoB 4803-1 correspond to almost five times the natural background. However, on average, the zinc contents of the upper polluted sediment layers are between 2.0 and 2.9 times higher than the natural background content (Table 2).

The Pb contents show a pattern very similar to the Zn contents, however, with lower values. The average Pb background contents in the lower parts of the cores range between 17 and 20 ppm (Fig. 4, Table 2), whereas average contents in the enriched layers are 29 to 55 ppm (Table 2). Thus, the Pb contents within the enriched layers are on average 1.5 to 2.7 times higher than the natural background contents (Table 2).

The Zn/Pb ratios in the background sediments range between 2.1 (GeoB 4805-1) and 2.5 (GeoB 4801-1). In the upper sediment layers enriched in Zn and Pb, this ratio changes in each core to higher values ranging from 2.3 (GeoB 4805-1) to 3.4 (GeoB 4801-1), thereby demonstrating a relatively larger increase in the Zn content compared to the Pb content.
Particle size results

The particle size distributions in cores GeoB 4801-1 and 4806-1 indicate similar patterns, with the sediment being dominated by mud (< 63 μm) with average contents of 87 wt.% and 76 wt.%, respectively (Fig. 5). In contrast, the sediments of core GeoB 4803-1 are generally coarser and show larger variations, with an average mud content of 50 wt.% (Fig. 5).

Radiocarbon ages

The AMS-14C radiocarbon dates based on mixed benthic foraminifera reveal ages going back as far as 253 BC (Table 3). In

![Calibrated absolute zinc (Zn; dark/light shading, cut-off at 60 ppm) and lead (Pb, black line) contents (ppm) of the sediment cores from the Helgoland Mud Area. Black stars mark the available 14C datings with the respective ages given in BC/AD notation. The thick grey line linked to core GeoB 4801-1 marks that part of the core for which the stratigraphy is based on correlation to historical (Scheurle et al. 2005) and instrumental (Scheurle et al. 2003) data.](image-url)
some of the cores, age reversals, indicating possible effects of some re-sedimentation and/or bioturbation, can be observed (Fig. 6). Nevertheless, when orienting along the youngest dates (see discussion) or along overlapping sigma ranges for all four investigated cores, average sedimentation rates result in rather consistent values (Fig. 6). For the past centuries, these range between 0.5 and 1.2 mm year$^{-1}$, close to the value of 1.4 mm year$^{-1}$ since AD 1600 for core GeoB 4801-1 taken from a detailed age model (Scheurle et al. 2005). Highest rates of > 1 mm year$^{-1}$ are being restricted to the central and north-western HMA.

Discussion

The spatial pattern of enhanced Zn and Pb contents in the Helgoland Mud Area

In a first spatial assessment, Irion et al. (1987) reported enhanced Zn contents, measured from the fraction < 2 μm, in the uppermost sediments of 21 sediment cores distributed all over the HMA (Fig. 7) and even beyond its limits as defined by von Haugwitz et al. (1988). The trend towards thicker Zn-enriched deposits in the eastern HMA reported by Irion et al. (1987) is also reflected in the new data presented here, based on the Zn content in the total sediment (Fig. 7). Of course, the use of sediments comprising different grain sizes in both studies can be problematical when comparing such different data sets directly, especially in terms of quantification (see, e.g., Birch 2017). Consequently, both data sets are not compared quantitatively here. In contrast, the spatially more extensive data (Irion et al. 1987) are only compared to the new data in a qualitative way to demonstrate the coherent spatial pattern of

Table 3 Overview of the AMS$^{14}$C dates in the sediments based on the analysis of mixed benthic foraminifera

| Core number GeoB | Depth (cm) | Laboratory identification | Conventional radiocarbon age (B.P.) | $1\sigma$ range of calendar age | $2\sigma$ range of calendar age | Median probability calendar age |
|------------------|-----------|---------------------------|-----------------------------------|-------------------------------|-------------------------------|-------------------------------|
| 4802-1           | 9         | Poz-65886                 | Modern                            | AD 1518 to 1626               | AD 1480 to 1660               | AD 1568                       |
| 4802-1           | 37        | Beta-361007               | 690 ± 30                          | AD 1544 to 1636               | AD 1498 to 1676               | AD 1591                       |
| 4802-1           | 45        | Beta-361008               | 660 ± 30                          | AD 1645 to 1728               | AD 1568 to 1827               | AD 1694                       |
| 4802-1           | 51        | Poz-65913                 | 560 ± 30                          | AD 1314 to 1393               | AD 1286 to 1427               | AD 1353                       |
| 4802-1           | 49        | Beta-361009               | 970 ± 30                          | AD 14 to 165                  | BC 324 to BC 195              | BC 253                        |
| 4803-1           | 93        | Poz-66014                 | 2200 ± 50                         | AD 702 to 805                 | AD 676 to 879                 | AD 761                        |
| 4803-1           | 102       | Beta-361010               | 1570 ± 30                         | AD 1724 to 1744; 1751 to 1790; 1800 to 1892; 1944 to 1950 | AD 1712 to 1950 | AD 1824 |
| 4805-1           | 4         | Poz-65914                 | 55 ± 30                           | Modern                        | Modern                        | Post 1950                     |
| 4805-1           | 15        | Beta-361011               | 450 ± 30                          | AD 1650 to 1762               | AD 1566 to 1847               | AD 1702                       |
| 4805-1           | 21        | Poz-65916                 | 555 ± 35                          | AD 1491 to 1594               | AD 1467 to 1649               | AD 1549                       |
| 4806-1           | 5.5       | Poz-75172                 | 315 ± 30                          | Modern                        | Modern                        | Post 1950                     |
| 4806-1           | 12        | Beta-361012               | 650 ± 40                          | AD 1540 to 1670               | AD 1500 to 1690               | AD 1640                       |
| 4806-1           | 25        | Poz-65918                 | 660 ± 40                          | AD 1539 to 1649               | AD 1489 to 1683               | AD 1590                       |
heavy metal contamination in the HMA. This spatial pattern of gradually increasing thicknesses of the Zn-enriched layers from NW to SE is a clear indicator for continuous, essentially undisturbed sedimentation in the HMA as already pointed out by Irion et al. (1987). The pattern provided by the Zn contents in the GeoB cores studied here is almost completely mirrored in the Pb contents (Fig. 4).

**Sedimentation rates in the Helgoland Mud Area**

It has to be admitted here that the radiocarbon data are partly ambiguous as they are marked by several age reversals, which, however, can be expected in such a dynamic, relatively shallow water setting. When in a sediment core older sediments/fossils/dates occur on top of younger ones, it is obvious that these are reworked. Otherwise, it cannot be explained how the younger material got deeper into the core. Of course, arguing for reworking at one point also allows questioning the ‘younger ages’, which also might be reworked. However, to put it to the other extreme, if reworking of everything is assumed, it would be difficult to explain (1) the consistent regional pattern of zinc enrichment in the HMA developing along smooth gradients and (2) the consistent pattern of sedimentation rates resulting from orienting on the youngest ages. Finally, the reliability of the 14C dates obtained from core GeoB 4801-1, supported by the comparison to historical and instrumental data (Scheurle and Hebbeln 2003; Scheurle et al. 2005), gives some credibility to the approach followed here, considering the resulting 14C ages as reliable in the first place, unless reworking becomes obvious by age reversals. Of course, we cannot exclude that the dated material revealing ‘youngest’ ages has been reworked, but this is a problem inherent to all non-laminated sediments.

At first glance, the observation of a coherent sediment layer enriched in heavy metals that covers almost the entire HMA may point to an isochronous onset of enhanced heavy metal
deposition as also proposed by Irion et al. (1987). Obviously, such an interpretation would require a significant increase in sedimentation rates from west (~ 20-cm-thick enriched layer) to east (> 100-cm-thick enriched layer; Fig. 7). However, this is in contrast to the newly obtained radiocarbon dates (Table 3), which reveal rather similar sedimentation rates at

Fig. 7 Zinc (Zn) distribution map (modified after Irion et al. 1987) showing the total Zn contents (ppm) in the upper Zn-enriched sediments of the six sediment cores from the Helgoland Mud Area (HMA) investigated here (marked in red) and the Zn content above the natural background in the particle size fraction < 2 \( \mu \)m of the sediment cores studied by Irion et al. (1987) (marked in black). In core GeoB 4804-1 from the north-eastern HMA, where the enriched layers are thinnest, no enriched layer could be found due to lacking data from the uppermost 8 cm of that core that probably contained the contaminant signal.

Fig. 8 Precise dating of the onset of enhanced zinc deposition at site GeoB 4801-1 in the Helgoland Mud Area with the stratigraphy and the resulting sedimentation rates for this core being based on correlation to instrumental (Scheurle et al. 2003) and historical (Scheurle et al. 2005) records.
all sites, actually with a slight increase to the northwest, i.e. opposite to the expected direction (Fig. 6). In the uppermost 50 cm of the sediment column, all cores show decreasing ages towards the top with a rather narrow range of average sedimentation rates of 0.5 to 1.2 mm year\(^{-1}\) close to the sedimentation rate of 1.4 mm year\(^{-1}\) obtained for the well-dated core GeoB 4801-1 (Scheurle et al. 2005; Fig. 6). Further downcore, sedimentation rates in cores GeoB 4801-1 and 4803-1 remain rather stable (Fig. 6a) and follow roughly the same trend as above. Actually, the spatial pattern revealed by the sedimentation rates mirrors the thickness of the HMA (cf. Fig. 1; Von Haugwitz et al. 1988 with highest sedimentation rates of >1 mm year\(^{-1}\) in cores GeoB 4801-1 and 4802-1 fitting HMA thicknesses of >18 m and vice versa for cores GeoB 4803-1, 4805-1 and 4806-1.

The observed difference in sedimentation rate between the south-eastern core GeoB 4803-1 (0.8 mm year\(^{-1}\)) and the two north-western HMA cores GeoB 4801-1 and 4802-1 (~1.3 mm year\(^{-1}\)) is also reflected in the grain-size distribution with lower amounts of mud in the slower accumulating setting. This would be in line with the assumption that at the slightly shallower site GeoB 4803-1 a higher hydrodynamic energy reduces the deposition of finer material and, thus, the overall sedimentation.

Considering temporal variability in sedimentation rates not resolved by the new data presented here, but which are obvious for site GeoB 4801-1 at which the most recent sediments got deposited at a rate of 2.6 mm year\(^{-1}\) (Scheurle et al. 2005; Fig. 8), these sedimentation rates are in agreement with radiocarbon-based sedimentation rate estimates of 3 mm year\(^{-1}\) for the central HMA obtained by Serna et al. (2010). Deviating from these results are sedimentation rates of 7.7 mm year\(^{-1}\) reported by Dominik et al. (1978). However, this sediment core from an unreported position somewhere south of Helgoland is marked by event-driven sedimentation (storm flood deposits) and erosion (Dominik et al. 1978) and, thus, it is probably not well comparable to the HMA records described here.

For core GeoB 4801-1 the onset of enhanced heavy metal deposition can be rather precisely dated to ~AD 1850 (Fig. 8). Based on new XRF scanning results presented here that move the increase in Zn counts approximately 12 cm further up in the core, the previously published earlier age for this onset (~AD 1750 in Hebbeln et al. 2003) has been identified as an artefact. Assuming an isochronous onset over the entire HMA at ~AD 1850 would stretch the range of sedimentation rates from 1.4 mm year\(^{-1}\) (e.g. core GeoB 4801-1) to >20 mm year\(^{-1}\) (for a sediment core with >300 cm of polluted sediments; Irion et al. 1987), which would account for a 15-fold increase in sedimentation rate from NW to SE over only ~10 km. As such high rates in the order of centimetres per year have been tentatively linked to dredging activities in the Elbe estuary during the past few decades, sedimentation rates must have increased tremendously compared to the natural setting, when only little amounts of river borne sediments are expected to have reached the German Bight (Irion et al. 1987). Furthermore, a dramatic increase in sedimentation rates triggered by dredging would not align with an onset of enhanced sedimentation already as early as AD 1850. The high-resolution stratigraphy of core GeoB 4801-1 reveals (1) no major increase in sedimentation rate paralleling the onset of enhanced heavy metal input and (2) only slightly varying sedimentation rates since AD 1300 not exceeding 3.2 mm year\(^{-1}\) (Fig. 8). Following the idea of Irion et al. (1987) that in the mid-twentieth century enhanced dredging activities in the estuaries might have caused enhanced sedimentation in the HMA, the minor increase in sedimentation rate in core GeoB 4801-1 around AD 1940 might relate to this; however, this occurred almost a century after the input of Zn and Pb increased (Fig. 8).

Based on these considerations, we conclude that the sedimentation rate pattern based on the available radiocarbon dates (Fig. 6) provides the best representation of the history of sediment deposition in the HMA. Consequently, in the following we argue for a diachronous onset of enhanced heavy metal input.

The temporal pattern of increasing Zn and Pb inputs to the Helgoland Mud Area

Based on the radiocarbon dates, this diachronous onset can be followed along the various core sites. The first enrichments of Zn and Pb occurred in core GeoB 4803-1 in the southeast as early as ~AD 760 (Fig. 4). A very early age for this onset is supported by two accompanying \(^{14}\)C-ages in this core, also providing rather old, though reworked ages (Fig. 6a). The close-by core no. 14 (Fig. 7; Irion et al. 1987) provided a pollen-based ‘pre-medieval’ (i.e. ~AD 500) age estimate for a sediment sample taken in 2 m core depth from directly below the polluted sediments (Irion et al. 1987; von Haugwitz et al. 1988). This rough age estimate would result in a sedimentation rate slightly above 1 mm/year, thus in a similar order to the sedimentation rate estimates provided here. From the south-easternmost part of the HMA, the enhanced input of Zn and Pb progressed north-westward to reach site GeoB 4806-1 in the southernmost and site GeoB 4802-1 in the central HMA by ~AD 1650–1750. Advancing further to the north-west, sites GeoB 4805-1 and GeoB 4801-1 were reached at ~AD 1750–1850, respectively. Thus, the enhanced input of heavy metals to the HMA follows a time-transgressive pattern advancing from site GeoB 4803-1 towards the other sites with an average velocity of ~10 m year\(^{-1}\) (range for the four core sites, 9.5–10.6 m year\(^{-1}\)).
Sources of the enhanced Zn and Pb input

The constant low background values in the Zn and Pb contents as well as their significant increase in historical times (Figs. 2 and 4) indicate that the enhanced heavy metal inputs are due to anthropogenic activities. This is further corroborated by the Zn/Pb ratios, which show a distinct increase from 2.25 in the background sediments to 2.72 in the polluted sediments. This shift indicates a change in the source area, with the source(s) for the enhanced Zn and Pb inputs most likely differing from those contributing to the older sediments.

Earlier studies attributed enhanced Zn and Pb contamination to a variety of anthropogenic sources, including fossil fuel burning and sewage effluent since the 1850s (Dominik et al. 1978), dumping of acidic iron waste from titanium dioxide production (Dominik et al. 1978) and dredging activities in the neighbouring estuaries (Irion et al. 1987) both since the 1950s. From these possible sources, only the dredging activities could have resulted in the necessary increase in sedimentation required to form a >300-m-thick polluted sediment package within decades. Seeing the well-documented onset of Zn and Pb pollution at AD 1850 in core GeoB 4801-1, this would require—at least in the southeastern HMA where the polluted layer is thickest—two different sedimentation processes to explain first slow aggrading polluted sediments from ~AD 1850 to ~AD 1950, and much higher sedimentation afterwards. However, the comparatively coarser sediments found at site GeoB 4803-1 (Fig. 5) characterise the older unpolluted as well as the younger polluted sediments and do not provide any hint to a dramatic change in the sedimentation process. Actually, as pointed out above, the continuous high sand content supports the interpretation of having comparably low sedimentation rates at this ‘shallower’ site, where higher energies hamper the sedimentation of finer material. Thus, the sources proposed by Dominik et al. (1978) and Irion et al. (1987) are inconsistent with the new data provided here, which clearly demonstrate a much earlier onset of heavy metal enrichment (Fig. 4).

However, the new data do support the interpretation by Hebbeln et al. (2003), which related the anthropogenic signal to early mining activities in the Harz Mountains and in the Erzgebirge, both of which are well-known, long-established mining areas along the tributaries of the Elbe and Weser Rivers. Although the new data indicate a much earlier onset (~AD 760) than estimated before (Hebbeln et al. 2003), evidence for mining activities, especially in the Harz Mountains, extends back to ~3500 years BP (Monna et al. 2000). Furthermore, findings of enhanced Zn and Pb contents in sediments from the western foothills of the Harz Mountains with an estimated Karolingian age (AD 825 to AD 890) (Matschullat et al. 1997) and in Weser river sediments since AD 800 (Ortlam 1989) clearly document a fluvial transport of heavy metals from the mining locations towards the German Bight. In particular, a fluvial link to the Harz Mountains is indicated by increasing heavy metal contents in Weser sediments downstream of the confluence with the Aller River, a Weser tributary draining parts of the Harz Mountains (Pache et al. 2008). Earlier mining activities in the Harz Mountains, dating back to 3500 years BP (Monna et al. 2000), may have been conducted at a smaller scale than more recent mining, without contributing a detectable level of Zn and Pb to the HMA. Thus, the timing of the onset of enhanced Zn and Pb deposition in the HMA at ~AD 760 probably is a function of mining intensity/methodology and on the velocity of the signal propagation from the source areas through the rivers to the southern North Sea. Remarkable is the rather fast increase of Zn and Pb input at site GeoB 4803-1 at a very early time soon after its initiation at ~AD 760 (Fig. 4). As independent of the timing one usually would expect a gradual increase in pollution, the sudden increase observed here most likely is the result of changes in the sedimentological setting, possibly related to the positioning and strength of the mesoscale eddy controlling the sedimentation in the HMA (Mayer 1995).

Interestingly, on the nearby island of Helgoland, copper (Cu) was mined and smelted, probably from the early Bronze Age to medieval times (Schulz 1981). However, average Zn/Pb ratios of 0.27 in copper slices found near Helgoland (Schulz 1981) are significantly different from those found in our cores (2.25 to 2.72), which show no sign of enhanced Cu deposition. As heavy metal-enriched Weser sediments reveal Zn/Pb ratios of >2 (Ortlam 1989; Monna et al. 2000) and rather low Cu enrichments (Pache et al. 2008), heavy metals in the HMA sediments are most likely not related to copper mining on Helgoland, but are derived from the mainland rivers.

Actually, deposition of large-scale lead pollution has occurred for at least 3000 years in Europe with metal production and smelting as the main sources until the twentieth century (Renberg et al. 2001). The first major anthropogenic increase occurred during Roman times, when Pb contents in a Spanish peat bog revealed a fivefold increase above natural background values (Martinez Cortizas et al. 1997). In Swedish lakes, isotopic traces of local mining occur as early as AD 960 (Bindler et al. 2009) followed by significant increases in Pb contents in Baltic Sea sediments at AD 1200 (Zillen et al. 2012) and in lake sediments by AD 1500 (Bindler et al. 2009). Therefore, early shedding of heavy metal pollution around AD 800 originating from the Harz Mountains as recorded in Harz creek sediments (Matschullat et al. 1997), in Weser river sediments (Ortlam 1989) and in the HMA (this study) fits well into a pan-European picture.

The time-transgressive pattern of enhanced Zn and Pb input

The time-transgressive SE to NW advance of enhanced Zn and Pb inputs with on average 10 m year⁻¹ across the HMA might be due to an increasing contribution of river-borne...
material to the sedimentation in the HMA. For the period \(\sim \text{AD} 1250\) to \(\sim \text{AD} 1650\), overlapping with the NW-ward advance of the higher heavy metal input, decreasing salinities over the HMA have been explained with increasing river discharge to the German Bight (Scheurle et al. 2005). As only little riverborne material is expected to leave the estuaries (Irión et al. 1987), a further seaward advection of riverine waters stimulated by a higher discharge might have been able to change the composition of the HMA sediments qualitatively (higher heavy metal content) without affecting them quantitatively (as reflected in core GeoB 4801 sedimentation rate record: Fig. 8). Such a relative increase of the heavy metal load with increasing transport routes also have been described for the recent German Bight setting (Hinrichs et al. 2002). In line with the decreasing salinities corresponding to an enhanced presence of riverine waters, this impact on the composition of the HMA sediments probably has advanced progressively NWward. However, other scenarios invoking changes in the palaeogeography and/or sea level also might be generally feasible, but appear less likely on the time scales considered here.

In this context, one might wonder why the 1970s peak in atmospheric lead pollution resulting from gasoline burning has not shown a strong expression in the HMA sediments, although its fast spreading makes it already detectable in the ocean waters of the entire North Atlantic Ocean and even in corals from Bermuda (Boyle et al. 2014). The lack of such a signal in the HMA most likely is caused by the relatively minor contribution of atmospheric Pb to the deposition there. Ottley and Harrison (1991) provided an estimate for the atmospheric Pb deposition in the German Bight of \(\sim 600 \mu \text{g m}^{-2} \text{month}^{-1}\) based on data from 1980, which would amount to an annual deposition of \(7.2 \text{ mg m}^{-2} \text{ year}^{-1}\). Taking 30 ppm as the average Pb content in the polluted HMA sediments in core GeoB 4801-1 (Table 2) and its average sedimentation rate of \(1.4 \text{ mm year}^{-1}\) (considering a dry bulk density of \(0.9 \text{ g cm}^{-3}\)), Hebbeln, unpublished data), the sedimentary Pb deposition is \(38 \text{ mg m}^{-2} \text{ year}^{-1}\) (using the \(^{210}\text{Pb}\)-dating derived sedimentation rate of 2.6 mm year\(^{-1}\) for the uppermost sediments, this number would increase to \(70 \text{ mg m}^{-2} \text{ year}^{-1}\)). Thus, in coastal depocentres such as the HMA the atmospheric Pb input plays a subordinate role, which is in agreement with results from the Gulf of Gdansk in the Baltic Sea for example, where the riverine contribution also clearly dominates the atmospheric input (Zaborska 2014).

As the heavy metal contents in such depocentres strongly depend on the local setting, absolute values for (sub-)recent polluted sediments can be quite variable, ranging for Pb from low values of \(30-50 \text{ ppm}\) in the HMA and off the Polish coast (Zaborska et al. 2017) to values of \(>100 \text{ ppm}\) in Baltic Sea sediments (Bindler et al. 2009; Zillen et al. 2012). Put into a sedimentation/accumulation rate context, even strong local variability becomes obvious as, e.g., for a set of six rather young (post-AD 1850) sediment cores collected from the Gulf of Gdansk, Baltic Sea. These show rather consistent average Pb contents ranging from 35 to 67 ppm, which, however, show a large spread of \(9-106 \text{ mg m}^{-2} \text{ year}^{-1}\) in the average Pb accumulation rates (Zaborska 2014), with both ranges covering the values found in the HMA.

**Total anthropogenic Zn and Pb input to the mud area**

Using average values for the thickness of the uppermost sediment layer characterised by elevated Zn and Pb contents, this layer’s average Zn and Pb contents above the natural background and the sediment’s dry bulk density of \(\sim 0.86 \text{ g cm}^{-3}\) (see Table 2), the total anthropogenic Zn and Pb input to the whole HMA was estimated. Applying these values to the overall extent of the HMA of \(500 \text{ km}^{2}\), a total volume of \(245,000,000 \text{ m}^{3}\) or \(0.245 \text{ km}^{3}\) of enriched sediments can be assumed. This volume in combination with the average Zn and Pb contents above the natural background of 59 ppm for Zn and of 19 ppm for Pb and the average bulk density of \(0.86 \text{ cm}^{3}\) adds up to a rough estimate of anthropogenic induced inputs of \(\sim 12,000 \text{ t of Zn and } \sim 4000 \text{ t of Pb to the HMA. Although the density obtained from only one core might vary within the HMA, such variability would not affect the order of magnitude of the estimated anthropogenic Zn and Pb inputs.}

At first glance, these numbers seem to be rather large. However, the loads of the suspended heavy metals in the Elbe showed peak values of \(4400 \text{ t of Zn and } 350 \text{ t of Pb per year at the station Magdeburg in } 1986\) (FGG Elbe 2010) putting the numbers from the HMA into a broader perspective. Furthermore, in only 1 year, in 2014, a total of \(4.5 \text{ Mm}^{3}\) of sediments containing \(\sim 180 \text{ t of Pb and } \sim 1600 \text{ t of Zn was dredged from the port of Hamburg. At the Tonne E3 dumping site located } \sim 15 \text{ km south of Helgoland, and close to the HMA, } \sim 440,000 \text{ t of this dredged material was dumped containing } \sim 15 \text{ t of Pb and } \sim 104 \text{ t of Zn (Hamburg Port Authority 2015). Thus, the heavy metals deposited in the HMA obviously only reflect a fraction of the heavy metals that are present in the marine system of the German Bight. Most likely, other depocentres like the estuaries of the Elbe and Weser rivers, and the Skagerrak, which is the most prominent fine sediment depocentre in the North Sea, also serve as sinks for anthropogenic heavy metal inputs to the German Bight (Hinrichs et al. 2002).}

Interestingly, the high amounts of Zn and Pb in the dredged material dumped close to the HMA in recent years/decades seem to not have caused any increase in contamination within the cores analysed here. Instead, most of our cores show a decline in the Pb and Zn contents in the youngest sediments. Therefore, it seems that these dumped sediments somehow do not enter the eddy-controlled depositional system of the HMA, maybe due to the particle dynamics linked to the processing that might release only a fraction of the dumped material into a sustained suspension. Alternatively, the
declining Zn and Pb contents in the youngest HMA sediments may reflect reduced river sourced SPM loads due to the construction of various dams along the Weser and Elbe rivers.

**Conclusion**

In contrast to earlier interpretations linking enhanced heavy metal deposition in the HMA to increasing inputs of estuarine sediments remobilised by dredging activities throughout the last few decades, new data reveal that Zn and Pb enrichment in the HMA sediments follows a regional, time-transgressive pattern beginning already as early as AD 760. At this time, enhanced Zn and Pb inputs first reached the south-eastern HMA, probably as a consequence of increased mining activities in the Harz Mountains and the Erzgebirge within the catchments of the Weser and Elbe rivers that drain into the German Bight close to the HMA. Increasing river discharges since at least AD 1250 probably have triggered the gradual advance of the heavy metal across the HMA from SE to NW reaching its north-western part by AD 1850. However, as river-borne sediments are expected to only contribute little to the overall sedimentation in the German Bight, they rather impact the HMA sediments in a qualitative way (enhanced heavy metal contribution) than in a quantitative way (enhanced sedimentation). Indeed, measured sedimentation rates are rather constantly in the order of millimetres per year, which is in contrast to a previously assumed sudden increase in sedimentation rates to the order of centimetres per year in association with enhanced Zn and Pb inputs. Irrespective of the timing, the entire anthropogenic Zn and Pb inputs to the HMA are estimated to 12,000 t of Zn and 4000 t of Pb.

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