Morphological effects of vegetation on the fluvial-tidal transition in Holocene estuaries

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Abstract. Vegetation enhances bank stability and sedimentation to such extent that it can modify river patterns, but whether similar strong biogeomorphological feedbacks exist in estuarine environments is poorly understood. On the one hand, tidal flats accrete faster in the presence of vegetation, reducing the flood storage and ebb-dominance over time, while flow-focusing effects of tidal floodplain elevated by mud and vegetation could lead to channel concentration and incision. Here we study isolated and combined effects of mud and saltmarsh vegetation on estuary dimensions. A 2D hydromorphodynamic sandy estuary model was developed, which was coupled to a vegetation model and used to simulate 100 years of morphological development. Vegetation settlement, growth and mortality were determined by the hydromorphodynamics. Eco-engineering effects of vegetation on the physical system are here limited to hydraulic resistance, which affects erosion and sedimentation pattern through the flow field. We investigated how vegetation, combined with mud, affects the average elevation of tidal flats and controls the system-scale planform. Results show that the vegetation reaches its largest extent in the mixed energy zone of the estuary. Here vegetation can cover more than 50% of the estuary width while it remains below 10-20% in the outer, tide dominated zone. Aerial image analysis shows general agreement with trends in natural estuaries. The presence of mud leads to stabilization and accretion of the intertidal area and a slight infill of the mixed energy zone, which acts as a bed load convergence zone at the fluvial-tidal transition. Without mud, the modelled vegetation has a limited effect, again peaking in the mixed energy zone. Combined modelling of mud and vegetation leads to mutual enhancement with mud causing new colonization areas and vegetation stabilizing the mud. While vegetation focusses the flow into the channels such that mud sedimentation in intertidal side channels is prevented on a timescale of decades, the filling of intertidal area and resulting reduction of tidal prism may cause infilling of estuaries over centuries.

1 Introduction

1.1 Problem definition

Estuaries are flanked by tidal marshes, which are unique ecosystems with a very high biomass that modify the local hydromorphodynamic conditions (Davidson et al., 1991; Meire et al., 2005; Friedrichs, 2010). It is well known that vegetation affects hydromorphodynamics in rivers (Corenblit et al., 2009; Oorschot et al., 2015), and this has also been shown on the scale of
individual tidal marshes (Bouma et al., 2005; D’Alpaos et al., 2006; Temmerman et al., 2007). The effect of vegetation on hydromorphodynamics in tidal marshes is therefore relatively well known on the patch or individual plant scale (Järvelä, 2002; Siniscalchi et al., 2012), while its effect on estuary scale morphodynamics has barely been studied. Incorporating vegetation in estuarine morphodynamic models is considered one of the three biggest challenges to overcome (Coco et al., 2013). A comprehensive but qualitative model suggests that marshes reach their largest extent in the mixed energy zone of the estuary (Dalrymple et al., 1992). Here we investigate whether plant species collectively can have eco-engineering effects that are significant enough to modify entire estuarine landscapes.

Our hypothesis results from a combination of three independent and complementary analyses. First, a reconstruction of the Holocene development of estuaries and tidal basins suggests that vegetation combined with mud can cause a positive feedback on estuary size. Through reduction of intertidal water storage at the system margins, due to vegetation-enhanced sedimentation, the tidal prism reduces and tends towards flood-dominant transport (Speer and Aubrey, 1985; Friedrichs and Perry, 2001; Friedrichs, 2010). Second, Leuven et al. (2017) showed on the basis of imagery of a large number of estuaries that all space wider than that covered by an ideal convergent estuary is filled with tidal bars. This analysis excluded tidal marshes but clearly a number of estuaries were larger in the past and have at least partly been filled by mud flats, saltmarsh or mangroves. A model study by Braat et al. (2017) on effects of mud on system-scale development of estuaries over millennia showed that mud decreases morphodynamics and decreases total system width depending on mud concentration. All three approaches, geological, remote sensing and numerical, point at system-scale effects of mud and vegetation in estuaries.

Our aims are to determine the combined effects of mud and vegetation on estuarine planform and morphodynamics, specifically in the setting of a sandy estuary with mud input from the river. To this end we will use a numerical model for century-scale simulation of flow, sediment transport, morphology and vegetation. We ignore binding of sediment by roots because of the relatively shallow rooting and only explore cohesive effects of mud, floodplain-filling effects of mud and flow resistance effects of vegetation. This allows us to pragmatically apply an existing model for riparian vegetation to the tidal environment. Two questions of specific interest are what explains the zonation of vegetation as found by Dalrymple et al. (1992), and what are the morphological and hypsometric changes as a result of presence of vegetation. First we will review known effects of vegetation and mud, which results in specific hypotheses for vegetation zonation and morphodynamic effects that are subsequently tested with a 2DH numerical model.

1.2 Review and hypothesis development

In rivers, riparian vegetation stabilizes channels by reducing floodplain flow and adding bank strength to the floodplains (Coenblit et al., 2009; Gurnell et al., 2012). These eco-engineering effects can be strong enough to cause the transition from braiding towards meandering or even sinuous rivers (Ferguson, 1987; Tal and Paola, 2007; Dijk et al., 2013; Oorschot et al., 2015). However, presence of vegetation can also cause bifurcation of channels by stabilizing bar tips, causing flow resistance on pointbars and diverging the flow from the channel onto the floodplain (Burge, 2005; Dijk et al., 2013). Furthermore this increased flow resistance drives an increase in water height, which may induce flooding events (Darby, 1999; Kleinhans et al., 2018). The presence of mud has a partly similar effect as vegetation because it can lead to stabilization of systems as well, and
Figure 1. Active and vegetated parts of estuaries, showing proportionally more vegetated area in the upstream transition from single-thread river to multi-thread estuary. The estuaries are the Dovey (UK), Columbia (USA) and Gannel (UK).
mud has shown to preferentially accumulate at vegetated areas (Kleinhans et al., 2018). Based on these insights and general similarities between rivers and the fluvial-tidal transition, it is well-conceivable that similar biogeomorphological interactions shape upstream parts of estuaries. While salinity is an important variable determining which species prevail, here we focus on a single and often dominant saltmarsh vegetation species.

Saltmarsh vegetation flanks estuaries from the brackish zone to the mouth. Saltmarsh enhances sedimentation both through reduced flow velocities and through particle capture, somewhat comparable to what happens on river floodplains, but saltmarsh is not considered a particularly effective channel and bank stabilizer (Lee and Partridge, 1983; French, 1993; Allen, 1994; D’Alpaos et al., 2006; Bouma et al., 2007; Mudd et al., 2010). If the hydroperiod, the time that marshes are submerged every day, gets longer the sediment supply to the marsh increases and therefore so does the sediment accretion. Several authors therefore found that marshes are most productive at a certain rate of sea level rise (SLR), because this keeps the hydroperiod more or less constant as SLR balances with accretion rates (Redfield, 1972; Orson et al., 1985). However, marshes may drown when sea level rise rate is too large relative to the sediment supply, which leads to vegetation loss and therefore marsh drowning at an enhanced rate (Kirwan and Temmerman, 2009). In general, tidal marshes are thought to approach an equilibrium level relative to the sea level whether rising or not (Friedrichs and Perry, 2001; Marani et al., 2013).

For saltmarsh to accrete, the supply of mud is essential as the source of inorganic accumulation. This mud may have a coastal or fluvial source, pointing at the importance of the boundary conditions (de Haas et al., 2017). Although mud is transported in suspension and thus reaches higher, low-energetic elevations and areas more distal from the main channel, it is not unlimited. The suspended sediment concentration quickly decreases with distance from the channels into the marsh (Townend et al., 2011). Nevertheless, cohesive mud is more difficult to erode than sand, so that on the estuary scale mud leads to narrower systems with reduced bar dynamics through mudflat accumulation (Braat et al., 2017). The logical hypothesis is that the added effect of vegetation leads to even more accretion at the flanks of the estuary (Brew and Williams, 2010).

The availability of mud is partly determined by the changing hydrodynamic energy along the river continuum, especially in shallow, well-mixed estuaries that we focus on (Fig. 1) (Dalrymple et al., 1992). A central zone of lower energy where the average grain size decreases has been observed where bedload converges (Johnson, 1982). Dalrymple et al. (1992) suggested that this area of bedload convergence often coincides with the relative largest marsh extent (Fig. 1). Furthermore, in many estuaries a turbidity maximum zone (TMZ) occurs in the same mixed energy zone of the estuary, which is characterized by elevated suspended sediment concentrations (e.g. Brenon and Le Hir, 1999). In other words, the fluvial-tidal transition appears to be a zone of sand and mud convergence, both of which are therefore conducive to saltmarsh establishment (Fig. 1). In turn, saltmarsh may enhance the accretion as described above.

2 Methods

To investigate whether the transition of dominantly fluvial energy to dominantly tidal energy is indeed the hotspot of sedimentation and tidal marsh formation, we combine a vegetation model with the morphological estuary model built in Delft3D by Braat et al. (2017) that includes cohesive sediment. Saltmarsh modelling will be based on the recently developed riparian
vegetation model by Oorschot et al. (2015). This model takes the vegetation cycle into account, which includes colonization, growth and mortality due to flooding, uprooting, scour, and high flow velocity. We modelled the combined effects of mud and vegetation to investigate feedback mechanisms between these two and compare the model results with measurements in nine natural systems.

The model consists of two interacting codes: the hydromorphological modelling package Delft3D version 4.01.00 and our Matlab-based vegetation module. To investigate the combined effects of mud and vegetation, an existing model schematisation was used that is loosely based on the Dyfi estuary in Wales (Braat et al., 2017). The large computation times of the interacting codes necessitated our model start from their well-developed morphology after 1000 years. To isolate the effect of vegetation in the simplest possible settings, we ignore salinity, waves, and tidal components other than M2. The marsh vegetation is represented by the settling, growth and mortality traits of *Spartina anglica* and the hydraulic resistance as a function of stem dimensions and density as detailed later. It could be argued that the pioneers arriving first are other species such as *Salicornia*, but the vegetation modelling here is simplified, given the large spatiotemporal scales and first application of a vegetation model. In our runs, the vegetation traits based on the commonly occurring *Spartina anglica* are to be seen as a generic saltmarsh plant species.

2.1 Hydromorphodynamic model

Delft3D is a widely tested, open source, model that can calculate both sand and mud transport. The 2DH (depth-averaged) version was used with a parameterisation for bend flow-effects on the direction of sediment transport.

The model is mainly based on two hydrodynamic equations, the first being the conservation of mass equation:

\[
\frac{\partial h}{\partial t} + \frac{\partial hu}{\partial x} + \frac{\partial hv}{\partial y} = 0
\] (1)

where \(h\) is the waterdepth, \(t\) is time, \(u\) is the flow velocity in the x-direction and \(v\) is the flow velocity in the y-direction. Equation 1 states that any change in water depth follows from a gradient in \(q_x\) in the x-direction or a gradient in \(q_y\) in the y-direction, for a 2-D model. Momentum conservation is calculated as:

\[
\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + v \frac{\partial u}{\partial y} + g \frac{\partial z_w}{\partial x} + \frac{guv}{C^2h} \frac{u^2 + v^2}{C^2h} - V \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right) + F_x = 0
\] (2)

\[
\frac{\partial v}{\partial t} + u \frac{\partial v}{\partial x} + v \frac{\partial v}{\partial y} + g \frac{\partial z_w}{\partial y} + \frac{guv}{C^2h} \frac{u^2 + v^2}{C^2h} - V \left( \frac{\partial^2 v}{\partial x^2} + \frac{\partial^2 v}{\partial y^2} \right) + F_y = 0
\] (3)

where \(z_w\) is the water surface height, \(C\) is the Chezy roughness (\(\sqrt{m/s}\)), which will be calculated by the vegetation model described below, \(V\) is the horizontal eddy viscosity and \(F_{x,y}\) is the streamline curvature-driven acceleration term (Schuurman et al., 2013). These two equations describe the velocity variations in the x-y plane in one grid cell over time under influence of advection, eddy diffusivity, friction, changing water depth and streamline curvature. Our grid was rectangular.
Sediment transport is calculated by separate equations for the different sediment constituents. Sand transport in case of a non-cohesive bed is calculated with the Engelund-Hansen sediment transport predictor:

$$ S = 0.05 \sqrt{u^2 + v^2} \sqrt{\frac{gC^3 \rho_s - \rho_w}{\rho_w D_{50}}} $$ \tag{4} 

where $\rho_s$ the sediment density, $\rho_w$ the water density and $D_{50}$ the median grainsize. The sediment transport of the mud fraction of the model is calculated by Partheniades-Krone equations (Partheniades, 1965) for erosion flux $E_m$:

$$ E_m = M_m \left( \frac{\tau_{cw}}{\tau_{cr,e}} - 1 \right) $$ \tag{5} 

and for deposition flux $D_m$:

$$ D_m = w_s c_b \left( 1 - \frac{\tau_{cw}}{\tau_{cr,d}} \right) $$ \tag{6} 

for $\tau_{cw} > \tau_{cr,e}$, where $\tau_{cw}$ is the maximum bed shear stress due to currents, $\tau_{cr,e}$ is the critical erosion shear stress, $M_m$ is an erosion parameter, $w_s$ is the mud settling velocity and $c_b$ the average sediment concentration in the near bottom layer. Above a critical mud content threshold ($p_m > p_{m,cr}$) the sand and mud flux are proportional to their respective fractions in the sediment bed. Mud erosion is the same in the cohesive and non-cohesive regime, but the sand erosion becomes dependent on the mud entrainment in the cohesive regime, when the mud content in the bed exceeds 40%. The transport of sand becomes fully dependent on the mud flux, as bedload transport is assumed to be zero in the cohesive regime. Once sediment is suspended following the Partheniades-Krone equation it is transported by the advection-diffusion equations. A constant mud settling velocity was assumed.

A parameterization is needed for helical flow due to streamline curvature in a depth-averaged simulation to create point bars in river bends and estuarine bars, and is included as follows. The bedload transport direction $\phi_t$ is given by the following equation:

$$ \tan(\phi_t) = \frac{v - \alpha_I \frac{u}{U} I_s}{u - \alpha_I \frac{u}{U} I_s} $$ \tag{7} 

where $U$ is the depth averaged flow velocity, $I_s$ is the spiral flow intensity factor, here taken at unity, and $\alpha_I$ is given by the following equation:

$$ \alpha_I = \frac{2}{\kappa^2} \left( 1 - \frac{1}{2} \sqrt{\frac{g}{\kappa C}} \right) $$ \tag{8} 

Lastly, bed slope effects are included in the model to simulate a deviation in sediment transport direction from the shear stress direction due to grains moving downslope. The sediment transport in the x and y direction under influence of the bed slope effect is given by:

$$ q_x = q_s \left( \cos(\phi_t) - \frac{1}{f(\theta)} \frac{\partial z_b}{\partial x} \right) $$ \tag{9} 

$$ q_y = q_s \left( \sin(\phi_t) - \frac{1}{f(\theta)} \frac{\partial z_b}{\partial y} \right) $$ \tag{10}
$f(\theta)$ is given by the following equation:

$$f(\theta) = \alpha \theta^\beta$$  \hspace{1cm} (11)

In this equation $\theta$ is the shields parameter and $\alpha$ and $\beta$ are calibration parameters specified later.

### 2.2 Vegetation model

A model programmed in Matlab was used to simulate the vegetation in the estuary (Oorschot et al., 2015). This model simulates vegetation colonization, growth and mortality and translates this to hydraulic roughness used in Delft3D as based on the Baptist et al. (2007) equation:

$$C = \frac{1}{\sqrt{C_b^2 + C_d^2 n h_v^2}} + \sqrt{\frac{g}{\kappa}} \ln \frac{h}{h_v}$$  \hspace{1cm} (12)

where $C$ is the Chezy roughness value due to the bed and vegetation roughness ($\sqrt{\text{m/s}}$), $C_b$ is the Chezy value for the bed without vegetation, $C_d$ is the drag coefficient, $n$ is the number of stems per square meter times the stem diameter, $h_v$ is the vegetation height and $\kappa = 0.41$ is the Von Karman constant. Vegetation of different ages and therefore with different characteristics can occur simultaneously in one grid cell up to a total fraction of 1. The Chezy value is calculated for each age class and afterwards a total Chezy coefficient is calculated based on the fraction coverage of each age class.

The vegetation model divides the morphological year in 24 ecological timesteps, which correspond with half a month of morphological development (Table 1). Following each ecological timestep the hydromorphodynamic calculations are stopped and the bed level changes, water levels and flow velocities are exported from Delft3D to the vegetation model. A two week interval, during which vegetation properties are assumed constant, was chosen to capture the dominant vegetation development processes. The vegetation has both general and life-stage specific characteristics (Table 2 and 3). General characteristics are the seedling dimensions, i.e. shoot length and diameter and root length, maximum age, growth factors for logarithmic shoot, root and diameter development, and seed dispersal timing (Oorschot et al., 2015). Life-stage specific characteristics are rules for mortality due to flooding and uprooting, number of stems per area, drag coefficient and fraction of the grid cell surface covered with vegetation. All the variables in the Baptist et al. (2007) equation are thus accounted for. The new vegetation characteristics are then used to update the Chezy roughness field in Delft3D.

Colonization takes place during the month of seed dispersal on every location where water has been present. This means that all cells in the intertidal zone are colonized with *Spartina anglica* by the predefined colonization density. Given that the tides in the model are simplified to M2, the supratidal zone where vegetation settles in nature can be seen as included as high intertidal. There is no seed dispersal module other than that we assume the seeds to spread through the water (hydrochorously) and neither do seeds end up above the water surface. This means that seedlings colonize lower intertidal areas after which mortality determines which plants survive such that the lower intertidal zone is not occupied by plants during the flow modelling. We do not model rhizomal growth since this is a process occurring at a much smaller spatial scale than the grid cell size.

The vegetation follows a logarithmic growth function dependent on age, which limits their growth once they mature:

$$G = F_v \log(a)$$  \hspace{1cm} (13)
in which $G$ is the length or diameter of the shoot or root, $F_v$ is a characteristic growth factor for the root or shoot, and $a$ is the vegetation age in years. The initial dimensions of the seedlings are defined in the general characteristics, after which plant growth is calculated yearly following the equation.

Mortality is calculated yearly as a function of burial, uprooting, maximum flow velocities, flooding and ageing. Burial and uprooting are determined by comparison of the plant dimensions and bed level change. If the erosion in an ecological timestep exceeds the length of the root, the plant is uprooted, and if the sedimentation exceeds the shoot length it is considered buried, both leading to mortality (Oorschot et al., 2015). The calculation of mortality due to flooding and flow velocity is slightly more complex: every timestep contains twelve hours of hydrology, which is approximately the M2 tide. A morphological scale factor of 30 is used, which implies that the morphological development is 30 times faster than expected based on the hydrodynamics. Therefore, one M2 tide is used for two weeks of morphodynamics. For each cell the maximum, minimum and average water depth are determined during the tidal cycle. Because marsh vegetation starts to occur above mean tide, and usually quickly accretes to the high tide mark, the subsequent days that the cells are flooded during mean tide are recorded. For flow velocity the maximum value during the tidal cycle in each cell is stored. Lastly, vegetation dies when its maximum age is reached. A dose-effect relation (Oorschot et al., 2015) is applied to model gradual plant demise as the fraction of plants that do not survive the hydrodynamic pressure. Until a threshold is exceeded no mortality occurs, while above this threshold an increasing portion of the plants start dying with increasing stress. The threshold value and the slope of the stress-mortality relation are user-defined and can vary between the life-stages of the plants. Mortality was applied to each age class in all grid cells (Oorschot et al., 2015).

2.3 Model setup

We set up four model scenarios based on our earlier work and about 30 preliminary test runs, where we balanced time efficiency and the processes that could be realistically represented.

The initial bathymetry is the final outcome of a model run that started from an idealised convergent shape (Braat et al., 2017). This avoids long computational time to develop sufficient bars and mud flats where vegetation can settle. The rectangular cell size varies from 50 by 80 in the estuary to 125 by 230 offshore. This is done to balance computational time and sufficient spatial resolution. A 0.2 minute timestep was used based on the Courant criterion. We applied a 1.5 m tidal amplitude defined by two harmonic water levels at the north and south coastal boundaries and a constant 100 $m^3/s$ discharge at the upstream river boundary. The bed is initially entirely composed of sand and has a sand supply equal to the transport capacity at the river boundary, which avoids sedimentation or erosion at the upstream boundary. Mud, on the other hand, is supplied as a constant concentration at the upstream boundary of 20 mg/l, the same as in the run by Braat et al. (2017) that led to large-scale equilibrium of the estuary planform. This model was run for 1000 years without vegetation in Braat et al. (2017) and the final bathymetry was used as the initial condition for further simulations including vegetation (Figure 2 B). Note that this bathymetry was the result of calculations including mud, while we initially apply it as a pure sand bed in order to isolate the effect of the addition of vegetation and mud through the upstream supply.
2.4 Parameters and scenarios

Several parameters for hydromorphodynamic processes, numerical processes and vegetation development were varied (Table 1) to study their effect on estuary developments. Model scenarios were run for a 100 years, which is about the minimum time required for morphological changes at the system scale to occur due to vegetation and the practical maximum time given computational and i/o costs of about two months on a single node in a fast desktop computer (Table 1). A small morphological scale factor of 30 was used, since preliminary testing showed that this allowed vegetation settlement, growth and mortality over a number of tidal cycles without significant morphological change. In contrast, for sandy estuaries without vegetation values up to 1000 have been used (Van der Wegen and Roelvink, 2008). In the vegetation model a balance is required between morphological and hydrological timescales, since these both affect the development of the plants. If the morphology changes significantly faster than the hydrodynamics, plants are subject to large scale burial and uprooting. A default Chezy value of 50 for bare sediment was chosen as in Braat et al. (2017). Vegetation traits of Spartina anglica were based on Nehring and Adersen (2006) and Deng et al. (2009) (Table 2,3).

2.5 Data collection of natural systems

For a first quantitative comparison of model results with natural systems, we mapped along-channel variability of unvegetated channel width and width of the vegetated zone in nine natural estuaries. The natural systems were selected from the dataset of Leuven et al. (2017) based on the presence of saltmarsh vegetation, and include one system with mangrove species (Table 4).
Table 1. The main hydromorphological parameter settings.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>value</th>
<th>unit</th>
<th>motivation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Timespan model run</td>
<td>100</td>
<td>year</td>
<td>sufficient time to have changes on estuary scale</td>
</tr>
<tr>
<td>Hydrodynamic timestep</td>
<td>0.2</td>
<td>minutes</td>
<td>to fulfill courant number criteria</td>
</tr>
<tr>
<td>Morphological spin up time</td>
<td>24</td>
<td>hours</td>
<td>two tidal cycles</td>
</tr>
<tr>
<td>Drying flooding depth</td>
<td>0.08</td>
<td>m</td>
<td>balance between capturing morphodynamics and time efficiency</td>
</tr>
<tr>
<td>Morphological acceleration factor</td>
<td>30</td>
<td>-</td>
<td>low value to allow vegetation processes</td>
</tr>
<tr>
<td>Active bed layer thickness</td>
<td>0.1</td>
<td>m</td>
<td>Braat et al. (2017)</td>
</tr>
<tr>
<td>Transverse bedslope parameter $\alpha$</td>
<td>0.2</td>
<td>-</td>
<td>Braat et al. (2017)</td>
</tr>
<tr>
<td>Transverse bedslope parameter $\beta$</td>
<td>0.5</td>
<td>-</td>
<td>Braat et al. (2017)</td>
</tr>
<tr>
<td>Vegetation timestep</td>
<td>21900</td>
<td>min</td>
<td>to capture settling, growth and mortality</td>
</tr>
</tbody>
</table>

Table 2. Parametrization of general characteristics of *Spartina anglica*.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Value</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vegetation type</td>
<td>-</td>
<td><em>Spartina anglica</em></td>
<td>Common European marsh species</td>
</tr>
<tr>
<td>Maximum age</td>
<td>yr</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>Initial root length</td>
<td>m</td>
<td>0.02</td>
<td>Based on <em>S. alterniflora</em> (Deng et al., 2009)</td>
</tr>
<tr>
<td>Initial shoot length</td>
<td>m</td>
<td>0.07</td>
<td></td>
</tr>
<tr>
<td>Initial stem diameter</td>
<td>m</td>
<td>0.001</td>
<td></td>
</tr>
<tr>
<td>Logarithmic growth factor root</td>
<td>-</td>
<td>0.19</td>
<td>Based on <em>S. alterniflora</em> (Deng et al., 2009)</td>
</tr>
<tr>
<td>Logarithmic grow factor shoot</td>
<td>-</td>
<td>1</td>
<td>Nehring and Adsersen (2006)</td>
</tr>
<tr>
<td>Logarithmic growth factor stem diameter</td>
<td>-</td>
<td>0.005</td>
<td>Nehring and Adsersen (2006)</td>
</tr>
<tr>
<td>Timing of seed dispersal</td>
<td>Month</td>
<td>April</td>
<td></td>
</tr>
</tbody>
</table>

The area of each estuary was visually classified as either unvegetated or vegetated in Google Earth. Here polygons were drawn around the unvegetated part of the estuary (as described in Leuven et al., 2017), and the dataset was extended with polygons of the vegetated area (Fig. 1). The vegetated area comprises the area that borders the active estuary and is covered with pioneering or fully-grown saltmarsh vegetation. The presence of sinuous tidal creeks and vegetation other than, for instance, forest, were used as an indicator for present-day or recent tidal influence and exclude older riparian vegetation. Tidal vegetation was distinguished by its different color compared to surrounding forests and grass fields and by its clumpy and patchy structure. The elevation data in Google Earth were used as further evidence for the outer boundary of the tidal vegetation area to avoid steep gradients and cliffs at the transition from supratidal elevation level to higher elevated areas bordering the estuary.
Table 3. Parametrization of life stage specific characteristics of *Spartina anglica*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Ls 1</th>
<th>Ls2</th>
<th>Ls3</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Numbers of years in life stage</td>
<td>yr</td>
<td>1</td>
<td>10</td>
<td>9</td>
<td>...</td>
</tr>
<tr>
<td>Number of stems</td>
<td>stems/m²</td>
<td>13.000</td>
<td>1500</td>
<td>600</td>
<td>Nehring and Adersen (2006)</td>
</tr>
<tr>
<td>Area fraction (0-1)</td>
<td>-</td>
<td>0.05</td>
<td>0.5</td>
<td>0.8</td>
<td></td>
</tr>
<tr>
<td>Drag coefficient</td>
<td>-</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>cylindrical stems</td>
</tr>
<tr>
<td>Desiccation threshold</td>
<td>days</td>
<td>360</td>
<td>360</td>
<td>360</td>
<td>No desiccation assumed</td>
</tr>
<tr>
<td>Desiccation slope</td>
<td>-</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Flooding threshold</td>
<td>days</td>
<td>20</td>
<td>40</td>
<td>40</td>
<td>No desiccation assumed</td>
</tr>
<tr>
<td>Flooding slope</td>
<td>-</td>
<td>0.75</td>
<td>0.75</td>
<td>0.75</td>
<td></td>
</tr>
<tr>
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<td>0.5</td>
<td>1</td>
<td>1</td>
<td></td>
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<tr>
<td>Flow velocity slope</td>
<td>-</td>
<td>0.75</td>
<td>0.75</td>
<td>0.75</td>
<td></td>
</tr>
</tbody>
</table>

Subsequently, centerlines of the polygons were constructed along the channel, which allowed width measurements perpendicular to this centerline (following the approach of Leuven et al., 2017). This resulted in along-channel profiles of the active channel width, summed width of vegetation and estuary width, in which the estuary width is defined as the active channel width including bars plus the summed width of vegetation. The along-channel distance from the mouth was normalized with the length of the estuary. Estuary length is defined as the length from the mouth up to the point where the estuary width is equal within a few percent to the active channel width, in our case the upstream river. By this normalization a direct comparison is possible between estuaries with different lengths and our modelled simulations.

Estimates of local tidal prism and total energy were made for each of the natural systems based on (Leuven et al., 2017). Local tidal prism was estimated by multiplying the along-channel width profile with the tidal range profile and integrating over the distance upstream of a given point. The volume added by the river was characterised by river discharge multiplied by tidal period. We then calculated a characteristic velocity by dividing the local prism $TP$ by the local active width $W_a$ and half the tidal M2 period $T_{M2}/2$. As a proxy for the total flow energy this velocity was taken to the power of three as this is also a common indicator of sediment movement (Aubrey and Speer, 1985), so that flow energy is here calculated as $2TP(W_aT_{M2})^{-3}$.  

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Figure 3. Results of the four scenarios after 100 years of simulation. (a) Morphology. Colors representing larger depths than -5 m were saturated to enhance contrast. (b) Tidal range. (c) Mean of absolute flow velocity during the tidal cycle. (d) Mud thickness in cm. (e) Vegetation cover at the surface, ranging from 0–1.
Table 4. Channel area, vegetation area and estuary length derived from polygons digitised in Google Earth, accessed October 2017.

<table>
<thead>
<tr>
<th>Name</th>
<th>Location</th>
<th>Date aerial photography</th>
<th>Channel area (km²)</th>
<th>Vegetation area (km²)</th>
<th>Estuary length (km)</th>
</tr>
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<td>196.6</td>
<td>84.7</td>
</tr>
<tr>
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<tr>
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<td>3.1</td>
<td>16.0</td>
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<td>Teign estuary</td>
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<td>1/12/2011</td>
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<td>12/31/2001</td>
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<tr>
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<td>1/12/2006</td>
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<td>6.5</td>
<td>10.2</td>
</tr>
<tr>
<td>Whitehaven beach</td>
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<td>1/12/2011</td>
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<td>3.4</td>
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</tbody>
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3 Results

In the following section we will first discuss the effects of vegetation and mud on the entire estuary in terms of hydrodynamics, vegetation development, mud cover development and biomorphological development. After that we focus analyses on the central, mixed energy, zone of the estuary, which is potentially the bed load convergence zone with the largest effects of the vegetation and mud.

3.1 Effects of mud and vegetation on the entire estuary

The mouth of the estuary has a 3 m tidal range, which decreases gradually in landward direction to disappear roughly 14 km into the estuary (Fig. 3). The flow velocity, on the other hand, increases in the outer part of the estuary because the convergence is more stronger than the friction. Further in the estuary the convergence decreases and the increase in friction begins to dominate, which results in a decreasing flood velocity. Therefore, there is an optimum in the flood flow velocity at roughly 5 km into the estuary (Fig. 4). The tidal range thus behaves as a hyposynchronous system while the current behaves as a hypersynchronous system (Fig. 4).

The simulation with vegetation only develops fringing marshes at the edges of the estuary. The marshes start from the estuary mouth up to the tidal limit, roughly 14 km upstream (Fig. 3). The relative width of the tidal marshes is fairly constant at $\approx 10\%$ of the estuary width in the outer zone. Between roughly 6 km and 11 km, however, the relative width of the marshes suddenly increases. The relative width of the tidal marshes can go up to $60\%$ of the estuary width. This area coincides with the area where the flood velocity and river velocity start to decrease due to friction and estuary shape respectively (Fig. 4). Beyond 14 km there is no vegetation anymore, this is because this is beyond the tidal limit and therefore there is no drying and flooding area where seeds are distributed and seedlings survive.

The simulation with mud only results in a fairly continuous mud cover along the entire estuary (Fig. 3). There are small amounts of mud which deposit on tidal bars, in the order of an accumulated 10 cm admixed in sand over 100 years, but the
more pronounced accumulations occur on the edges of the system. Similar to the simulation with vegetation the relative mud abundance starts to increase landward of the maximum flood velocity, which occurs at roughly 6 km. The relatively large mud extent in the central zone of the estuary is due to the low flow velocities in this zone (Fig. 3,4). Unlike the vegetation cover, however, the relative mud abundance does not decrease to zero at the tidal limit, but approaches a roughly constant value of approximately 30% of the system width (Fig. 4). This is because the system is very small in this area, as the river is only several cells wide, and not because there are large extensive mudflats.

In the simulation without mud and vegetation, i.e. the reference scenario, channels and shoals are moving, but no system scale changes occur as the initial system seems to be close to dynamic equilibrium. During this simulation a slight change in hypsometry occurs. The roughly medium heights are slightly eroded, while the higher parts are slightly sedimentated (Fig. 5).

The morphology in the simulation with vegetation but without mud shows little differences compared to the reference simu-
Figure 5. Hypsometry of the entire estuary after 100 years. Dashed lines indicate the tidal range at the seaward boundary. Around 70% of the estuary area is intertidal in all scenarios, indicating that the model represents a shallow system. The hypsometry is determined over the surface occupied by the estuary of the initial condition, which excludes new areas formed by bank erosion that is modelled rather simplistically in Delft3D.

This indicates that the vegetation is unable to enhance sedimentation in absence of suspended sediment, and that it predominantly colonizes locations which are not prone to erosion because there is no significant reduction of the erosion of the intertidal area (Fig. 5). When mud is supplied to the simulations, intermediate hypsometric heights show a slight aggradation (Fig. 5). Addition of vegetation to the simulation with mud further enhances the aggradation of the upper hypsometric heights, and thus the intertidal area.

Furthermore, there is a positive feedback between mud and vegetation. Not only do mud and vegetation occur in the same area, their relative abundance also increases compared to simulations where one of them is absent (Fig. 3,4). This is emphasized by the total mud and vegetation cover in the estuary, which are almost identical after 100 years (Fig. 7a). There is an especially strong feedback in the beginning of the simulation when vegetation cover increases strongly after which mud cover starts to increase faster (Fig. 7a).
Figure 6. Estuary width over time for of the entire system and for zones along the estuary. Width is normalised by average initial width. See Fig. 2 for locations of zones.

3.2 Effects of mud and vegetation in the mixed energy zone

Vegetation presence affects the location and thickness of mud deposits mainly in the central estuary (Fig. 7b) and to a lesser degree in the outer area (Fig. 8). The vegetation cover develops faster than the mud cover, but afterwards stimulates the mud sedimentation which reaches a higher final area (Fig. 7). A major difference in hypsometry is, however, that the outer estuary has a concave profile while the central and river reach have a convex profile. This has direct consequences for the available area for vegetation. Because the effect of vegetation is largest in the central part of the estuary, a series of close-up images is provided (Fig. 9). The bathymetry of the reference simulation shows limited changes (Fig. 9 a). Vegetation colonizes the edges of the area in the simulation without mud, but remains distal from the main ebb-channel and the bathymetry develops similar to that of the reference simulation (Fig. 9 c). Larger differences occur in simulations where mud is present. When mud is
Figure 7. Interaction of mud and vegetation. (a) The development of the total mud and vegetation cover over time in the simulation where both are present, where the simulation begins in the origin of the plot. Black line indicates equality of mud and vegetation cover. (b) The average mud cover in vegetated cells and in the entire model, showing substantially higher cover in vegetated cells.

When vegetation is added to the simulation it first focuses the main ebb-channel, but afterwards the entire area starts to gradually fill and becomes shallower (Fig. 9 b).

When vegetation is added to the simulation with mud the infill of the deeper parts of the estuary is stopped. Instead the vegetation captures mud in the intertidal area and the vegetation expands laterally towards the main channel while focusing the flow (Fig. 9 d). Vegetation traps the mud in the higher intertidal areas and through this redistribution decreases the siltation of the deeper parts of the estuary. Simultaneously the accumulation of mud increases the bed level in the central part of the estuary, which enables the vegetation to laterally expand in the direction of the channel. Because mud enables vegetation to expand laterally and because mud accumulation increases within vegetated areas, the total mud and vegetation cover increases when both are present. Also the vegetation causes the deposition of mud on bars in the middle of the estuary (Fig 9d) where mud barely occurs when vegetation is absent (Fig 9c). In other words, the combined effect of vegetation and mud in the central estuary is to raise the intertidal areas and deepen the subtidal areas relative to the run with mud alone, but the overall depth compared to the control run and vegetation run is reduced. This means that the vegetation acts to focus flow into the channels, but the dominant effect is the filling of intertidal area that reduces the overall tidal prism over time.
Figure 8. Development of hypsometry of three zones in the modelled estuaries. The outer estuary has a concave shape while the central and river area have a convex shape. The middle part shows significant deposition compared to the outer estuary in simulations with mud and vegetation. Blue lines indicate initial minimum and maximum water surface elevation.

The water elevation and mean flow velocity in the middle of the estuary were plotted over time to test the hypothesis that the system becomes flood dominant when vegetation (and mud) are present (Fig. 10). The system is ebb dominant from the start. The tidal asymmetry does not change much over time for the four scenarios, but the tidal range decreases for the scenario with mud and vegetation and both simulations with vegetation cause a decreased average flow velocity (Fig. 10 b). Furthermore, the effect of combined vegetation and mud is disproportionally larger than that of vegetation or mud alone, confirming the idea of interaction. Moreover, the effect of reduction of tidal prism that determines overall flow energy dominates over the effect of reduction of intertidal area that determines the tendency of flood-dominance.

3.3 Natural systems

In the model simulations, we found that the relative vegetation abundance increases in the mixed energy zone of the estuary. This is in close agreement with observations in nine natural systems (Table 4). In natural systems, vegetation increases in
Figure 9. The development of the central zone of the estuary. (a) Simulation without mud and vegetation. (b) Simulation with only vegetation. (c) Simulation with only mud. (d) Simulation with both mud and vegetation. The mud maps belong to the simulation above it.

abundance from the estuary mouth towards a short distance before the tidal limit, while landward of the tidal limit the vegetation cover decreases quickly towards zero (Fig. 11). Similar to the modelled scenarios, the landward vegetation cover increase coincides with the decrease of the flow energy. The upper limit of the vegetation is slightly beyond the tidal limit, but this is probably because we included old marshes, which are rarely flooded.

5 Discussion

In the discussion first the location of tidal marshes is assessed, second their effect on morphology is investigated, thirdly we look into their effect on the tidal wave, then we compare our model outcome with natural systems and last, the implications for further research are given.

Modelled marshes reach their largest extent in the central part of the estuary, where the tidal energy is the lowest in agreement with the qualitative model of Dalrymple et al. (1992). The marsh expands mostly landward from the maximum flood current velocity. This is also where the bedload convergence zone begins, and in natural estuaries where a the turbidity maximum zone may occur (Fig. 12). The main reason for the increase in marsh extent is the combination of flow velocities being low...
The final tidal cycle in the central estuary at 6 km from the mouth, showing the strongest reduction for the scenario with combined mud and vegetation. (a) Tidal water level. (b) Width-averaged flow velocities over the cycle.

enough, with the presence of suitable bed elevations. The establishment of tidal marshes requires a window of opportunity with long enough mild hydrodynamic stress (Bouma et al., 2014). However, the modelled marshes develop primarily landward and not seaward of the maximum flood velocity, which shows that the hydrodynamics are not the only limiting factor. In reality, however, the hydrodynamic stresses will be larger in the outer part as well as wave magnitude is more significant there (Dalrymple et al., 1992) and waves are a major limiting factor for seedling establishment in marsh and mangrove landscapes (Balke et al., 2013). Waves would result in a further reduction in tidal marsh extent in the outer estuary but will have limited effect on the central part of the estuary and therefore strengthen the trends in our model.

The importance of sediment accumulation in the central part for marsh development is shown in the scenario with mud and vegetation. This simulation shows a further extent of the marshes because mud preferably accumulates in the central part of the estuary, regardless of the fact that no preferential establishment of vegetation on a muddy substrate is included in the model. While it is known that suspended sediment is a requirement for tidal marshes to keep up with sea level rise (D’Alpaos et al.,
Figure 11. (a) The total, active and marsh width along three natural estuaries, partitioned by the method of Leuven et al. (2017). (b) The vegetated part as a percentage of the total width. (c) Tidal prism, discharge and energy taken as width-averaged tidal prism (see for method Leuven et al., 2017).
Figure 12. Comparison of mud flats and tidal marsh vegetation in a modelled and natural system. Here, velocity magnitude to the power 3 is plotted as an indication for hydrodynamic energy.
Figure 13. Relative vegetated width along the estuary averaged for nine natural estuaries compared to the simulation with mud and vegetation. Distance along the estuary is normalised by the approximate distance between coastline and tidal limit.

2006, 2007; Murray et al., 2008; Fagherazzi et al., 2012), the present model results show that suspended sediment is also a requirement for significant lateral marsh progradation into the estuary. We show that the presence of vegetation increases the mud deposition in the upper intertidal area in agreement with observations (Larsen et al., 2007; Zong and Nepf, 2011; Follett and Nepf, 2012), but also that this reduces accumulation in the lower intertidal area. Once the vegetation starts to expand and approaches the main channel (Fig. 9) it starts to focus and concentrate the flow (Fig. 3). After vegetation settlement and stabilization, vegetation causes flow focussing, similar to the fluvial environment (Tal and Paola, 2007; Dijk et al., 2013).

Despite the reduction of intertidal flood storage, the central zone barely becomes more flood dominant and the tidal limit shifts seaward. This is in contrast to expected tidal dynamics (Friedrichs, 2010), probably because the river in this part of the estuary already dominates over the tidal influence. The seaward shift of the tidal limit implies that the inundation time, and therefore stress, of the marshes decreases, explaining why vegetation density increases in the central estuary. Regardless, the river flow, if large enough to move sediment, will keep a channel open even if the floodplains fill up, such that an equilibrium tidal river may develop. This amounts to progradational filling of the estuary as observed in the Holocene (de Haas et al., 2017).

The general agreement between trends in the natural systems and the numerical model indicates that the overall pattern of tidal marsh and mud flats along the estuary is determined mainly by the tidal hydromorphodynamics and the interaction with mud and vegetation. Figure 13 shows the mean relative vegetation abundance for nine alluvial systems along the fluvial-tidal transition with pronounced marshes. The relative extent of the vegetation can be higher in natural systems, which has three main causes. First, the modelled system started as a narrow convergent estuary while many natural systems start from unfilled basins. Second, natural systems are to a much larger degree infilling than our ebb-dominant system with little sediment import from the sea and they had a much longer time to fill gradually. Third, many natural estuaries develop pronounced turbidity
maximum zones (TMZ) under effect of density driven currents, tidal currents and river currents. Such a TMZ would develop roughly at the mixed energy zone, and a pronounced TMZ can be hypothesized to enhance marsh expansion and accretion of the central part of the estuary that already occurs without a turbidity maximum zone (Braat et al., 2017).

Present limitations of our model study leave open the question what the effects would be of sediment supply coming from the sea on marsh establishment in the outer estuary. Further refinement could also include a longer duration of the simulation and the inclusion of multiple tidal components, which may reduce the ebb dominance. Regardless, the fundamental feedback mechanism between mud and vegetation would still affect the larger scale estuary development as mud facilitates the expansion and survival of marshes while vegetation facilitates the capture of mud.

5 Conclusions

Numerical modelling of estuaries shows that vegetation follows mud accumulation patterns and simultaneously enhances mud accumulation rates. A positive feedback mechanism emerged in the model between the mud sedimentation and vegetation settlement. Mud sedimentation leads to higher elevated intertidal areas suitable for vegetation settling and development. The vegetation then increases local flow resistance which enhances sedimentation of mud that would otherwise be resuspended again.

Through this biomorphological feedback loop vegetation has a strong effect on morphodynamics in the middle estuary while its effect in the outer estuary is marginal due to larger flow energy. The relative extent of marsh vegetation increases from the outer estuary towards the inner estuary and can increase from 10% to 50% of the estuary width or probably even more, which is in agreement with observations in natural systems. In particular, the feedback enhances the sedimentary trend in what has been recognised in the literature as the Bedload Convergence Zone in the mixed-energy fluvial-tidal transition. The main effect of the overall intertidal space filling is to reduce the tidal prism and progressively fill the estuary in agreement with observations of Holocene systems. The focussing of flow between flanking marsh vegetation has only a limited effect on channel depth, in contrast to observed effects in saltmarsh channels and rivers. The reduction of flood storage has a negligible effect on the flood dominance of the estuary, in contrast to idealised modelling results in the literature. These results are, however, mainly valid for shallow sandy estuaries.

The effect of vegetation alone on the hypsometry of the entire estuary is limited. This is mainly because its effect on the outer estuary is marginal, where it occupies only a small portion of the estuary surface. In the central part of the estuary vegetation occupies a much larger fraction of the width so that its effects are most pronounced here. When mud is present and forms new intertidal area, the vegetation expands towards the channel, which drives further accretion and forces the system into a single main channel. When mud is absent vegetation lacks an accreting effect because the sand does not reach the vegetated areas for lack of energy in the shallowest flows.
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Competing interests. The authors declare that they have no competing financial interests.

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References


