Identification of the natural background of phosphorus in the Scheldt river using tidal marsh sediment cores
Florian Lauryssen 1, Philippe Crombé 2, Tom Maris 3, Elliot Van Maldegem 2, Marijn Van de Broek 4, Stijn Temmerman 3, Erik Smolders 1

1Division of Soil and Water Management, Department of Earth and Environmental Sciences, KU Leuven, Kasteelpark Arenberg 20 bus 2459, 3001 Leuven, Belgium
2Department of Archaeology, Ghent University, Sint-Pietersnieuwstraat 35, 9000, Ghent, Belgium
3University of Antwerp, Ecosystem Management Research Group, Campus Drie Eiken, D.C.120, Universiteitsplein 1, 2610 Wilrijk, Belgium
4Sustainable Agroecosystems group, Department of Environmental Systems Science, Swiss Federal Institute of Technology, ETH Zürich, Zürich, Switzerland

Correspondence to: Florian Lauryssen (florian.lauryssen@kuleuven.be)

Abstract. Elevated phosphate (PO₄) concentrations can harm the ecological status in water by eutrophication. In the majority of surface waters in lowland regions such as Flanders (Belgium), the local PO₄ levels exceed the limits defined by environmental policy and fail to decrease, despite decreasing total phosphorus (P) emissions. In order to underpin the definition of current limits, this study was set up to identify the pre-industrial background PO₄ concentration in surface water of the Scheldt river, a tidal river in Flanders. We used the sedimentary records preserved in tidal marsh sediment cores as an archive for reconstructing historical changes in surface water PO₄. For sediment samples at different depths below the sediment surface, we dated the time of sediment deposition and analysed the extractable sediment-P. The resulting time series of sediment-P was linked to time series of measured surface water PO₄ concentrations (data 1967-present). By combining the sediment-P and water-PO₄ data, the sorption characteristics of the sediment could be described. Those sorption characteristics allowed us to estimate a pre-industrial background surface water PO₄ levels, based on deeper sediment-P that stabilised at concentrations smaller than the modern. In three out of the four cores, the sediment-P peaked around 1980, coinciding with the peak in surface water PO₄. The estimated pre-industrial (~1800) background PO₄-concentration in the Scheldt river water was 62 [57; 66 (95%CI)] µg PO₄-P/L. That concentration exceeds the previously estimated natural background values for lakes in Flanders (15-35 µg TP/L) and is about half of the prevailing limit in the Scheldt river (120 µg PO₄-P/L). In the 1930s, river water concentrations were estimated at 140 [128; 148] µg PO₄-P/L, already exceeding the current limit. The method developed here proved useful for reconstructing historical, background PO₄ concentrations of a lowland tidal river. A similar approach can apply to other lowland tidal rivers to provide a scientific basis for local, catchment specific PO₄ backgrounds.
1 Introduction

Elevated phosphorus (P) concentrations can harm the ecological status in water by promoting excess levels of primary production, i.e. eutrophication, which yields subsequent anoxia, blooms of toxic blue-green algae and generally affects biodiversity (Correll, 1998). Therefore, limiting P concentrations in the surface water is crucial to ensure a good ecological status. Phosphorus in surface waters is present in solution (dissolved P) and in the suspended matter. The dissolved P can exist as inorganic phosphate (PO₄), organic P forms, or PO₄ sorbed to mineral colloids. The reactive P, determined by a colour reaction, is almost identical to PO₄. The environmental limits are either expressed as reactive P (equated to PO₄-P limits), as total P (TP) limits or both.

Many lowland regions, like Flanders (Belgium), do not achieve good water quality mainly due to the excess of nutrients. Currently, the average PO₄ in Flanders stabilised at 290 μg PO₄-P/L, well above the river type-specific limits of 70-140 μg PO₄-P/L (Smolders et al., 2017; VMM, 2018). Despite the current net-zero P-balance in agricultural soils, the situation did not improve since 2004. Therefore, the question arises when or, even, if these limits can be achieved. Natural background concentrations of lowland river PO₄-P may be higher than those of upland rivers because of biogeochemical processes typical for such waters. Ferric iron- (Fe(III)) and aluminium-oxyhydroxides have a high affinity for PO₄-anions and adsorb PO₄, limiting the P-solubility (Borggaard, 1990; Holtan et al., 1988). However, anoxic conditions lead to reductive dissolution of Fe(III) minerals. As a result, the associated P is released again to the overlying water when the sediment is strongly reduced (Baken et al., 2015). It is now well established that such conditions explain the typical summer peaks in PO₄ and that regional differences in sediment Fe concentrations explain regional differences in surface water PO₄ concentrations in Flanders (Smolders et al., 2017).

Since 2000, the European Union regulates surface water quality with the Water Framework Directive (WFD). The WFD does not prescribe limits for P in rivers and lakes but provides a framework for local regulations. The local maximum P concentrations are based on identifying pristine environments with minimal anthropogenic disturbance, i.e. the natural background (EU-Parliament, 2000). Flanders is densely populated, and it was not possible to locate pristine or reference lakes in the current environment. Instead, natural background TP concentrations for Flanders have been inferred from reference lakes sampled in Central and Baltic states in Europe that were identified for the WFD (Cardoso et al., 2007). Based on that study, background TP concentrations in lakes of Flanders were set at 15-35 μg P/L, selected for lakes with representative depth and alkalinity. For rivers, no TP or PO₄ natural background has been established.

The definition of the natural background has been subject to debate in many river basins (Matschullat et al., 2000; van Raaphorst et al., 2000). The natural background concentrations defined here are those concentrations found in the environment in the absence of any human activity, reflecting only natural geochemical processes (Laane, 1992; Reimann and Garrett, 2005). This definition implies that concentrations have to be estimated before the presence of human activity. This is not always feasible and, therefore, a pre-industrial background can be defined instead, which is inferred from samples dating before the industrial revolution (Reimann and Garrett, 2005).
The pre-industrial background is logically affected by anthropogenic processes. In Belgium, the industrial revolution started around 1800 (Vanhaute, 2003). At that time, the Belgian population was about three million, and large scale agriculture dates back to the middle ages and the Roman period. However, the most substantial increase in nutrient emissions occurred after the 1950s due to sewer infrastructure, mineral fertilisers and P-loaded detergents (Billen et al., 2005; Van Raaphorst et al., 2000). Because P emissions mainly originate from point sources due to domestic loading, the increase of surface water P between 1950 and 1975 relates to the rise in population connected to sewer systems (Billen et al., 2005). Since 1985, the increase of wastewater treatment allowed a significant improvement to the situation (Billen et al., 2005).

River floodplains provide sediment archives from which region-specific background concentrations could be derived. The sediments deposited onto floodplains bonded with surface water PO₄ and are deposited at a predictable rate. Those sediments can serve as an archive for reconstructing historical P emissions trends and provide useful spatial and temporal information on historical P concentrations in adjacent water bodies (Birch et al., 2008). Boyle et al. (2015) used P profiles from lake sediments in the UK to infer the historical evolution in population density in catchments over 10,000 years. Similarly, banded iron formations found in deep oceanic waters allowed to infer oceanic P concentrations from over two billion years ago (Bjerrum and Canfield, 2002).

In lowland rivers with tidal influence, also called estuaries such as the Scheldt estuary in Flanders, tidal marshes are the analogue of floodplains along fluvial rivers. Tidal marshes are vegetated ecosystems located along the tidal portion of rivers or coastlines, which periodically flood during high tide and storm events. Due to their net accumulation of sediments, tidal marshes increase in elevation over time (Temmerman et al., 2003a). Tidal marshes directly adjacent to tidal rivers are regularly flooded during high tides, so river sediments and associated elements like P are deposited within the dense marsh vegetation (Friedrichs and Perry, 2001; De Swart and Zimmerman, 2009; Temmerman et al., 2004a). Researchers have used tidal marshes as sediment archives of deposited substances other than P, such as organic carbon (Van de Broek et al., 2019) and silicon (Struyf et al., 2007).

However, it remains to be investigated to what extent P concentrations measured in tidal marsh sediment archives can be used to reconstruct historical changes in PO₄ concentrations in the adjacent estuary. Therefore, in this manuscript, we test and evaluate a methodology to estimate the historical surface water PO₄ concentrations. We described the tidal marsh sediment sorption characteristics by linking the P concentration in dated tidal marsh sediments to historical measurements of PO₄ in the Scheldt river water. Those sorption characteristics allowed us to estimate river water PO₄ concentrations based on a sediment analysis. The underlying assumption is that P remains immobile in these sediments and that the historical trend of PO₄ in the Scheldt river is reflected in this depth profile. Thereby, we argue that the sediment P-composition in deeper sediment layers of tidal marshes will reconstruct the historic PO₄ concentration of adjacent water bodies. A database containing measurements of the PO₄ concentration in the Scheldt river’s surface water (1967-current) will verify this assumption. The hypothesis we test is that the previously estimated natural background P in this major lowland river is larger than that what was estimated earlier for lakes (15-35 µg P/L) because these previous estimates did not account for the internal loading processes typical for lowland rivers.
2 Materials and methods

2.1 Study area

Freshwater tidal marshes were sampled at four locations along the Scheldt river (Fig. 1, Table S1). The Scheldt estuary is located in northern Belgium and the south-western Netherlands, where it flows into the North Sea. The river basin of the Scheldt covers a large part of Flanders (71%) and the adjacent region of Northern France; the total catchment area is approximately 22,000 km². The population living in the river basin is about 10 million (Meire et al., 2005). The tidal wave extends from the mouth (Vlissingen) to 160 km upstream near Ghent, where sluices stop the tidal wave. The estuary's freshwater tidal zone reaches from Ghent to Rupelmonde (Fig. 1). This research focused on freshwater tidal marshes, i.e. located in this freshwater tidal zone of the estuary.

Figure 1: Map of the Scheldt Estuary, triangles indicate the locations of the sampled tidal marshes, Old1 and Young1 were only 250 m apart, and on the scale of the map, they overlap

Sediment accreting in tidal marshes originates from the deposition of riverine suspended matter, including inorganic mineral sediment and organic matter (Callaway et al., 1996). For the period 1931-2002, the sediment accretion rates were 0.32-3.22 cm/y for freshwater tidal marshes (Temmerman et al., 2004b). We discriminate old and young tidal marshes, hereafter referred
to as marshes. Old marshes have a higher elevation compared to young marshes. As a general mechanism, young marsh surfaces accumulate sediments quickly and increase their elevation asymptotically up to an equilibrium level around the mean high water level (MHWL) (Pethick, 1981; Temmerman et al., 2003a). Temmerman et al. (2003a) defined an old marsh as visible on topographic maps of Ferraris (1774 - 1777), so it was formed before the 19th century (Temmerman et al., 2003a). Young marshes in the Scheldt estuary were formed more recently, by the natural establishment of pioneer marsh vegetation on formerly bare tidal mudflats, generally after 1944. During the last decades, the young marshes had a surface elevation below MHWL. Young marshes experienced more frequent inundations and therefore had larger sediment accretion rates than old marshes. Old marshes built up slower, at a rate comparable to the rise of the local MHWL (Temmerman et al., 2004a).

This study analysed depth profiles of sediment cores originating from tidal marshes along the freshwater Scheldt river. The analysis contained two old and two young marshes, referred to as Old1, Old2, Young1 and Young2 (locations indicated in Fig. 1). In total, we analysed eight cores; three replicate cores for both sites Old1 and Young1 and one core each for Old2 and Young2.

### 2.2 PO$_4$ concentration in surface waters

The IMIS (Flanders Marine Institute) provided surface water phosphate-P (PO$_4$-P) data measured colourimetrically on a filtered water sample and total phosphorus (TP) by acid digestion and a segmented flow analyser. Data of PO$_4$ concentrations in Scheldt river were available from 1967 up to 2019 compiled by the OMES-monitoring, who did additional quality controls on the data (ECOBE - UA; The Flemish Waterway, 2019). The PO$_4$ data originated from different sources described in supplementary information (SI.V) (De Pauw, 2007; ECOBE - UAntwerpen, 2007; Institute for Hygiene en Epidemiology (IHE), 2007; OMES: Monitoring physical-chemical water quality in the Zeeschelde, 2016; Van Meel, 1958). The open-source software R (R Core Team, 2020) was used to compile all available datasets for PO$_4$, closest to the study sites (Temse) and to calculate annual means by averaging all observations within each year for which data was available. The annual means of PO$_4$ were used to visualise the evolution of PO$_4$ in the Scheldt river (Fig. 2).
Figure 2. Concentrations of phosphate in the Scheldt River at Temse, annual means and standard deviation (error bar) around the annual mean. Samples were taken in Temse close to tidal marsh sites. (data sources: (ECOBE - UAntwerpen, 2007; Institute voor Hygiëne en Epidemiologie (IHE), 2007; Van Meel, 1958; Nv, 2016; De Pauw, 2007)

2.3 Sediment sampling

The sediment samples used in this study had been collected during a previous study in the Scheldt estuary about carbon sequestration in tidal marsh sediments (Van de Broek et al., 2018; Van De Broek et al., 2016). Collection of undisturbed sediment profiles on the tidal marshes took place between July and September 2016 (Old1, Young1, Old2, Young2; Fig. 1). Undisturbed sediment cores were taken using a gauge auger (0.06 m diameter) at each sampling location. In the field, the cores were divided into subsamples with a 0.03 m interval. The sediment samples were dried at a maximum temperature of 50 °C for 48 hours and crushed to a 2 mm grain size. Macroscopic vegetation residues were removed manually using tweezers (Van de Broek et al., 2018). Bulk density, grain size distribution and organic carbon (OC) content were analysed by Van de Broek et al. (2018). For detailed information, we refer to Van de Broek et al. (2016, 2018).
2.4 Sediment analysis

The dried sediment samples were analysed for oxalate-extractable P, Fe, Al and Mn (P$_{ox}$, Fe$_{ox}$, Al$_{ox}$, Mn$_{ox}$; Schwertmann, 1964). That acid oxalate extractant at pH = 3 targets poorly crystalline oxyhydroxides of Fe, Al and Mn and the associated P. The poorly crystalline oxyhydroxides are the most reactive due to their large specific surface area (Hiemstra et al., 2010). A sub-sample of each sediment sample was taken, and the extraction was made over two hours at a solid-liquid ratio of 1 g in 50 ml in an end-over-end shaker at 20°C (26 rpm). The suspension was filtered through a 0.45 µm membrane filter (CHROMAFIL® Xtra PET-45/25). The extract was diluted 20 times and measured by inductively coupled plasma optical emission spectrometry (ICP-OES). The degree of P-saturation (DPS; %) was calculated as in Eq. (1). The DPS represents the ratio of extractable (P$_{ox}$) to the P sorption capacity of the sediment. This P sorption capacity is estimated as half of the sum of oxalate Fe$_{ox}$ and Al$_{ox}$, because not all the Fe and Al in a soil is available for P sorption with Fe$_{ox}$, Al$_{ox}$ and P$_{ox}$ in molar units.

$$DPS = \frac{P_{ox}}{0.5(F_{e_{ox}} + A_{l_{ox}})} \times 100\%.$$  

(1)

The DPS is expressed in percentage and can be interpreted as the ratio of sorption sites on the sediment occupied by P. Previous research used the DPS to identify agricultural areas sensitive to phosphate leaching and showed a good correlation with pore water P concentrations (Breeuwsma et al., 1995; Lookman et al., 1995; Schoumans and Chardon, 2015; Schoumans and Groenendijk, 2000; van der Zee, 1988).

2.5 Age-depth model

The sediment analysis and the surface water PO$_4$ data had to be linked by a corresponding date and location to fit a sorption model. Therefore, an age-depth model was used to calculate the time since deposition of each sediment sample. Temmerman et al. (2004b, 2004a) developed a model (MARSED) to estimate sediment deposition rates and the resulting evolution of the sediment surface elevation in the tidal marshes of the Scheldt estuary. Hence, we could also use that model to determine the time since deposition of sediments throughout the sampled sediment profiles. The MARSED model simulates the tidal supply of suspended sediments and the settling of these sediments to the marsh surface during tidal inundation cycles integrated over the years to decades. The model was calibrated and validated against measured data on sediment deposition rates on the Scheldt estuary tidal marshes from 1945 until 2002 (Temmerman et al. 2003; 2004). The empirical data on sediment deposition rates were derived from radiometric and paleoenvironmental dating of sediment cores at the same locations as sampled for the present study (Temmerman et al., 2004a, 2004b). For our present study, we extrapolated the model simulations of sediment accretion from 2002 until 2016, the sampling date of the sediment cores (Van de Broek et al. 2018). We found that the MARSED model overestimated the observed marsh surface elevation 2016 (observed by RTK GPS surveying; Van de Broek et al. 2018; Poppelmonde, 2017) by 25 cm for sampling location Old1, 29 cm for Young1, 19 cm for Old2, and 8 cm for Young2. While the MARSED model was initially designed to simulate the overall behaviour of sediment accretion and surface elevation changes in tidal marshes in response to sea level rise scenarios (for which we accepted errors in the orders mentioned
in the previous sentence), for the present study, we wanted to estimate the time of sediment deposition throughout the sediment profiles as good as possible. Therefore, we rescaled the modelled marsh surface elevation by fitting it through the observed depth-age points from the GPS measured marsh surface elevation in 2016 and previously published radiometric and paleoenvironmental dating (Temmerman et al. 2003; 2004). This rescaling procedure is explained in the Supporting Information (Fig. S1, S2, S3, S4). The observed age-depth points were available starting from 1958 for sampling site Old1, 1947 for Young1, 1963 for Old2 and 1968 for Young2 (Temmerman et al. 2004).

An approximate extrapolation procedure was used to estimate the time of deposition of sediment from depths below the oldest measured age-depth points (mentioned in previous sentence). This extrapolation procedure was only applied for old marshes, which were defined as marshes that existed at least since the end of the 18th century (Temmerman et al. 2003a; 2004). Two sediment cores originated from such old tidal marshes (Old1 and Old2). We know from observed age-depth points that old marshes reached equilibrium with the MHWL before 1944, and that they built up their elevation after 1944 at a rate comparable to the rate of local MHWL rise (Temmerman et al., 2003a). Here, we assumed that between 1800 and 1944, these old marshes also accreted at a rate comparable to the MHWL rise. Historical tide gauge data of MHWL rise was available from 1901 for site Old1 and from 1930 for site Old2 (ScheldeMonitor Team and VNSC, 2020; Temmerman et al., 2003a) and a linear regression of the MHWL against time was used to estimate the marsh surface elevation before 1944 (Fig. S6, S7). However, the accuracy of the dating will be lower going further back in time. Such extrapolation to earlier dates is not appropriate for young marshes, as they only formed after 1950 by pioneer vegetation establishment on formerly bare mudflats (Temmerman et al. 2003a; 2004). Those mudflat sediment profiles do not have continuous sedimentary records as tidal marshes and are likely to be disturbed by erosion and sedimentation alternations (Belliard et al., 2019). Therefore, for the young marsh sampling locations, the sediment deposition time could not be extrapolated before the oldest available measured age-depth points, dating back to 1947 on location Young1 and 1968 on location Young2.

2.6 Relating surface water PO$_4$ with sediment P: the sediment-water model

The age-depth model and linear regression of MHWL provided a deposition year for each sediment sample. Through that age-depth relation the dataset of water PO$_4$ between 1967-2016 was linked to the sediment DPS for each core. The resulting dataset contained all available surface water PO$_4$ readings between 1967 and 2016 closest to the tidal marshes in Temse (n = 1932) and a corresponding DPS value of a sediment sample originating from one sediment core or, when available for Old1 and Young1, a mean of the replicate sediment samples. This dataset allowed to fit a sorption model further termed the sediment-water model. Schoumans and Groenendijk (2000) presented a Langmuir-type sorption model to predict PO$_4$ concentration leaching from a soil layer based on the DPS Eq. (2).

$$[PO_4] = K^{-1} \frac{DPS}{100-DPS},$$

(2)

With [PO$_4$] phosphate concentration in (kg L$^{-1}$), K the sorption constant (L kg$^{-1}$), DPS (degree of P-saturation; %). This model adequately described P sorption in soil across a wide range of pH values, including the Scheldt river pH (Schoumans and
Groenendijk, 2000; Warrinnier et al., 2018). The model relies on surface complexation between PO$_4$ and Fe-, Al-oxyhydroxides in the sediment, determined by a chemical equilibrium between solid (adsorbed) and dissolved PO$_4$ phase (Warrinnier et al., 2019). We did not use the existing sorption models for soils directly, and instead, the parameter K, in Eq. (2), was calibrated to the most recent Scheldt water data. So that, the K-value adapted to the geochemistry of the tidal marsh sediments.

We explored 16 different scenarios to fit the sediment-water model Eq. (2). These scenarios illustrate the statistical uncertainty surrounding the estimated PO$_4$ concentrations. The model was either fitted separately for each sediment core or on the combined cores Old1 and Young1 (SLVI). Every sediment sample had between one and three replicates, depending on the depth and the core. We entered either the average value of these replicates or the individual replicate DPS values. One sediment sample covered several deposition years, so multiple PO$_4$ observations corresponded with each sediment sample. Again, the average of all corresponding PO$_4$ readings was taken, or all available values were used separately, resulting in 16 models (Table S2). For each of these, the parameter estimation of Eq. (2) was fitted by nonlinear regression with JMP Pro (Version 15.1.0. SAS Institute Inc., Cary, NC, 1989-2019). Non-linear least squares regression to the PO$_4$-DPS data was used to estimate the model parameter (K) yielding the lowest sum of squared errors.

2.7 Evaluation Model Performance

The predictions of the sediment-water model were evaluated by plotting the measured surface water PO$_4$ against predicted PO$_4$ between 2007 and 2016 (Fig. S10). Additionally, the actual over-predicted median ratio was calculated for data points between 2007 and 2016. The prediction of recent years is interesting to evaluate the model’s performance because of two reasons. First, the most recent surface water PO$_4$-concentrations are relatively low and more representative of background concentrations. Second, the monitoring data have a high temporal resolution, and the age-depth model is more accurate at shallow depths.

3 Results

3.1 History of surface water PO$_4$ concentrations

The Scheldt PO$_4$-concentrations varied greatly over the past decades, with the peak in surface water PO$_4$-concentrations between 1975 and 1985 (Fig. 2). In Temse, the annual mean concentrations rose from 410 µg PO$_4$-P/L in 1967 and peaked in 1980 with 1570 µg PO$_4$-P/L. Between 1990 and 2003, a decrease followed the peak and over the last decade concentrations stabilised between 160 and 200 µg PO$_4$-P/L in Temse. Current PO$_4$-levels are a factor two lower than in 1967 and almost a factor ten lower than the peak in 1980 (Fig. 2; Table 1).
3.2 Sediment cores

The $P_{ox}$ in the sediments ranged between 370 mg P/kg and 13,000 mg P/kg, while the DPS ranged between 13% and 94% (Table 1). In all soil cores starting at the surface, the DPS and $P_{ox}$ increased with depth and peaked at about 0.5 m depth (Fig. S7, Fig. S8). In deeper (>1.0 m) sediment layers, $P_{ox}$ and DPS decreased and stabilised for Old1, Young1 and Young2 (Table 1). Overall, the $P_{ox}$ increased by, on average, a factor of 3.5 between the surface and the maximum concentrations (Fig. S8, Table 1). Based on the age-depth model these peak DPS were deposited between 1960 and 1985 in three of the four sediment cores (Fig. 2). Only the core Old2 reached the peak earlier (ca. 1940-1950). Most importantly for this work, DPS for Old1 showed an apparent stabilisation in deeper or older layers, which indicated undisturbed sediment layers (Fig. 3, Fig. S7).

Table 1: The sediment oxalate extractable P ($P_{ox}$) and its Degree of Phosphate Saturation (DPS) of the top, bottom, peak sediment layers at four different tidal marsh locations. Top layers are the sediments closest to the surface, peak layers had maximal $P_{ox}$ and DPS, and bottom layers are those sediments sampled at greatest depth. Values of $P_{ox}$ and DPS are means (± standard deviation) of N sediment samples, between top and bottom (cm) depth.

| Location | N | Top – Bottom (cm) | $P_{ox}$ (mg/kg) | DPS (%) |
|----------|---|-------------------|------------------|---------|
| Old1     |   | Top               | 4 0 – 9          | 2300 (± 2400) | 36 (± 3) |
|          |   | Peak              | 7 27 – 57        | 5400 (± 1300) | 70 (± 8) |
|          |   | Bottom            | 8 147 – 180      | 540 (± 110)  | 20 (± 4) |
| Young1   |   | Top               | 4 0 – 9          | 2700 (± 320)  | 37 (± 2) |
|          |   | Peak              | 6 27 – 57        | 8500 (± 3200) | 85 (± 15)|
|          |   | Bottom            | 6 129 – 144      | 910 (± 440)  | 40 (± 4) |
| Old2     |   | Top               | 3 0 – 9          | 2800 (± 90)   | 45 (± 1) |
|          |   | Peak              | 3 54 – 69        | 8000 (± 1600) | 94 (± 2) |
|          |   | Bottom            | 3 132 – 147      | 1700 (± 620) | 43 (± 6) |
| Young2   |   | Top               | 3 0 – 9          | 2700 (± 410)  | 23 (± 1) |
|          |   | Peak              | 3 48 – 63        | 7000 (± 1200) | 55 (± 7) |
|          |   | Bottom            | 3 144 – 183      | 3200 (± 110) | 31 (± 2) |
Within the first meter, Fe$_{ox}$ was stable in the three soil cores (Old1, Young1, Old2) with concentrations around 20,000 mg/kg, except for Young2 for which Fe$_{ox}$ was a factor two larger (Fig. S9). For Young1 and Young2 a decrease in Fe$_{ox}$ concentration occurred at depths > 1 m. For Old1, Fe$_{ox}$ showed a steady decline from 20,000 mg/kg at the surface to 10,000 mg/kg at the bottom of the profile (Fig. S9). The Al$_{ox}$ concentrations showed a similar trend as the P$_{ox}$ concentrations, with an initial increase followed by a decrease with depth. The strong correlations of Al$_{ox}$ and Fe$_{ox}$ with P$_{ox}$ ($r_{Al} = 0.73$ and $r_{Fe} = 0.65$) illustrate the positive effect of Fe and Al oxyhydroxides on P sorption.

![Figure 3: The Degree of Phosphate Saturation (DPS) timeline based on four tidal marsh sediment sites. Each dot represents a sediment analysis, the year assigned to each sediment analysis was calculated with the age depth model. Before 1930, no model dates are available, therefore a linear regression of the MHWL was used to extrapolate the dates for the old marshes. Dates before 1930, are increasingly uncertain going further back in time. For young marshes such extrapolation was not possible, the points before formation of the marshes are indicated with a question mark.](https://doi.org/10.5194/bg-2021-100)
3.3 Sediment core selection

Under the assumption that PO₄ does not migrate, the tidal marsh sediment cores can provide an archive for river water PO₄. However, that assumption may be most violated at two locations, Old2 and Young2. Considering P-migration, evaluating the distance from a creek within the tidal marsh is crucial. That distance is essential for two reasons. First, within 10 to 20 m of the creeks, the groundwater table fluctuates largely with the tides (Van Putte et al., 2020), which can induce vertical P-migration. Secondly, sediment accretion is more difficult to predict at closer distances to the creeks and can affect the age-depth relation (Temmerman et al., 2003b). The distance to the nearest creek within the tidal marshes are 21 m for Old1, 56 m for Young1, 35 m for Young2 and 5 m for Old2. Furthermore, the profile of Old2 had a peak of Pox at an earlier date (1950) than was expected from surface water data (1980), indicating P-migration (Fig. 2). Consequently, Old2 was not taken up for interpretation of the relation between DPS and PO₄. For core Young2, deeper sediment layers had a larger DPS compared to the surface layers (Table 2). Additionally, the age estimation of sediments older than 1968 was not possible due to this tidal marsh’s young age, which hampers the interpretation of DPS values from deeper layers as representative for background values. Furthermore, Feox concentrations were a factor two larger than the other cores (Fig. S9) and a factor two larger than the average sediment Fe-concentration of the Upper-Scheldt basin (VMM, 2019). The local enrichment in iron lowers the DPS values and makes the core less representative of the average situation in the Scheldt. These observations made Young2 inappropriate to fit the relation between DPS and PO₄.

The two remaining soil cores, Old1 and Young1, originate from the same tidal marsh area named The Notelaer, located near the city of Temse (Fig. 1) and has been the topic of multiple studies on sediment accretion (Temmerman et al., 2004b, 2003a) and soil OC stocks (Van de Broek et al., 2018; Van De Broek et al., 2016). The sediment profiles of both sites Old1 and Young1 rise and fall in DPS comparable to dynamics in surface water PO₄-concentrations (Fig. 2, Fig. 3). In deeper sediment layers, DPS and Pox stabilise below levels of recent deposits (Fig. S7, S8). These observations indicated a well-preserved Pox and DPS profile, essential for the DPS-PO₄ relation. Therefore, Old1 and Young1 are considered the best profiles for applying the sediment-water model and interpretation of background concentrations.
Table 2: The predicted pre-industrial background concentrations of phosphate (PO₄-P µg/L) in the Scheldt river based on the Degree of Phosphate Saturation (DPS) in the sediment layers of marsh Old1, dating back to 1800 (pre-industrial), where DPS values stabilised with depth at 20%, and the predicted background dated to 1930 where DPS stabilised at 36%. Conversion of DPS to river phosphate concentration based on the association of DPS with PO₄-P calibrated to data 1967-2016, thereby using different calibration for sediment-water models; the details of models are in Table S2. Model 3b (in bold), is proposed as the most accurate one (see text).

| Model # | K (L/kg) [95% CI] | Pre-industrial background µg PO₄-P/L [95% CI] | µg PO₄-P/L 1930 [95% CI] | Median ratio predicted / observed-PO₄ (2007-2016) |
|---------|------------------|---------------------------------------------|---------------------------|---------------------------------------------|
| 1b      | 2.1×10⁶ [2.0×10⁶; 2.3×10⁶] | 120 [110; 130] | 270 [245; 281] | 1.67                                      |
| 2b      | 4.9×10⁶ [4.6×10⁶; 5.2×10⁶] | 51 [49; 54] | 120 [109; 122] | 0.75                                      |
| 3b      | 4.1×10⁶ [3.8×10⁶; 4.4×10⁶] | 62 [57; 66] | 140 [128; 148] | 0.89                                      |

3.4 Sediment-water model fit

The sediment-water model Eq. (2) was fitted on DPS-PO₄ data from the different sediment cores both individually and for a combination of cores (Table S2). Two observations were omitted because the DPS values were too large (0.99 – 1.02) and produced artefacts in the results. The Nash-Sutcliffe model efficiency (E) of the models ranged between 0.04 and 0.85 depending on the input data (Table S2; Nash and Sutcliffe, 1970). The sediment-water model was fitted on the data of each core separately and for the combination of the data from Old1 and Young1, as they came from the same tidal marsh location. The models fitted on data from sites Old2 and Young2 were not considered as migration possibly affected those cores (cf. section 3.3).

The models fitted on an average DPS (across replicates) associated with individual PO₄ readings were considered most suitable (Models 1b, 2b, 3b; Table S2). Models using unique DPS associated with single PO₄-data duplicated or even triplicated the PO₄-data, artificially creating more degrees of freedom (Model 1c, 2c, 3c; Table S2). Using mean PO₄-values artificially reduced the degrees of freedom, compromising the model predictions, by increasing RSE and widening confidence intervals (Models 1a, 2a, 3a; Table S2). The fitted parameter K (L/kg) ranged between 1.0×10⁶ and 5.4×10⁶ for the different input datasets, with the 95% confidence intervals ranging between 0.8×10⁶ and 7.2×10⁶. The variation of parameter K for the different input datasets was larger than the individual confidence limits variation (Table S2). Thus, the uncertainty was more pronounced due to the variability in sediment samples than due to the model fit.
3.5 Model performance

The sediment-water model performance was evaluated by comparing the actual by predicted PO$_4$ concentrations over the last decade, as those PO$_4$ concentrations are more comparable to the background. Therefore, we calculated the ratio of predicted over observed PO$_4$ in the surface water of the Scheldt river between 2007 and 2016 and plotted actual by predicted plots (Fig. S10; Table 2). A ratio of predicted over observed below one of indicates that the model underestimates ambient PO$_4$-concentrations (mean Temse [2007-2016] = 170 µg PO$_4$-P/L). Model 1b overestimated recent observations, which is unwanted for calculating the background, and was therefore considered unsuitable. In contrast, half of the observations were underestimated by at least 25% for model 2b and by 11% for model 3b (Table 2). The actual by predicted plots illustrate the same (Fig. S10). Based on these observations, model 3b was considered as the best model, although the residual standard error (RSE) was lower for model 2b (Table S2). Model 3b predicted recent PO$_4$-concentrations best, with median underestimation of only 11% (Table 2). The selected model 3b succeeds to reconstruct the rise and fall in surface water PO$_4$-concentrations based on the sediment characteristic DPS (Fig. 4).

Figure 4: Measured (black points) and predicted (grey crosses; +) of PO$_4$-P concentrations (µg/l) in the Scheldt river in Temse. The concentrations are calculated from the sediment phosphate saturation (DPS) of the tidal marshes at Old1 and Young1, using sediment-water model 3b.
Maxima of monitored and predicted PO₄-concentrations coincide in time and have a similar size. For example, in 1973, the average PO₄ concentration predicted by the model was 1200 µg PO₄-P µg/L and measured concentrations was on average 1300 µg PO₄-P µg/L. The maximal predicted PO₄ concentration was 2200 µg PO₄-P/L, while the maximal observed was 3000 µg PO₄-P/L. Predictions for recent years are within 15% of the observed data (e.g. 2015: Model: 133 µg PO₄-P/L, Measured 155 µg PO₄-P/L). Between 1940 and 1990, the modelled PO₄-concentrations show more variation. Likewise, monitored PO₄-data are spread more between 1967 and 1990 (Fig. 2). Before 1930, modelled PO₄-concentrations stabilise at levels below current observations (Fig. 4).

3.6 Estimating background PO₄ concentrations in the Scheldt river

The deepest sediment layers are most suitable for predicting background PO₄ concentrations of the Scheldt river water. These layers are the oldest and expected to have experienced the least impact of P additions from anthropogenic sources. The Old1 marsh site was appropriate for this purpose as it developed before 1774, which dates before the industrial revolution in Belgium. The average DPS for the bottom sediments, dated between 1800 and 1840, was 20% for core Old1 (Table 1; Fig. 2); these samples are considered to represent the pre-industrial background. That DPS value produced PO₄-concentrations of 62 µg PO₄-P/L [95%CI (57; 66)] for the pre-industrial background, using sediment-water model 3b. The sediment dated to 1930 had a DPS of 36%, for that value the same sediment-water predicted a PO₄-concentration of 140 µg PO₄-P/L [95%CI (128; 148)] (Table 2).

4 Discussion

4.1 Background vs ambient PO₄ concentrations

This work presents a novel approach to reconstruct background surface water PO₄ concentration in a tidal river using the DPS of adjacent tidal marsh sediments. The background concentration is essential in the context of developing local nutrient limits. The predicted pre-industrial background concentration (62 µg PO₄-P/L; Table 2) is about half of the current surface limit of the Scheldt (120 µg PO₄-P/L; Flemish Government, 1995). Remarkably, the predicted background concentrations are about a factor two larger than the background estimates of lake waters for Flanders today (15-35 µg PO₄-P/L; Cardoso et al., 2007), suggesting that the internal loading related to summer anoxia in lowland rivers, contributes naturally to larger PO₄ concentrations in lowland rivers (see introduction).

Our analysis suggests that the pre-industrial PO₄ concentration was about three times lower than the current concentrations in the Scheldt. The mean PO₄ concentration of the Scheldt in Temse was 170 µg PO₄-P/L between 2007 and 2016. However, in the 1930s, the concentration was estimated at 140 µg PO₄-P/L and larger than current limits, at a time before widespread connection to sewer systems, P-loaded detergents, and application of mineral fertilisers.
4.2 Limitations of the model

Care needs to be taken with background extrapolations to ensure that post-depositional processes have not modified the biogeochemical patterns and that the area represents the area of interest (Reimann and Garrett, 2005). Several factors can obscure the reconstructed background concentrations. First, vertical migration of P can enrich deeper sediment layers, thereby causing an overestimation of the background. The sediment profiles at the tidal marshes are almost permanently saturated, and the intrusion of P-rich groundwater could affect the P concentrations in the tidal marsh sediment. Moreover, periodic flooding occurs at an approximate range of 300-350 inundations per year, depending on the tidal marsh elevation (Temmerman et al., 2003b). These conditions could favour phosphorus migration due to reductive dissolution of Fe (oxy)hydroxides (Baken et al., 2015). We removed two cores with indications of PO$_4$ migration to address the issue (Old2 and Young2). These cores were identified by comparing the peak in the DPS age-profile with the known peak in surface water PO$_4$ concentrations in the 1980s and by considering the distance from nearby creek (Fig. 2; Fig. 1). Additionally, the DPS levels of the deepest sediment layers were compared with layers at the surface. For one core the surface layers had lower DPS levels than the deepest layers (Young 2). The two remaining cores (Old1, Young1) had lower DPS-levels in deeper sediment layers (Fig. S7). More importantly, the modelled peak in PO$_4$ concentrations based on the cores Old1, Young1 were found within two years of the monitored peak and had a similar magnitude (Fig. 4). The coinciding peaks illustrate little migration of PO$_4$ in Old1 and Young1, thereby justifying these cores as an archive for water-PO$_4$

The limited migration is also logical: at the average DPS of 90 % in sediment showing at the peak, the sorption models predict that the solid-liquid P concentration ratio is 2900 L/kg, at the average K value of models of Table 2. That ratio can be converted to dimensionless retardation factor, representing the ratio of the distance migrated by the PO$_4$ compared to the distance traveled by percolating water, of 7500 at a bulk density ($\rho_b$) of 1.3 and porosity ($\theta$) of 0.5. With a net vertical annual water percolation of about 2 meters, this corresponds to a net vertical P migration rate of 2.5 cm over 100 years, i.e. vanishingly small (calculation details not shown).

Secondly, there is uncertainty on the age-depth estimation of the sampled sediment profiles. We expect that the age-depth model is most reliable for the Young1 sediment core, as it is based on a fitting of a modelled age-depth relation to four observed age-depth points, while we only had two observed age-depth points available for the other cores. Additionally, observed age-depth points were not older than 1944. Hence, extrapolation of the age-depth model to periods before the older available age-depth points is increasingly uncertain.

4.3 Pre-industrial and natural background values

The population increase between 1800 and 1930 can provide a first, and very crude estimate of the population-DPS relation in the Scheldt basin. In 1800 the population in Belgium was around 3 million. In 1930, this number has more than doubled to 7 million (Vanhaute, 2003). A linear relation between both suggests that the DPS is 8% for the pre-anthropogenic pristine environment, corresponding with a PO$_4$-concentration 19-41 µg PO$_4$-P/L, i.e. close to what researchers have suggested for
pristine lakes. Clearly, such predictions need to be corroborated with older sediment observations and other archaeological information. The Scheldt river is logically more aerated than several smaller lowland rivers where summer anoxia are naturally more present, i.e. the pristine PO$_4$-P values will be likely higher.

5 Conclusions

Our study illustrated that tidal marsh sediments can provide an estimate of pre-industrial background PO$_4$-concentrations of the freshwater rivers like the Scheldt river. A sediment assessment can record and time-integrate environmental events, which provides useful spatial and temporal information. Our data estimated that the pre-industrial background concentration is about half of the environmental limits set for surface waters in Flanders and neighbouring countries. Around 1930, the PO$_4$ levels were only about 20% lower than today, which is a remarkably large concentration at a time before the massive application of mineral fertilisers, with lower population density and limited connection to sewer systems. The current PO$_4$ concentrations decreased by a factor ten from the peak found about 40 years ago, reflecting wastewater treatment efforts and reduced diffuse P emission. It is also clear from this study that the pristine, pre-anthropogenic PO$_4$-P concentrations in the Scheldt river are well below the current ambient ones.

Data availability

The results of the sediment data analysis and age depth model are provided in the supplement as csv format. Results of Surface water data are available upon request at the The IMIS (Flanders Marine Institute).

Author contribution

FL, ES and PC designed the research. FL conducted the investigation process, and developed the methodology under supervision of ES. MVDB carried out the fieldwork and conceptualized the use of the samples. ST provided the methodology for the age-depth model and software. TM validated the use of the surface water data. EVM and FL placed the results in perspective with historical data. All the authors contributed to discussion and data interpretations, review and editing of the work.

Competing interests

The authors declare that they have no conflict of interest
Acknowledgements

This project was supported by the Research Fund Flanders (FWO), project G089319N. The results of this research greatly depended on the data collected by the OMES-monitoring and The Flemish Waterway. Many years of intensive data collection and quality assessment of the Scheldt river resulted in a unique and valuable phosphate time series. We have the utmost respect for their work and are thankful we could apply the dataset for this research. We acknowledge Dries Grauwels and Kristin Coorevits for technical assistance. Finally, thanks to the Scheldt in providing this beautiful sediment archive, to travel back in time and explore environmental history.

References

Baken, S., Verbeeck, M., Verheyen, D., Diels, J. and Smolders, E.: Phosphorus losses from agricultural land to natural waters are reduced by immobilization in iron-rich sediments of drainage ditches, Water Res., 71, 160–170, doi:10.1016/j.watres.2015.01.008, 2015.

Belliard, J. P., Silinski, A., Meire, D., Kolokythas, G., Levy, Y., Van Braeckel, A., Bouma, T. J. and Temmerman, S.: High-resolution bed level changes in relation to tidal and wave forcing on a narrow fringing macrotidal flat: Bridging intra-tidal, daily and seasonal sediment dynamics, Mar. Geol., doi:10.1016/j.margeo.2019.03.001, 2019.

Billen, G., Garnier, J. and Rousseau, V.: Nutrient fluxes and water quality in the drainage network of the Scheldt basin over the last 50 years, Hydrobiologia, doi:10.1007/s10750-004-7103-1, 2005.

Birch, G. F., McCready, S., Long, E. R., Taylor, S. S. and Spyrrakis, G.: Contaminant chemistry and toxicity of sediments in Sydney Harbour, Australia: Spatial extent and chemistry-toxicity relationships, Mar. Ecol. Prog. Ser., doi:10.3354/meps07445, 2008.

Bjerrum, C. J. and Canfield, D. E.: Ocean productivity before about 1.9 Gyr ago limited by phosphorus adsorption onto iron oxides, Nature, doi:10.1038/417159a, 2002.

Borggaard, O. K.: Dissolution and adsorption properties of soil iron oxides., Royal Veterinary and Agricultural University., 1990.

Breeuwsma, A., Reijerink, J. G. A. and Schoumans, O. F.: Impact of manure on accumulation and leaching of phosphate in areas of intensive livestock farming, in Animal waste and the land-water interface., 1995.

Van de Broek, M., Vandendriessche, C., Poppelemonde, D., Merckx, R., Temmerman, S. and Govers, G.: Long-term organic carbon sequestration in tidal marsh sediments is dominated by old-aged allochthonous inputs in a macrotidal estuary, Glob. Chang. Biol., doi:10.1111/gcb.14089, 2018.

Van de Broek, M., Baert, L., Temmerman, S. and Govers, G.: Soil organic carbon stocks in a tidal marsh landscape are dominated by human marsh embankment and subsequent marsh progradation, Eur. J. Soil Sci., doi:10.1111/ejss.12739, 2019.

Van De Broek, M., Temmerman, S., Merckx, R. and Govers, G.: Controls on soil organic carbon stocks in tidal marshes along
an estuarine salinity gradient, Biogeosciences, doi:10.5194/bg-13-6611-2016, 2016.

Callaway, J. C., Nyman, J. A. and DeLaune, R. D.: Sediment accretion in coastal wetlands: A review and a simulation model of processes, Curr. Top. Wetl. Biogeochem., 1996.

Cardoso, A. C., Solimini, A., Premazzi, G., Carvalho, L., Lyche, A. and Rekolainen, S.: Phosphorus reference concentrations in European lakes, in Hydrobiologia, vol. 584, pp. 3–12., 2007.

Correll, D. L.: The Role of Phosphorus in the Eutrophication of Receiving Waters: A Review, J. Environ. Qual., 27(2), 261–266, doi:10.2134/jeq1998.00472425002700020004x, 1998.

ECOBE - UAntwerpen: AZ monitoring water quality of the Scheldt. [online] Available from: http://www.vliz.be/en/imis?module=dataset&dasid=1468 (Accessed 28 September 2020), 2007.

EU-Parliament: DIRECTIVE 2000/60/EC OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 23 October 2000 establishing a framework for Community action in the field of water policy., 2000.

Flemish Government: VLAREM II, Vlarem II, EMIS Navig. [online] Available from: https://navigator.emis.vito.be/mijn-navigator?woId=263 (Accessed 23 December 2020), 1995.

Friedrichs, C. T. and Perry, J. E.: Tidal Salt Marsh Morphodynamics: A Synthesis, J. Coast. Res., 2001.

Hiemstra, T., Antelo, J., Rahnemaie, R. and Riemsdijk, W. H. va.: Nanoparticles in natural systems I: The effective reactive surface area of the natural oxide fraction in field samples, Geochim. Cosmochim. Acta, 74(1), 41–58, doi:10.1016/j.gca.2009.10.018, 2010.

Holtan, H., Kamp-Nielsen, L. and Stuanes, A. O.: Phosphorus in soil, water and sediment: an overview, Hydrobiologia, 170(1), 19–34, doi:10.1007/BF00024896, 1988.

Institute voor Hygiëne en Epidemiologie (IHE): Scheldt water quality data. [online] Available from: http://www.vliz.be/en/imis?module=dataset&dasid=1438 (Accessed 28 September 2020), 2007.

Laane, R. W. P. M.: Background concentrations of natural compounds in rivers, sea water, atmosphere and mussels, The Hague. [online] Available from: http://publicaties.minienm.nl/documenten/background-concentrations-of-natural-compounds-in-rivers-sea-wat (Accessed 19 October 2020), 1992.

Lookman, R., Vandeweert, N., Merckx, R. and Vlassak, K.: Geostatistical assessment of the regional distribution of phosphate sorption capacity parameters (FeOX and AlOX) in northern Belgium, Geoderma, 66(3–4), 285–296, doi:10.1016/0016-7061(94)00084-N, 1995.

Matschullat, J., Ottenstein, R. and Reimann, C.: Geochemical background - Can we calculate it?, Environ. Geol., 39(9), 990–1000, doi:10.1007/s002549900084, 2000.

Van Meel, L.: hydrobiology of the Sea-Scheldt near Liefkenshoek. [online] Available from: http://www.vliz.be/en/imis?module=dataset&dasid=1412 (Accessed 28 September 2020), 1958.

Meire, P., Ysebaert, T., Van Damme, S., Van Den Bergh, E., Maris, T. and Struyf, E.: The Scheldt estuary: A description of a changing ecosystem, Hydrobiologia, 540(1–3), 1–11, doi:10.1007/s10750-005-0896-8, 2005.

Nv, E.-U. D. V. W.: OMES: Monitoring fysical-chemical water quality in the Zeeschelde. [online] Available from:
De Pauw, C.: The environment and plankton of the WesterScheldt estuary, Ghent. [online] Available from: http://www.vliz.be/en/imis?module=dataset&dasid=1069, 2016.

Pethick, J. S.: Long-term accretion rates on tidal salt marshes., J. Sediment. Petrol., doi:10.1306/212F7CDE-2B24-11D7-8648000102C1865D, 1981.

Poppelmonde, D.: Organic carbon dynamics in tidal marshes of the Scheldt estuary A combined field and modelling approach, KULeuven, VUB., 2017.

R Core Team: R: A language and environment for statistical computing, [online] Available from: https://www.r-project.org/, 2020.

van Raaphorst, W., de Jonge, V. N., Dijkhuizen, D. and Frederiks, B.: Natural background concentrations of phosphorus and nitrogen in the Dutch Wadden Sea, Rapp. voor Kust en Zee, 53pp., 2000.

Reimann, C. and Garrett, R. G.: Geochemical background - Concept and reality, Sci. Total Environ., doi:10.1016/j.scitotenv.2005.01.047, 2005.

ScheldeMonitor Team and VNSC, R. & M.: Data downloaded from ScheldeMonitor: a data portal with information, data and products on the Scheldt Estuary, Data downloaded from ScheldeMonitor a data portal with information, data Prod. Schelt Estuary [online] Available from: https://rshiny.scheldemonitor.org/waterniveauschelde/ (Accessed 12 April 2021), 2020.

Schoumans, O. F. and Chardon, W. J.: Phosphate saturation degree and accumulation of phosphate in various soil types in The Netherlands, Geoderma, 237, 325–335, doi:10.1016/j.geoderma.2014.08.015, 2015.

Schoumans, O. F. and Groenendijk, P.: Modeling Soil Phosphorus Levels and Phosphorus Leaching from Agricultural Land in the Netherlands, J. Environ. Qual., 29(1), 111–116, doi:10.2134/jeq2000.00472425002900010014x, 2000.

Schwertmann, U.: Differenzierung der Eisenoxide des Bodens durch Extraktion mit Ammoniumoxalat-Lösung, Zeitschrift für Pflanzenernährung, Düngung, Bodenkd., 105(3), 194–202, doi:10.1002/jpln.3591050303, 1964.

Smolders, E., Baetens, E., Verbeeck, M., Nawara, S., Diels, J., Verdievel, M., Peeters, B., De Cooman, W. and Baken, S.: Internal Loading and Redox Cycling of Sediment Iron Explain Reactive Phosphorus Concentrations in Lowland Rivers, Environ. Sci. Technol., 51(5), 2584–2592, doi:10.1021/acs.est.6b04337, 2017.

Struyf, E., Temmerman, S. and Meire, P.: Dynamics of biogenic Si in freshwater tidal marshes: Si regeneration and retention in marsh sediments (Schelde estuary), Biogeochemistry, doi:10.1007/s10533-006-9051-5, 2007.

De Swart, H. E. and Zimmerman, J. T. F.: Morphodynamics of tidal inlet systems, Annu. Rev. Fluid Mech., doi:10.1146/annurev.fluid.010908.165159, 2009.

Temmerman, S., Govers, G., Meire, P. and Wartel, S.: Modelling long-term tidal marsh growth under changing tidal conditions and suspended sediment concentrations, Scheldt estuary, Belgium, Mar. Geol., doi:10.1016/S0025-3227(02)00642-4, 2003a.

Temmerman, S., Govers, G., Wartel, S. and Meire, P.: Spatial and temporal factors controlling short-term sedimentation in a salt and freshwater tidal marsh, scheldt estuary, Belgium, SW Netherlands, Earth Surf. Process. Landforms, doi:10.1002/esp.495, 2003b.
Temmerman, S., Govers, G., Wartel, S. and Meire, P.: Modelling estuarine variations in tidal marsh sedimentation: Response to changing sea level and suspended sediment concentrations, Mar. Geol., doi:10.1016/j.margeo.2004.10.021, 2004a.

Temmerman, S., Govers, G., Meire, P. and Wartel, S.: Simulating the long-term development of levee-basin topography on tidal marshes, Geomorphology, 63(1–2), 39–55, doi:10.1016/j.geomorph.2004.03.004, 2004b.

Vanhaute, E.: en arbeid in België in de ’ lange negentiende eeuw ’, 118(2001), 153–178, 2003.

Warrinnier, R., Goossens, T., Braun, S., Gustafsson, J. P. and Smolders, E.: Modelling heterogeneous phosphate sorption kinetics on iron oxyhydroxides and soil with a continuous distribution approach, Eur. J. Soil Sci., 69(3), 475–487, doi:10.1111/ejss.12549, 2018.

Warrinnier, R., Goossens, T., Amery, F., Vanden Nest, T., Verbeeck, M. and Smolders, E.: Investigation on the control of phosphate leaching by sorption and colloidal transport: Column studies and multi-surface complexation modelling, Appl. Geochemistry, 100, 371–379, doi:10.1016/j.apgeochem.2018.12.012, 2019.

van der Zee, S. E. A. T. M.: Transport of reactive contaminants in heterogeneous soil systems., 1988.