UNRAVELING THE COUPLED LARGE-SCALE SUSPENDED SEDIMENT AND PHYTOPLANKTON DYNAMICS IN A TURBID AND TIDE-DOMINATED ESTUARY
Unraveling the coupled large-scale suspended sediment and phytoplankton dynamics in a turbid and tide-dominated estuary

Het ontraadselen van de gekoppelde grootschalige dynamiek van zwevend sediment en fytoplankton in een troebel en getijde-gedomineerd estuarium

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Estuaries often show regions in which suspended particulate matter (SPM) and/or phytoplankton accumulate. Predicting the location of these regions and the corresponding magnitude of the SPM and phytoplankton concentration is of great importance for managing the estuary; it may prevent the system from evolving towards a (hyper-)turbid state, a condition in which phytoplankton growth is very limited. Such an evolution may have drastic consequences for the ecological state of the estuary as phytoplankton forms the basis of the food chain.

To predict the location and magnitude corresponding to accumulation of SPM and phytoplankton, we have to understand the interconnection between SPM and phytoplankton concentration. On the one hand, SPM is often a limiting factor for estuarine phytoplankton growth by deteriorating the light climate and thereby limiting photosynthesis. On the other hand, various authors showed that phytoplankton may determine the SPM concentration by, for example, the excretion of sticky substances. These substances may impact flocculation and thereby the settling velocity and dynamics of SPM flocs. Additionally, they may stabilize the bed and thus change the erosion properties, which also affects the SPM concentration by adjusting the sediment flux from the bed. While most literature focuses on the small-scale impact of biological flocculants on the formation of larger aggregates and erosion properties of the bed, its influence on the SPM profiles on the large spatial- and temporal scale is still largely unknown.

In this thesis, we study the interconnection of SPM and phytoplankton on the large temporal- and spatial scale, applied to a turbid, tide-dominated, and nutrient-rich estuary, being the Scheldt estuary. To this end, we combine a model approach and analysis of multi-annual observations covering the entire domain of the Scheldt estuary.

We first study the impact of flocculation on the formation of estuarine turbidity maxima (ETM), which are regions in which the concentration of SPM is larger than landward and seaward of these regions. To this end, an idealized, width-averaged hydro-sediment transport model is utilized and extended by a flocculation model. The model is applied to the Scheldt estuary in winter conditions. We show that spatial and temporal variations in settling velocity due to flocculation may promote land-inward sediment transport and result in additional ETM. They are essential to capture the observed magnitude of the SPM concentrations and their dependence on river discharge.

Secondly, to study the importance of biotically-induced seasonality in flocculation and erosion on seasonality in ETM formation, the same model framework is applied to both a winter and summer case in the Scheldt estuary. By calibrating the model to multi-annual, in situ turbidity observations, we show that we do not require seasonality
in biotically-affected flocculation and erosion parameters to reproduce the seasonality in ETM formation. Our findings are supported by multi-annual, in situ observations in turbidity, floc size, Chlorophyll-a (cf. phytoplankton abundance), and sticky substances excreted by phytoplankton; the abiotic observations show seasonality, while seasonality in the sticky substances is absent on the estuary scale.

Thirdly, before moving towards a fully operational phytoplankton-sediment transport model, we study the impact of temporal variability in factors affecting phytoplankton growth through light availability. These factors are solar irradiance at the water surface, exponential light extinction in the water column, and water depth (cf. ratio between euphotic/mixing depth). This study is important because, within our model approach, we aim to reduce computation time as it enables an extensive sensitivity analysis. We thus want to know whether we may neglect temporal variability in these factors and, if not, how we can correct for the introduced error by excluding this temporal variability. To this end, we construct a conceptual model and apply it to the Scheldt estuary. Here, we show that our conceptual model correctly predicts the magnitude and sign of the introduced error. Moreover, we demonstrate that variability in solar irradiance has the largest impact on time-averaged primary production (PP) in dynamic equilibrium, resulting in a 30% decrease compared to time-invariant PP. Furthermore, temporal variability in solar irradiance may significantly decrease unbound exponential phytoplankton growth and may delay the onset of a phytoplankton bloom by $\sim 2$ weeks.

Finally, we combine our previous results to construct a phytoplankton-sediment transport model. To show its applicability, we apply the model to investigate the multi-annual evolution of the estuary-scale phytoplankton distribution in spring in the Scheldt estuary. More specifically, we study what change in factors affecting phytoplankton growth may have dominantly contributed to the appearance and disappearance of a spring phytoplankton bloom in the Scheldt estuary between 2004-2018. Our results suggest that the observed change in SPM alone cannot explain the multi-annual evolution in phytoplankton blooms in spring. Instead, a change in phytoplankton mortality rate and the corresponding grazing by zooplankton and phytoplankton community characteristics may have significantly contributed to this multi-annual estuary-scale evolution.

To conclude, we combined a model approach and analysis of multi-annual observations in the Scheldt estuary to study the coupled SPM-phytoplankton dynamics on a large spatial- and temporal scale. First, we extended a hydro-sediment transport model by a flocculation model and showed that flocculation might significantly impact the estuary-scale SPM distribution. Next, we applied the model to study the impact of biotically-induced flocculation and erosion on the estuary-scale seasonality in the SPM distribution. Here, we showed that biotically-induced seasonality in flocculation and erosion only has a minor impact on the seasonality in SPM on the estuary scale; seasonality in freshwater discharge might explain the observed seasonality in SPM. Next, before constructing a fully operational phytoplankton-sediment transport model, we constructed a conceptual model to assess the relative impact of temporal variability in light climate (cf. SPM) on phytoplankton growth. We found that temporal variability in light climate may drastically reduce the time-averaged PP and exponential phytoplankton growth and delay the onset of a spring bloom by $\sim 2$ weeks. Last, we combined our previous results to show that
a multi-annual change in mortality rate, and not in the SPM alone, may explain the observed multi-annual evolution in phytoplankton blooms in spring in the Scheldt estuary. Although we applied our modeling framework to the Scheldt estuary, it can also be applied to other turbid, tide-dominated, and nutrient-rich estuaries for which our assumptions are valid.

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Samenvatting

Estuaria vertonen regelmatig regio’s waarin gesuspendeerd sediment (SPM) en fytoplankton accumuleert. Voor het beheer van het estuarium is het belangrijk om de locaties van dergelijke regio’s te voorspellen en de bijhorende magnitude van de SPM en fytoplankton concentratie; zo kan men immers voorkomen dat het systeem evolueert naar een hypertroefbele toestand, een conditie waarin fytoplankton zeer gelimiteerd is. Een dergelijke evolutie kan drastische gevolgen hebben voor de ecologische waarde van het estuarium aangezien fytoplankton de basis vormt van de voedselketen.

Om de locatie te voorspellen waar SPM en fytoplankton accumuleert en de bijhorende concentraties dient het verband tussen SPM en fytoplankton begrepen te worden. Aan de ene kant is SPM vaak limiterend voor fytoplankton groei aangezien SPM zorgt voor een verslechtering van het lichtklimaat en daarmee de fotosynthese bemoeilijkt. Aan de andere kant toonden verschillende auteurs aan dat fytoplankton de SPM-concentratie kan bepalen door bijvoorbeeld de uitscheiding van plakkerige substanties. Deze substanties hebben immers een invloed op het vlokvormingsproces en bepalen daarmee de valsnelheid en dynamiek van SPM vlokken. Daarbovenop kunnen ze de bodem stabiliseren en zo de erosie eigenschappen aanpassen, wat dan weer een impact heeft op de sediment flux afkomstig van de bodem. Terwijl de meeste literatuur focust op de impact van biologische flocculanten op de vorming van grotere vlokken en op de bodemeigenschappen op kleine schaal, is de invloed op grote spatiale en temporele schaal nog voornamelijk onbekend terrein.

In deze thesis bestuderen we het verband tussen SPM en fytoplankton op grotere temporele en spatiale schaal, toegepast op een troebel, getijde-gedomineerd en nutriëntrijk estuarium, zijnde het Schelde estuarium. Om dit te bekomen combineren we een model aanpak en analyse van meerjarlijkse observaties die het gehele domein van het Schelde estuarium omvatten.

Ten eerste bestuderen we de impact van vlokvorming op het ontstaan van estuarine turbiditeitsmaxima (ETM). Dit zijn regio’s waarin de SPM-concentraties groter zijn dan land en zeewaarts van deze regio’s. Met dit als doel gebruiken we een geïdealiseerd breedtegemiddeld hydro- en sediment transport model. Het model wordt uitgebreid en toegepast op het Schelde estuarium in winter condities. We tonen aan dat spatiale en temporele variaties in de valsnelheid ten gevolge van vlokvorming kunnen resulteren in landinwaarts sediment transport en het ontstaan van extra ETM. Bovendien zijn deze variaties essentieel om de geobserveerde magnitude van de SPM-concentraties en bijhorende afhankelijkheid van rivierafvoer te modeleren.

Ten tweede bepalen we de bijdrage van seizoenaliteit in vlokvorming en erosie met
een biotische oorsprong aan seizoenaaliteit in het vormen van ETM. Hiervoor passen we dezelfde modelaanpak toe op zowel een winter als zomer geval in het Schelde estuarium. Door het model te kalibreren aan meerjaarlijkse in situ turbiditeitsobservaties tonen we aan dat seizoenaaliteit in parameters gelinkt aan vlokvorming en erosie met een biotische oorsprong niet noodzakelijk is om de seizoenaaliteit in de ETM-dynamiek te verklaren. Onze bevindingen worden bevestigd door meerjaarlijkse in situ observaties van turbiditeit, vlokgrootte, Chlorophyll-a (cf. fytoplankton aanwezigheid) en plakkerige substanties uitgescheiden door fytoplankton; terwijl abiotische observaties een seizoenaal patroon vertonen, zien we dit niet in de observaties van de plakkerige substanties op estuariene schaal.

Ten derde bestuderen we de impact van temporele variaties in lichtklimaat op fytoplankton groei alvorens we overgaan op een uitgebreid fytoplankton en sediment transport model. Hier focussen we op variaties in lichtinstralning aan het wateroppervlak, exponentiële licht-extinctie in de waterkolom en waterdiepte (cf. ratio tussen euotische- en mengdiepte). Deze studie is belangrijk aangezien we in onze modelaanpak de rekentijd zo veel als mogelijk trachten te beperken om zo een uitgebreide gevoeligheidsanalyse mogelijk te maken. We wensen dus te bepalen of we bovengenoemde temporele variaties al dan niet mogen verwaarlozen. Als dit niet kan, wensen we te bepalen welke correcties we kunnen opleggen om de geïntroduceerde fout te beperken. Om dit te doen stellen we een conceptueel model op en passen we dit model opnieuw toe op het Schelde estuarium. We tonen aan dat ons conceptueel model de magnitude en het teken (i.e., afname of toename) van de geïntroduceerde fout juist voorspelt. Bovendien laten we zien dat variabiliteit in lichtinstralning aan het wateroppervlak de grootste impact heeft op tijds gemiddelde primaire productie (PP). Het resulteert in een vermindering van tijdgemiddelde PP van ongeveer 30 %, zorgt voor een significante vertraging van fytoplankton in exponentiële groei en kan resulteren in het uitstellen van een fytoplankton voorjaarsbloei met twee weken.

Ten slotte combineren we al onze eerdere resultaten om een uitgebreid fytoplankton-sediment transport model op te stellen. Om de toepasbaarheid van dit model aan te tonen, gebruiken we dit model om de meerjaarlijkse evolutie van de fytoplankton verdeling op estuariene schaal in het Schelde estuarium te bestuderen. Meerbepaald bekijken we welke verandering in factoren die fytoplankton groei beïnvloeden een belangrijke voetdruk zou kunnen hebben op het ontstaan en verdwijnen van een fytoplankton voorjaarsbloei in het Schelde estuarium in 2004-2018. Onze resultaten suggereren dat de geobserveerde verandering in SPM niet voldoende is om de meerjaarlijkse evolutie in voorjaarsbloei te verklaren. Daartegenover, een verandering in sterfte van fytoplankton en daaraan gelinkte begrazing door zooplankton en de soorteneigenschappen van het fytoplankton zouden wel eens een belangrijke verklaring kunnen zijn voor deze meerjaarlijkse evolutie op estuariene schaal.

Samengevat, we hebben een modelaanpak en analyse van meerjaarlijkse observaties in het Schelde estuarium gecombineerd om de gekoppelde SPM en fytoplankton dynamica te bestuderen op grote spatiale en temporale schaal. Ten eerste hebben we een hydro-sediment transport model uitgebreid met een vlokvormingsmodel. We hebben aangetoond dat vlokvorming een belangrijke impact kan hebben op de SPM-verdeling op estuariene schaal. Vervolgens hebben we het model toegepast om zo de invloed van vlokvorming
en erosie met biotische oorsprong op seizoenaliteit in de SPM-verdeling op estuariene schaal te bestuderen. Hier toonden we aan dat biotische-geïnduceerde seizoenaliteit in vlokvorming en erosie slechts een beperkte impact heeft op seizoenaliteit in SPM op estuariene schaal. Seizoenaliteit in rivierafvoer kan grotendeels de geobserveerde seizoenaliteit in de SPM-verdeling verklaren. Vervolgens hebben we een conceptueel model opgesteld om de relatieve impact van temporele variabiliteit in lichtklimaat (cf. SPM) op fytoplankton groei te bepalen. We vonden dat temporale variabiliteit in lichtininstraling aan het wateroppervlak de tijdsgemiddelde PP en fytoplankton in exponentiële groei drastisch kan verlagen. Bovendien kan het resulteren in het uitstellen van een fytoplankton bloei met twee weken. Ten slotte hebben we al onze eerdere resultaten samengevoegd om aan te tonen dat een meerjaarlijkse verandering in mortaliteit van fytoplankton, en niet enkel SPM, de geobserveerde meerjaarlijkse evolutie in de voorjaarsbloei van fytoplankton in het Schelde estuarium kan verklaren. De toepassing van onze modelaanpak is niet beperkt tot het Schelde estuarium. Het model kan ook toegepast worden op andere troebele getijde-gedomineerde en nutriëntrijke estuaria.

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Chapter 1
Introduction
1.1 Why study sediment and phytoplankton dynamics in estuaries?

The term estuary originates from the Latin adjective austuarium, meaning tidal (Fairbridge, 1980); as defined by Dionne (1963), an estuary is an inlet of the sea reaching into a river valley as far as the upper limit of tidal rise, usually being divisible into three sectors: (a) a marine or lower estuary, in free connection with the open sea; (b) a middle estuary, subject to strong salt and freshwater mixing; and (c) an upper or fluvial estuary, characterized by freshwater but subject to daily tidal action.

The specific features of estuaries make them valuable from multiple perspectives. Being the interconnection between land and sea makes estuaries often densely populated. Consequently, estuaries are necessary for flood protection as they reduce the tidal wave energy in the land-ward direction. They are also crucial for the economy through, for example, food production, water regulation, and wastewater treatment (Costanza et al., 1997). Furthermore, estuaries form unique habitats (Davidson et al., 1991) and thereby are essential to maintain biodiversity from an ecological perspective.

In many estuaries, we have two crucial indicators regarding their ecological condition: primary production of phytoplankton and the suspended particulate matter (SPM) concentration. The reason is that first of all, phytoplankton forms the base of the estuarine pelagic food web (Brett and Müller-Navarra, 1997; Sobczak et al., 2005; Falkowski, 2012). Additionally, the fact that estuaries are often turbid and SPM may significantly deteriorate the light climate in the water column (Wofsy, 1983) makes the SPM concentration an important limiting factor for phytoplankton growth (Sverdrup, 1953; Colijn, 1982; Cloern, 1987) compared to other limiting factors such as nutrient limitation and grazing.

In the past, specific anthropogenic measures had drastic consequences on both the phytoplankton and SPM concentration. For example, the Ems estuary evolved towards hyper-turbid conditions after deepening the system, prohibiting primary production of phytoplankton, the base of the estuarine food web, and even impeding navigation because of increased friction force (Winterwerp and Wang, 2013; Winterwerp et al., 2013). Another example, the notable increase in wastewater treatment capacity in the Scheldt estuary around 2006 resulted in significantly higher oxygen concentrations and intense phytoplankton blooms (Maris and Meire, 2017).

To summarize, why study sediment and phytoplankton dynamics in estuaries? Estuaries are unique systems with substantial economic value and are essential to maintain biodiversity and ensure flood protection. However, they are under continued anthropogenic pressure. To assess the impact of specific management measures on phytoplankton and SPM dynamics, often two crucial factors for a good ecological state of an estuary, insight into the fundamental functioning of the SPM and phytoplankton dynamics and their interconnection is required.
1.2 The link between SPM and phytoplankton dynamics

This section briefly introduces the complex interconnection between SPM and phytoplankton dynamics we focus on in this thesis (Fig. 1.1). To reduce complexity, we mainly study the SPM and phytoplankton dynamics on the estuary scale. Here, regions with locally elevated phytoplankton and SPM concentrations are particularly interesting.

Figure 1.1: Illustration of the interconnection between SPM and phytoplankton studied in this thesis. SPM limits phytoplankton growth by deteriorating the light climate in the water column. Phytoplankton may affect the SPM dynamics by altering the erosion properties of the bed, floc size density and structure of flocs, and by excreting sticky substances which stimulate flocculation of SPM.

Focusing on phytoplankton, such elevated concentrations result from a complex interplay between physical, transport-related processes and chemical-biotic factors that determine net local phytoplankton growth. Such processes include water temperature (Eppley, 1972), river flushing (Filardo and Dunstan, 1985) [Liu and de Swart, 2015], salinity (Lucas et al., 1998), phytoplankton grazing (Alpine and Cloern, 1992) [Lionard et al., 2005], nutrients (Tilman et al., 1982) [Cira et al., 2016], and the light climate in the water column (Sverdrup, 1953) [Desmit et al., 2005].

The light climate in the water column is determined by the SPM concentration, especially in turbid estuaries (Wofsy, 1983). To understand the phytoplankton dynamics, we thus have to study the SPM dynamics.

The SPM dynamics also result from complex interactions of various sediment transport mechanisms. Examples of such transport mechanisms are, amongst others, related to tidal asymmetries in water motion, water density gradients, and transport mechanisms related to temporal variability in settling velocity of SPM [see, for example, Burchard et al. (2018) for a review]. One mechanism resulting in temporal and spatial variability of settling velocity is flocculation. Flocculation results in the aggregation and break-up
of cohesive SPM, including both organic and mineral solids, thus changing the density, floc size, and settling velocity.

The fact that the flocculation process impacts the SPM dynamics has important consequences. Many authors showed that the presence of biota might have a significant impact on the flocculation process. Biotic characteristics that influence flocculation are, for example, sticky biotic substances excreted by phytoplankton and the organic content of SPM. The organic content directly alters the differential density and structure of flocs (Kranenburg, 1994; van Leussen, 1994; Fall et al., 2021), which affects the averaged floc size (Mietta et al., 2009), floc strength and collision efficiency (Winterwerp and van Kesteren, 2004), and floc break-up (Alldredge et al., 1990). Finally, in situ observations show a correlation between Chlorophyll-a (Chl-a), which is an indicator for phytoplankton abundance, and flocculation efficiency (Verney et al., 2009), and sticky biotic substances excreted by phytoplankton [i.e., Transparent Exopolymer Particles (TEP)] and floc strength (Fettweis et al., 2014).

Besides the flocculation process, biotic factors may also impact the erosion properties of the sediment bed, which also affects the SPM dynamics. On the one hand, various authors found that phytoplankton activity, which typically peaks in spring and summer, often stabilizes the sediment bed (Frostick and McCave, 1979) and induces the formation of bedforms (Malarkey et al., 2015) by, for example, the excretion of extracellular polymeric substances (EPS), of which TEP are a particulate form (Passow, 2002), and which bind the sediment together (Stal, 2010). On the other hand, the subsequent grazing of phytoplankton by bioturbatory macroheterotrophs may increase the erodibility (Paterson and Black, 1999).

To summarize, both the SPM and phytoplankton dynamics are complex and coupled. On the one hand, the SPM concentrations impact phytoplankton growth by deteriorating the turbidity of the water column, especially in turbid estuaries. On the other hand, phytoplankton may affect SPM dynamics by altering the erosion properties of the bed and flocculation characteristics.

### 1.3 Introduction to the Scheldt estuary

Depending on the domain of interest, estuaries can be classified based on, for example, their physiography (e.g., shallow, funnel-shaped), hydrodynamics (e.g., freshwater discharge, tidal range), climate (e.g., tropical, non-tropical), and biological condition (nutrient-rich, light-limited for phytoplankton growth) (Fairbridge, 1980).

This thesis focuses on shallow, tide-dominated systems, which are systems in which the tidal energy is large compared to the freshwater discharge. Consequently, the system can be considered to be well-mixed. This assumption allows us to approximate the vertical distribution of salinity and phytoplankton as homogeneous. Moreover, we assume that the system is nutrient-rich and light-limited, allowing us to neglect nutrient limitation. These
assumptions result in a drastic simplification of the dynamics. This is recommended given the complexity of phytoplankton and SPM dynamics, as demonstrated in the previous section.

An example of an estuary for which these assumptions are valid is the Scheldt estuary (Belgium, Netherlands). The Scheldt estuary is the system we focus on in this thesis. In each chapter, we apply our theoretical results to this estuary. However, our theoretical results are not limited to the Scheldt estuary but are also applicable to other systems with similar characteristics. In the following, we briefly introduce the Scheldt estuary by focusing on the features of interest in this thesis.

The Scheldt estuary is an approximately 160 km long, funnel-shaped estuary (Fig. 1.2). It flows through Belgium into the North Sea near Vlissingen (Netherlands). Because of its relatively small freshwater discharge, the Scheldt estuary can be considered a tide-dominated and well-mixed estuary (Baeyens et al., 1997; Meire et al., 2005). The total time-averaged freshwater discharge over the years 2015-2018 shows distinct seasonality and equals 40 m$^3$ s$^{-1}$, 174 m$^3$ s$^{-1}$, and 72 m$^3$ s$^{-1}$ in summer (Jun.-Aug.), winter (Jan.-Mar.), and spring (Apr.-May), respectively. The main tributaries of the Scheldt estuary are the Rupel and the Dender. The Upper Sea Scheldt boundary (i.e., the upstream boundary of the Scheldt estuary), the Rupel, and the Dender are responsible for 16.4, 77.2, and 6.4 % of the total river discharge in summer, 35.4, 53.1, and 11.5 % in winter, and 27.1, 63.3, and 9.6 % in spring, respectively (Waterinfo.be, 2019, deducted from gauge stations at the upstream boundary zes57a-1066, at the Dender tributary den02a-1066, and downstream from the Rupel tributary zes29f-1066 assuming conservation of mass). The dissolved nitrogen, phosphorous, and silicate concentrations in spring range from 0.1 mmol L$^{-1}$, 0.001 mmol L$^{-1}$, and 0.005 mmol L$^{-1}$ at the seaside boundary to 0.4 mmol L$^{-1}$, 0.007 mmol L$^{-1}$, and 0.13 mmol L$^{-1}$ at the upstream boundary, respectively. These concentrations are at least one order of magnitude larger than the half-saturation constants at which we expect nutrient depletion (Billen and Garnier, 1997; Lancelot et al., 2005; Arndt et al., 2011; Naithani et al., 2016).

The maintenance of the navigation channel to the port of Antwerp (first four red dots from the Dutch-Belgian border in Fig. 1.2) requires intensive dredging activities. To minimize the risk of flooding by altering the total volume of the estuary, the dredged material is often dumped back into the Scheldt estuary. This dumping comes with significantly higher SPM loads ($10^6$ ton year$^{-1}$) than the fluvial input of SPM (Dijkstra et al., 2019b). At the main dumping locations at km 73 and 78 from the mouth, the time-averaged dumped material between 2001-2015 corresponds to a high SPM input of approximately 60 and 100 kg s$^{-1}$, respectively. Consequently, these dumping activities impact the SPM concentration locally, which was confirmed by a multivariate regression analysis of SPM observations between 1996-2016. This analysis showed that the SPM concentrations at the dumping sites are dominantly correlated to dumping activities (IMDC, 2016). The yearly fluvial input of SPM at the upstream boundary and tributaries is of the order of $10^4$-$10^5$ ton year$^{-1}$ (Plancke et al., 2017; Dijkstra et al., 2019b).

The Scheldt estuary is an interesting example in view of the interplay between phytoplankton, SPM, and other factors. Firstly, the flocculation process, which links phyto-
plankton to SPM, is important in the Scheldt estuary (Peters, 1972; Gourgue et al., 2013). Moreover, Cox et al. (2019) reported a long-term estuary-scale change in SPM dynamics in the Scheldt estuary. From 2009 onwards, a change in the estuarine turbidity maximum dynamics and an overall increase in SPM concentration was observed, reaching typical SPM concentration at the water surface of $\sim 100$ mg L$^{-1}$, which may significantly limit photosynthesis (Wofsy, 1983). Simultaneously, the water quality in the Scheldt estuary improved drastically mainly because of a significant increase in wastewater treatment capacity in Brussels around 2006 (Brion et al., 2015). This resulted in increasing oxygen concentrations and changes in the zooplankton community and abundance. More specifically, calanoid copepods, in casu *Eurytemora affinis*, dominated in the downstream brackish region and were quasi absent in the freshwater region at the beginning of the observations in 1996. Next, they gradually developed more upstream to also become dominant there from 2009 (Appeltans, 2003; Mialet et al., 2010, 2011; Chambord et al., 2016). This could affect the phytoplankton abundance through grazing. These changes in the SPM dynamics and zooplankton abundance coincide with long-term changes in phytoplankton dynamics, such as the appearance and disappearance of a phytoplankton spring bloom in the brackish region. From 2004 until 2007, we observed almost no spring bloom in the brackish region. Such a spring bloom was consistently observed between 2008-2014 but disappeared after 2015.
1.4 Approach and research questions

In this thesis, we combine two approaches to study the SPM and phytoplankton dynamics in the Scheldt estuary. Firstly, we analyze long-term in situ observations in the Scheldt estuary. Secondly, we develop a model to study the interconnection between SPM and phytoplankton dynamics. To show the applicability of our model approach, we apply our model to the multi-annual observations in the Scheldt estuary. The following sections introduce the monitoring campaigns resulting in the multi-annual observations and the model approach.

1.4.1 Monitoring campaigns in the Scheldt estuary

Both the Belgian and Dutch part of the Scheldt estuary have been monitored intensively over the last two decades. In the Belgian region, various variables, including Chl-a (cf. phytoplankton), SPM, salinity, and phytoplankton characteristics, such as the maximum photosynthetic production rate and growth efficiency, have been measured within the multi-annual OMES (Dutch: “Onderzoek Milieu Effecten Signaplan”) monitoring campaign (Maris and Meire, 2017), independently of the tidal phase and spring-neap tide biweekly or monthly at 16 fixed stations (see Fig. 1.2). In the Dutch region, we use biweekly or monthly observations of Chl-a and SPM by Rijkswaterstaat at three stations in the main channel (see Fig. 1.2). For a detailed methodological description, we refer the reader to the OMES reports (Maris and Meire, 2017) and the website of Rijkswaterstaat (Rijkswaterstaat, 2020).

1.4.2 A suitable model: extending the iFlow model

The model we use is the process-based, width-averaged, idealized model known as iFlow (Dijkstra et al., 2017). The model solves the width-averaged shallow water equations in tide-dominated estuaries by neglecting the effects of Coriolis and assuming that density variability is small compared to the average density, allowing for the Boussinesq approximation. The hydrodynamics are forced at the upstream boundary and two main tributaries by a fixed water inflow and at the mouth by a tidal signal. These equations and corresponding boundary conditions result in the dynamics of the water level elevations, horizontal- and vertical water velocity. For a detailed description, we refer the reader to Chernetsky et al. (2010) and Dijkstra et al. (2017). Following Warner et al. (2005), the longitudinal salinity profile is implemented as a tide- and depth-independent profile. This assumption is consistent with the Scheldt estuary being a well-mixed estuary (Baeyens et al., 1997; Meire et al., 2005). The SPM and phytoplankton dynamics are resolved using the SPM and phytoplankton mass balance equations in equilibrium conditions. For completeness and because these dynamics form the basis of this thesis, we repeat the latter equations and corresponding boundary conditions. The width-averaged sediment mass balance equation and related boundary conditions read as (Chernetsky et al., 2010)
\[
\frac{\partial}{\partial t} c + u \frac{\partial}{\partial x} c + w \frac{\partial}{\partial z} c - \frac{1}{B} \frac{\partial}{\partial z} (BK_h \frac{\partial}{\partial x} c) = 0, \quad (1.1)
\]

\[
\begin{cases}
  w_s c + \partial_z (K_v c) = 0, & \text{at the water surface (no flux)}, \\
  w_s c + K_v \partial_z c = D - E, & \text{at the bed (flux equals } D - E), \\
\end{cases} \quad (1.2)
\]

in which \( t \) represents time, \( x \) and \( z \) are the coordinates in longitudinal and vertical direction, \( c \) is the suspended sediment concentration, \( u \) and \( w \) are the water velocity in the \( x \)- and \( z \)-direction, \( w_s \) is the settling velocity, \( B \) is the time-invariant width of the Scheldt estuary, \( K_v \) is the constant vertical eddy diffusivity coefficient, \( K_h \) the constant horizontal diffusivity coefficient and \( D \) and \( E \) are the deposition and erosion of sediment defined as

\[
D = w_s c, 
E = M |\tau_b| f(a), \quad (1.3)
\]

where \( M \) is the constant erosion parameter, \( \tau_b \) is the shear stress at the bed, and \( f(a) \) is the erodibility.

The phytoplankton model consists of a dynamic model that describes the spatial and tidal evolution of phytoplankton concentration and corresponding nutrients nitrogen and phosphorus ([Dijkstra et al., 2019a](#)). We do not focus on the equation for the nutrient dynamics because the Scheldt estuary can be considered a nutrient-rich system, as mentioned above. The model consists of a single-phytoplankton class and thus does not differentiate between diatoms and non-diatoms. Consequently, Si-dynamics and salinity stress are not included, and phytoplankton characteristics are assumed constant in the system. The width-averaged equation for the phytoplankton concentration \( P \) and corresponding boundary conditions read ([Dijkstra et al., 2019a](#))

\[
\begin{aligned}
\frac{\partial}{\partial t} P + u \frac{\partial}{\partial x} P + (w - w_P) \frac{\partial}{\partial z} P - \frac{1}{B} \frac{\partial}{\partial z} (BK_h \frac{\partial}{\partial x} P) & = (\mu - m) P, \\
\text{advection-diffusion} & = \text{balance between local growth and mortality} \\
\end{aligned} \quad (1.5)
\]

\[
\begin{cases}
  w_P P + K_v \partial_z P = 0, & \text{at the bed and water surface (no flux)}, \\
  \frac{1}{H} \int_{-\zeta}^{\zeta} P dz = P_{\text{sea}}, & \text{at the seaside boundary (constant concentration)}, \\
  B \int_{-H}^{\zeta} (uP - K_h \partial_z P) dz = QP, & \text{at the upstream boundary (constant influx)}, \\
\end{cases} \quad (1.6)
\]

in which \( w_P \) is the constant settling velocity of phytoplankton cells, \( \langle \cdot \rangle \) denotes averaging over a long time scale (i.e., larger than a tide or day), \( \zeta \) and time-invariant \( -H \) are the \( z \)-coordinates of the water surface and bed, \( P_{\text{sea}} \) is the constant phytoplankton concentration at the seaside boundary, \( QP \) is the constant influx of phytoplankton at the upstream boundary, and \( \mu \) and \( m \) are the growth and (constant) mortality rate of phytoplankton.
The model only focuses on the estuary-scale hydro-, SPM, and phytoplankton dynamics by approximating the estuary’s bathymetry and width by smooth profiles. The model resolves the tidal and subtidal water motion and cohesive SPM concentration and provides approximate solutions of the complex and non-linear set of equations for hydro and SPM dynamics using a scaling and perturbation approach. This approach simplifies the interpretation of the results, allows us to focus on individual processes, and drastically decreases computation time, enabling an extensive sensitivity analysis. For the technical details, we refer the reader to Chernetsky et al. (2010), Dijkstra et al. (2017), and Dijkstra et al. (2019a).

In this thesis, we implement three major extensions to the iFlow model. Firstly, we extend the iFlow model by a flocculation model to resolve flocculation of cohesive sediment, which forms an important link between phytoplankton and SPM. Here, we also apply a scaling and perturbation approach following Chernetsky et al. (2010) and Dijkstra et al. (2017). Secondly, we use an alternative approximated light limitation function for local phytoplankton growth [cf. \( \mu \) in Eq. (1.5)]. This alternative allows us to further decrease the computational cost and include the impact of temporal variability in water depth on phytoplankton productivity, which is important in shallow regions. Thirdly, we consider two phytoplankton groups (i.e., freshwater and marine diatoms), thereby implementing salinity stress. Additionally, we make the mortality rate dependent on two (meso)zooplankton groups: calanoids and non-calanoids.

1.4.3 Research questions and outline

The extended model combined with the long-term observations in the Scheldt estuary is used to answer the main research questions, which we present in this section. Each research question corresponds to a Chapter. The outline of this thesis is illustrated in Fig. 1.3.

- What is the impact of flocculation on the large-scale SPM dynamics?

In Chapter 2, we determine whether flocculation can change the estuary-scale SPM distribution. To this end, we extend the iFlow model by a flocculation model and apply the model to a winter case in the Scheldt estuary.

- Can we detect a seasonal biotic impact on the SPM dynamics through flocculation and erosion?

Assuming a strong biotic impact on flocculation and erosion and knowing that biota typically show strong seasonal behavior, we expect to detect seasonality in biotically-induced flocculation and erosion. In Chapter 3, we estimate the impact of biota on the seasonality of the sediment distribution through flocculation and erosion on the seasonal and estuary scale using the results from Chapter 2. This insight is crucial when constructing a
Figure 1.3: Illustration of the interconnection between SPM and phytoplankton studied in this thesis and corresponding chapters. SPM limits phytoplankton growth by deteriorating the light climate in the water column. Phytoplankton may affect SPM dynamics by altering the floc size and structure of flocs and excreting sticky substances that stimulate flocculation of SPM. In Chapter 2, we study the impact of flocculation on SPM. In Chapter 3, we focus on the importance of biotically-induced flocculation on SPM. In Chapter 4, we study the impact of temporal variability in light climate (cf. SPM) on phytoplankton. We combine all results and study a specific case regarding phytoplankton-SPM dynamics in the Scheldt estuary in Chapter 5.

• Can we construct a model to easily assess the relative impact of temporal fluctuations in SPM on phytoplankton growth?

Before proceeding to a fully complex phytoplankton-SPM model, we develop a tool to locally study the impact of temporal variability in light climate on primary production in Chapter 4 without solving the complex SPM dynamics explicitly. The tool allows us to determine the relative impact of temporal variability in the SPM dynamics without a priori assumptions regarding these dynamics. We aim to know whether we may neglect temporal variability in these parameters and, if not, how we can correct for the introduced error by excluding this temporal variability. We again apply the tool to the Scheldt estuary.

• Can we link the disappearance of the spring bloom in the brackish region in the
Scheldt estuary between 2004-2018 to multi-annual changes of processes impacting phytoplankton growth and their corresponding model parameters?

In Chapter 5, we combine all the previous results from Chapters 2, 3, and 4 to construct a sediment transport - phytoplankton model. To show its applicability, we again apply the model to the Scheldt estuary. We focus on the appearance and consequent disappearance of phytoplankton blooms in spring in the brackish region in the Scheldt estuary in the last two decades, which we introduced at the end of Section 1.3.

List of symbols

Latin

- $a$: Availability of easily erodible fine sediment at the sediment bed
- $B$: Width of the estuary
- $c$: Suspended sediment concentration
- $D$: Deposition flux of suspended sediment
- $E$: Erosion flux of sediment from the sediment bed
- $f(a)$: Erodibility
- $H$: Magnitude of the z-coordinate of the river bed
- $K_h$: Horizontal diffusivity coefficient
- $K_v$: Vertical eddy diffusivity coefficient
- $M$: Erosion parameter
- $m$: Mortality rate of phytoplankton
- $P$: Phytoplankton concentration
- $P_{\text{sea}}$: Phytoplankton concentration at the seaside boundary
- $QP$: Influx of phytoplankton at the upstream boundary
- $t$: Time
- $u$: Water flow velocity in the longitudinal dimension
- $w$: Water flow velocity in the vertical dimension
- $w_P$: Settling velocity of phytoplankton cells
- $w_s$: Settling velocity of suspended sediment
- $x$: Spatial coordinate in the longitudinal dimension
- $z$: Spatial coordinate in the vertical dimension

Greek

- $\zeta$: z-coordinate of the water surface
- $\mu$: Growth rate of phytoplankton
- $\tau_b$: Shear stress at the river bed
Bibliography


Chapter 2

The impact of flocculation on the large-scale SPM dynamics

Abstract

Sediment transport in estuaries and the formation of estuarine turbidity maxima (ETM) highly depend on the ability of suspended particulate matter (SPM) to flocculate into larger aggregates. While most literature focuses on the small-scale impact of biological flocculants on the formation of larger aggregates, the influence of the flocculation process on large-scale estuarine SPM profiles is still largely unknown. In this paper, we study the impact of flocculation of SPM on the formation of ETM. For this, a semi-analytical width-integrated model called iFlow is utilized and extended by a flocculation model. Starting from a complex one-class flocculation model, we show that flocculation may be described as a linear relation between settling velocity and suspended sediment concentration to capture its leading-order effect on the ETM formation. The model is applied to a winter case in the Scheldt Estuary (Belgium, Netherlands) and calibrated to a unique, long-term, two-dimensional set of turbidity (cf. SPM) observations. First, model results with and without the effect of flocculation are compared, showing that the spatial and temporal variations of the settling velocity due to flocculation are essential to reproduce the observed magnitude of the suspended sediment concentrations and its dependence on river discharge. Second, flocculation results in tidally averaged land-inward sediment transport. Third, we conduct a sensitivity analysis of the freshwater discharge and floc break-up parameter, which shows that flocculation can cause additional estuarine turbidity maxima and can prevent flushing of the ETM for high freshwater inflow.

2.1 Introduction

Estuaries often contain regions in which the concentration of suspended particulate matter (SPM) is larger than landward and seaward of these regions. These regions are called estuarine turbidity maxima (ETM) and are caused by fine sediments being trapped as a result of the complex interactions of the water motion and sediment dynamics. Examples of such transport mechanisms are, amongst others, related to tidal asymmetries in water motion, water density gradients, and transport mechanisms related to temporal variability in settling velocity of SPM (see Burchard et al. (2018) for a recent review). One mechanism resulting in temporal and spatial variability of settling velocity is flocculation. Flocculation results in the aggregation and break-up of cohesive SPM, thus changing the floc size and hence the settling velocity.

Two main classes of models describing flocculation can be distinguished, the Lagrangian flocculation (LF) models and the population balance equation (PBE) models (Lai et al., 2018). PBE flocculation models typically consist of multiple discrete size classes and compute the evolution of the number of flocs within each class over time (Smoluchowski, 1918; Verney et al., 2011). Although PBE models can calculate the evolution of particle size distribution over time, they come with high computational costs. In contrast, the LF models are dynamic models that resolve the dynamics of particle size and settling velocity by computing a balance between floc aggregation and floc break-
Floc aggregation and break-up are implemented using empirical formulations, which depend on both abiotic and biotic factors (Dyer, 1989; van Leussen, 1994; Lai et al., 2018).

A first important abiotic factor that impacts flocculation is the suspended sediment concentration. Both in situ measurements (Pejrup and Mikkelsen, 2010) and laboratory experiments (Tran et al., 2018) show that floc size typically increases with increased suspended sediment concentration. Another important abiotic driver is turbulence, which both promotes aggregation through enhanced mixing and break-up by increasing the shear stresses on the flocs. Turbulence promoting break-up is observed in various estuaries such as the Yangtze estuary (Guo et al., 2017) and the Scheldt estuary (Manning et al., 2007; Schwarz et al., 2017). Also, laboratory experiments show that turbulence can decrease the averaged floc size (Mietta et al., 2009). A third abiotic environmental condition that impacts flocculation is salinity. Edzwald et al. (1974) conducted laboratory experiments in which they showed that salinity promotes aggregation of clay particles and very little salt ($\sim 5$ ppt) is already sufficient to reach a maximum impact of salinity on flocculation. In contrast, Eisma et al. (1980) and Verney et al. (2009) found that salt flocculation is not a crucial factor in the Rhine estuary and Seine estuary, respectively. Biotic characteristics that impact flocculation are the organic content that directly alters the differential density and structure of flocs (Kranenburg, 1994; van Leussen, 1994). Furthermore, organic content impacts, for example, the floc strength and collision efficiency (Winterwerp and van Kesteren, 2004), averaged floc size (Mietta et al., 2009), and floc break-up (Alldredge et al., 1990). Finally, in situ observations show a correlation between Chlorophyll-a and flocculation efficiency (Verney et al., 2009) and sticky biotic substances (i.e., Transparent Exopolymer Particles) and floc strength (Fettweis et al., 2014).

Although temporal and spatial variations in settling velocity, which is related to floc size, are recognized as potentially important mechanisms for sediment trapping, most models for estuarine sediment transport use either a constant settling velocity or empirical flocculation relationships. Only a few studies included full floc dynamics in a sediment transport model (Ditschke and Markofsky, 2008; Xu et al., 2010; Sherwood et al., 2018; Shen et al., 2018), but the importance of flocculation on sediment transport mechanisms cannot be clearly identified from these model results.

This study aims to model and gain an understanding of the impact of varying settling velocity on large-scale sediment transport and the corresponding development of ETM. As a first step, we consider estuaries in which the suspended sediment concentration is lower than 1 g L$^{-1}$, allowing us to focus on flocculation process and neglect hindered settling effects, which also result in a temporally and spatially varying settling velocity.

To be able to gain an understanding of the underlying mechanisms that result in sediment trapping and quantify the relative importance of flocculation on ETM formation, we extend the width-averaged, hydrodynamics and sediment transport model known as ‘iFlow’ (Dijkstra et al., 2017) to include flocculation processes. The iFlow model is specifically geared toward gaining an understanding of the water motion, sediment transport, and trapping in tidally dominated estuaries and allows for extensive sensitivity analysis. The model is extended with the one-class LF model of Winterwerp (2002) in which the
impact on flocculation of suspended sediment concentration, turbulent shear, and floc size and structure are directly parameterized in the floc aggregation and break-up terms. Other effects, such as salinity and biotic factors, are indirectly included by fitting to observations.

We apply the extended iFlow model to the Scheldt estuary. The Scheldt estuary is located in Belgium and the Netherlands and debouches into the North Sea. Being a well-documented estuary (Meire et al., 2005), there have been intensive monitoring campaigns (Maris and Meire, 2016) and modeling experiments. Moreover, the iFlow model without flocculation has already successfully been applied to the Scheldt estuary (Brouwer et al., 2018; Dijkstra et al., 2019a). The flocculation process is expected to be important in the Scheldt estuary (Peters, 1972; Gourgue et al., 2013).

In this paper, we extend the iFlow model to include the effect of flocculation in Section 2.2. In Section 2.3, we calibrate the model to a unique, long-term turbidity (cf. SPM) dataset of the Scheldt estuary. In Section 2.4, we apply the coupled flocculation-sediment transport model to the Scheldt estuary and study the effect of flocculation on the large-scale sediment distribution and the underlying sediment transport mechanisms.

2.2 Model

In this section, we first present the sediment transport model (i.e., iFlow) and flocculation model of Winterwerp (2002). Next, we use scaling and perturbation methods to simplify the coupling of the two models. Then, we describe the theoretical impact of flocculation on sediment transport. Last, we introduce the (numerical) implementation of our model.

2.2.1 iFlow

The iFlow model (Dijkstra et al., 2017) was developed to obtain the width-averaged water motion, sediment transport, and trapping in a tidally dominated estuarine system by solving the width-averaged shallow water equations, suspended sediment concentration equation, and a dynamic equation for the erodibility of the bed (Brouwer et al., 2018). The water motion is assumed to be forced by an M2 and M4 tidal signal at the mouth, while a river discharge is prescribed at the upstream tributaries. It is assumed that the M2 tidal signal dominates over the M4 and riverine signals. We decompose the water motion into a longitudinal and vertical velocity component, denoted by \( u \) and \( w \). The suspended sediment concentration follows from an advection-diffusion equation and is forced by a constant suspended sediment concentration at the mouth and prescribed inflow of sediment at the upstream tributaries. The bathymetry and width are approximated by smooth polynomials, thereby neglecting the impact of small-scale gradients, as illustrated in Fig. 2.1. A diagnostic longitudinal salinity gradient is prescribed, consisting of a depth- and time-independent sigmoid function.
By assuming that the dynamics consist of strong tidal signals beside a subtidal part, the iFlow model solves the problem in frequency space for subtidal, M2, and M4 components. The water surface elevation is assumed small compared to the subtidal water depth, resulting in a small dimensionless parameter, given by the ratio of water surface elevation and water depth. Using a scaling procedure and relating the various dimensionless numbers to this small parameter allows for identifying the relative importance of individual processes (e.g., river, tide, density forcing, advection). Next, a perturbation approach is employed to solve the equations at different order. For further details on the iFlow model, we refer the reader to Dijkstra et al. (2017).

Using this information and truncating frequencies larger than M4 (i.e., M6, M8, etc. are neglected and, hence, not discussed), the leading-order water motion $u^0$ and suspended sediment concentration $c^0$ consist of an M2 tidal signal and subtidal and M4 tidal components, respectively (Chernetsky et al., 2010; Dijkstra et al., 2017), denoted by $u^{02}$ and $c^{00}$, $c^{04}$ (in which the first superscript denotes the order and the second one denotes the frequency) and defined as

$$u^0(x, z, t) = \text{Re} \left[ \tilde{u}^{02}(x, z) e^{i\omega_{M2} t} \right]$$

and

$$c^0(x, z, t) = \text{Re} \left[ \tilde{c}^{00}(x, z) + \tilde{c}^{04}(x, z) e^{2i\omega_{M2} t} \right].$$

with $x$ and $z$ being the longitudinal and vertical coordinates and $t$ being the time. At first order, the water motion $u^1$ and suspended sediment concentration $c^1$ consist of a subtidal and M4 tidal signal and an M2 tidal signal, respectively (Chernetsky et al., 2010; Dijkstra et al., 2017).
et al., 2017), denoted by $u^{10}$, $u^{14}$, and $c^{12}$ and defined as

$$u^{1}(x, z, t) = \text{Re} \left( \hat{u}^{10}(x, z) + \hat{u}^{14}(x, z)e^{2i\omega_{M2}t} \right)$$  \hspace{1cm} (2.3)$$

$$c^{1}(x, z, t) = \text{Re} \left( \hat{c}^{12}(x, z)e^{i\omega_{M2}t} \right).$$  \hspace{1cm} (2.4)$$

The net sediment transport $T$, i.e., the sediment transport averaged over a tidal period (denoted by angle brackets), is given by the sum of the advective and diffusive sediment transport integrated over the cross section:

$$T = \left\langle B \int_{-H}^{R+\xi} uc - K_{h}(\partial_{x}c)dz \right\rangle$$

$$\approx \left\langle B \int_{-H}^{R} \left( u^{0}c^{0} + u^{1}c^{0} + u^{0}c^{0} - K_{h}(\partial_{x}c)dz + u^{0}\partial_{x}c^{0} \right) \bigg|_{z=R} \right\rangle$$

$$= \left\langle B \int_{-H}^{R} \left( u^{10}c^{00} + u^{14}c^{04} + u^{12}c^{12} - K_{h}(\partial_{x}c)dz + u^{0}\partial_{x}c^{0} \right) \bigg|_{z=R} \right\rangle,$$  \hspace{1cm} (2.5)$$

where $B$ is the local channel width, $R$ is the reference level of the water surface elevation, $H(x)$ is the magnitude of the local z-coordinate of the river bed, and $K_{h}$ is the horizontal eddy diffusivity coefficient. Moreover, we used that only subtidal-subtidal, $M2$-$M2$, and $M4$-$M4$ tidal interaction terms result in net sediment transport, which is a direct result of the orthogonality of the complex exponential functions.

In iFlow, parameters such as the total freshwater discharge $Q$, horizontal eddy diffusivity coefficient $K_{h}$, and settling velocity $w_{s}$, are usually prescribed as constant parameters [although this is not needed, see Dijkstra et al. (2017)], allowing for a quick solution procedure. To include flocculation in the model, the settling velocity $w_{s}$ has to be coupled to the suspended sediment concentration $c$, which makes the model nonlinear, requiring an iterative solution procedure (see Section 2.2.5).

### 2.2.2 Flocculation model

The Winterwerp flocculation model employed in this paper reads (Winterwerp, 2002) as

$$\partial_{t}N + \partial_{x}[uN] + \partial_{z}[(w - w_{s})N] - \partial_{x}[K_{h}\partial_{x}N] - \partial_{z}[K_{v}\partial_{z}N]$$

$$= -k'_{A}GD_{f}^{3}N^{2} + k_{B}G^{3/2}(D_{f} - D_{p})D_{f}N,$$  \hspace{1cm} (2.6)$$

where $N$ is the number of flocs per unit volume, $w_{s}$ is the settling velocity, $K_{v}$ is the vertical eddy diffusivity coefficient, $G$ is the shear rate, $D_{f}$ and $D_{p}$ are the floc and primary
particle size, and $k'_A$ and $k_B$ are the empirical aggregation and break-up parameters. To acquire Eq. (2.6), we assumed a fractal dimension $n_f = 2$ and we set model calibration parameters $q = 0.5$ and $p = 1$, following Winterwerp (2002).

The aggregation parameter $k'_A$ and break-up parameter $k_B$ define the efficiency of the flocculation process and represent the intrinsic flocculation kinetics of the system of interest. Laboratory flocculation experiments show that floc kinetics depend, amongst others, on salinity (Peters, 1972; Edzwald et al., 1974). To include a salinity dependence in our model, we allow the flocculation kinetics to depend on the depth- and tidally averaged salinity profile, resulting in $k'_A$ and $k_B$ being functions of the longitudinal coordinate $x$. In Section 2.2.3, we show that net sediment transport only depends on the ratio of $k'_A$ and $k_B$ and not on the individual parameters. From the experiments carried out by Edzwald et al. (1974), we therefore postulate the following dependence of the ratio of $k'_A$ and $k_B$ on salinity $S$:

$$k'_A/k_B \sim \left\{ 1 + S_1 [1 + \tanh(S(x) - S_2)] \right\},$$  

(2.7)

with $S_1$ and $S_2$ being empirical parameters that are calibrated to a salinity $S(x)$ dataset. For more details, the reader is referred to Appendix 2.A.

Following Pejrup and Mikkelsen (2010), the shear rate $G$ reads as

$$G(x, \tilde{z}_1) = \sqrt{\frac{\langle u_* \rangle^3 (1 - \tilde{z}_1)}{\nu H \kappa' \tilde{z}_1}},$$  

(2.8)

where $\kappa'$ is the Von Karman constant, $\nu$ is the kinematic viscosity, $\tilde{z}_1 = z_1/H$ is the relative water depth, and $\langle u_* \rangle$ is the subtidal friction velocity:

$$\langle u_* \rangle(x) = \frac{\kappa'}{\ln \frac{z_1(x)}{s_{f0}}} \left( |u^0(x, z_1) + u_{riv}^{10}(x, z_1)| \right),$$  

(2.9)

where $z_1$ is the distance in the logarithmic layer above the bed, $s_{f0}$ is the bed roughness coefficient, $u^0$ is the leading-order, longitudinal flow velocity [see Eq. (2.1)], and $u_{riv}^{10}$ is the subtidal, first-order, longitudinal, river-induced flow velocity [see Eq. (2.3)]. For simplicity, we only consider the impact of subtidal shear velocity $u_*$ at leading order. The first-order, subtidal, riverine contribution is also included (i.e., $u_{river}^{10}$) because this constituent is dominant at the upstream border. Apart from neglecting tidal variations of $G$, we also approximate $G$ by its value at $z_1 = H/2$ and use this $G(x, \tilde{z}_1 = 1/2)$ as a proxy for the whole water column. Working with a depth-averaged value of $G$ instead of a midpoint value has no major impact on our results and conclusion because vertical variations in $G$ are relatively small (typically < 10% in our case study) in the logarithmic layer. Although turbulence is an important driver of flocculation, we do not focus on this variable in our case study because we assume a tide-dominated system with low stratification. This results in a shear rate that has a similar order of magnitude within the ETM.

The Winterwerp model depends on both the number of flocs per unit volume $N$, floc size $D_f$, and settling velocity $w_s$. These three variables are not independent. We
Table 2.1: Definition of $\beta$, $\kappa$, $\tau$ and $\gamma$, along with their units.

<table>
<thead>
<tr>
<th>variable</th>
<th>definition</th>
<th>units</th>
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</thead>
<tbody>
<tr>
<td>$\gamma$</td>
<td>$18 \mu g^{-1} (\rho_s - \rho_w)^{-1}$</td>
<td>m s $^{-1}$</td>
</tr>
<tr>
<td>$\beta$</td>
<td>$\frac{1}{18 \mu f_s} (\rho_s - \rho_w) \frac{g}{\sqrt{G}} k'_{A}$</td>
<td>m$^{4}$ s$^{-1}$ kg$^{-1}$</td>
</tr>
<tr>
<td>$\kappa$</td>
<td>$k'<em>{A} D</em>{p}^{2} f_{s} \rho_{s} \sqrt{G}$</td>
<td>kg m$^{-3}$</td>
</tr>
<tr>
<td>$\tau$</td>
<td>$\left(\frac{\rho_s - \rho_w}{18 \mu} g D_{p} \right)^{2} k'_{B} G^{-3/2}$</td>
<td>m$^{2}$ s$^{-1}$</td>
</tr>
</tbody>
</table>

rewrite Eq. (2.6) such that it solely depends on the settling velocity $w_s$. First, we express the number of flocs per unit volume $N$ in terms of the floc size $D_{f}$ (Kranenburg, 1994; Winterwerp and van Kesteren, 2004):

$$N = \frac{1}{f_s \rho_s D_{p} D_{f}^{2}},$$

(2.10)

with $f_s$ being the floc shape factor and $\rho_s$ the floc density.

Next, to obtain an equation for the settling velocity, we use the generalized Stokes formulation to translate floc size $D_f$ to settling velocity $w_s$ (Winterwerp and van Kesteren, 2004):

$$w_s = \frac{\rho_s - \rho_w}{18 \mu} g D_{p} D_{f},$$

(2.11)

where $\rho_w$ is the water density, $g$ is the gravitational acceleration, and $\mu$ is the viscosity of water. We assumed spherical flocs ($f_s = \pi/6$), a fractal dimension $n_f = 2$, and that the floc Reynolds number $\text{Re}_f = w_s D_f / \nu \ll 1$.

Combination of Eqs. (2.6), (2.10), and (2.11) gives a differential equation for $w_s$ that results in the evolution of $w_s$ over time and space:

$$
\partial_t cw_{s}^{-2} + \partial_x [ucw_{s}^{-2}] + \partial_x [(w - w_s)cw_{s}^{-2}] - \partial_x [K_h \partial_x cw_{s}^{-2}] - \partial_z [K_v \partial_z cw_{s}^{-2}] = \gamma w_{s}^{-1} cG \left[ -k'_{A} D_{p}^{-2} f_{s} \rho_{s}^{-1} + k_{B} G^{1/2} (\gamma D_{p}^{-2} w_{s} - 1) \right] \tag{2.12}
$$

where $\gamma$ is defined in Table 2.1. In the following section, we approximate this equation and write the settling velocity $w_s$ as an explicit function of the suspended sediment concentration $c$.

### 2.2.3 Ordering of the flocculation model

We apply scaling and perturbation analysis to the Winterwerp flocculation model shown in Eq. (2.12) to gain more insight into this highly complex equation. We refer the reader to Appendices 2.B, 2.C, and 2.D for the derivation.
2.2.3.1 Leading order

At leading order, the flocculation model reduces to the balance between floc aggregation and floc break-up:

\[
0 = -k'_A c^0 D_p^{-2} f_s^{-1} \rho_s^{-1} + k_B G^{1/2} (\gamma D_p^{-2} w_s^0 - 1) .
\]  

(2.13)

This can be interpreted as if the flocculation process is instantaneous and local: there is no inertia and transport. As a result, the settling velocity scales linearly with the suspended sediment concentration:

\[
w_s^0 = \beta (c^0 + \kappa),
\]  

(2.14)

where \(\beta\) and \(\kappa\) are defined in Table 2.1 and depend on several parameters including the shear rate and ratio between \(k'_A\) and \(k_B\). Using the definitions of \(\beta\) and \(\kappa\) (Table 2.1), we see that the \(\beta \kappa\) term equals the settling velocity \(w_{s,\text{min}}\) corresponding to the settling velocity of primary particles. Indeed, it equals the Stokes formulation for massive \((n_f = 3)\) primary particles with primary particle size \(D_p\). This ensures that for \(c \to 0\), \(w_s^0 \to w_{s,\text{min}}\). Moreover, our leading-order result in Eq. (2.14) is equivalent to the equilibrium floc size \(D_e\) presented in Winterwerp (2002):

\[
D_e = D_p + \frac{k'_A c}{k_B \sqrt{(G)}}.
\]  

(2.15)

The positive correlation of \(w_s\) with suspended sediment concentration in the leading-order result complies with observations. Indeed, Pejrup and Mikkelsen (2010) assuming a power relationship between settling velocity and suspended sediment concentration, found exponents ranging between 0.47 and 2.9 with an average of 1.29, using 18 different tidal systems based on real measurements. Recently, Tran et al. (2018) studied the influence of SPM concentration on floc size in laboratory mixing tanks. They found a linear relationship between floc size and SPM concentration, which again results in the linear relationship between settling velocity and SPM concentration as given in Eq. (2.14).

2.2.3.2 First order

At first order, our scaling procedure shows that local inertia (cf. \(\partial_t N\) term), settling, and vertical diffusion become significant [see Eq. (2.55)]. Advection and horizontal diffusion are still negligible at this order. The solution then reads as

\[
w_s^1 = \beta c^1 \, \text{break-up/aggregation} - 2\tau \frac{\partial_z c^0}{c^0} \left[ 1 + \frac{K_0}{w_s^0} \frac{\partial_z c^0}{c^0} \right],
\]  

(2.16)

where \(\tau\) is defined in Table 2.1. Apart from a contribution to \(w_s^1\) that again scales linearly in \(c\) (\(\beta c^1\) term), we obtain additional terms when compared to the leading-order result.
Because of inertial effects, settling, and vertical diffusion, we obtain an additional contribution that is constant in time and one that varies on the M4 time scale, no additional M2 contribution is generated [see Eqs. (2.86) and (2.87)]. As we show below, we are only interested in the M2 contribution $w_{s12}$ because this is the only first-order contribution that is important for net sediment transport. The only M2 signal in the first-order settling velocity is related to the first-order balance between aggregation and break-up and reads as

$$w_{s12} = \beta c_{12}. \quad (2.17)$$

### 2.2.4 Impact of flocculation on net sediment transport

In the previous section, we discussed the impact of suspended sediment concentration $c$ on settling velocity $w_s$, which is a direct consequence of flocculation. In the following, we present the influence of settling velocity $w_s$ on suspended sediment concentration $c$. As flocculation alters the leading and first-order suspended sediment concentration $c^0$ and $c^1$, the corresponding net sediment transport $T$ from Eq. (2.5) is also impacted. We rewrite Eq. (2.5) as

$$T_{floc} = \left\langle B \int_{-H}^{R} \left[ u_{10}^{c_{floc}} + u_{14}^{c_{floc}} + c_{floc}^{12} - K_h (\partial_x c) dz + u_0^0 c_0^0 \left|_{z=R} \right. \right] \right\rangle, \quad (2.18)$$

and examine the relation of $c_{floc}^{00}$, $c_{floc}^{04}$, and $c_{floc}^{12}$ with the settling velocity $w_s$.

A careful analysis shows that at leading order, both the settling velocity and suspended sediment concentration consist of a subtidal and M4 tidal signal. This can be inferred from the following argument: if at leading order, the (truncated) suspended sediment concentration $c^0$ consists of only a subtidal and M4 tidal signal as found for a time-independent $w_s$ [see Eq. (2.2)], the settling velocity also gets an M4 tidal component [see Eq. (2.14)]. This M4 signal impacts the leading-order suspended sediment concentration $c^0$ through the nonlinear interaction term $w_0^s c_0^1$ in the leading-order differential equation for $c^0$ [see Eq. (2.44)]. Interestingly, the nonlinear interaction of $c^0$ and $w_0^s$ results in an adjustment of the subtidal and M4 tidal signal in $c^0$, that is, $c_{floc}^{00}$ and $c_{floc}^{04}$, and generates an infinite number of additional tidal frequencies in $c^0$ because of its nonlinearity:

$$w_0^s \times \left( \frac{c_0^0 + c_0^{04} + \kappa}{c_{floc}^{00} + c_{floc}^{04}} \right) \text{generates } c_{floc}^{00} + c_{floc}^{04}. \quad (2.19)$$

As stated before, frequencies larger than M4 are truncated.

Additionally, the subtidal and M4 tidal signal in $w_0^s$ impacts the M2 signals in the first-order differential equation for $c^1$ through the $\sim w_0^s c_1^1$ forcing term [see Eq. (2.47)], which again results in an adjustment of net sediment transport through $c_{floc}^{12}$:

$$w_0^s \times \left( \frac{c_1^0 + c_1^{12} + c_1^{04}}{c_{floc}^{10} + c_{floc}^{12} + c_{floc}^{14}} \right) \text{generates } c_{floc}^{10} + c_{floc}^{12} + c_{floc}^{14}. \quad (2.20)$$
Consequently, at leading order both the subtidal and M4 tidal signal in $w^0_s$ have an impact on net sediment transport through its impact on $c_{\text{floc}}^{00}$ and $c_{\text{floc}}^{04}$ and $c_{\text{floc}}^{12}$.

At first order, only the M2 tidal signal in $w^1_s$, i.e., $w^{12}_s$, results in net sediment transport through the forcing term $\sim w^1_s c^0$ in the first-order differential equation for $c^1$ [see Eq. (2.47)]:

$$w^1_s \begin{cases} w^{12}_s \\ w^{10}_s + w^{14}_s \times c^0(z) \end{cases} \text{ generates } \begin{cases} c_{\text{floc}}^{12} \\ c_{\text{floc}}^{10} + c_{\text{floc}}^{14} \end{cases} .$$

Consequently, when interested in net sediment transport, the first-order settling velocity simplifies to the linear relationship in Eq. (2.17) and the Winterwerp model up to first order is thus equivalent to the equilibrium floc size result in Eq. (2.15).

### 2.2.5 Solution method

iFlow contains a numerical sediment module that solves for suspended sediment concentration $c$ numerically in $x$ and $z$ using a grid of 200 cells in the $x$ direction and 50 cells in the $z$ direction. To do so, the module requires a leading-order settling velocity $w^0_s$ and first-order settling velocity $w^1_s$ as input. A new module is added to iFlow containing the explicit analytical expression of $w^0_s$ and $w^1_s$ [Eqs. (2.14) and (2.17)]. The coupling of $w_s$ to $c$ results in a $w_s$ which is a function of time and space. Because the coupling is nonlinear, we solve the coupling iteratively using the Picard method. We start with an initial condition for $w^0_s$ and $w^1_s$ as input to the sediment module to acquire a suspended sediment concentration at leading order $c^0$ and first order $c^1$. Next we use $c^0$ and $c^1$ to recalculate $w^0_s$ and $w^1_s$ using Eqs. (2.14) and (2.17). We use the recalculated $w^0_s$ and $w^1_s$ as new input to the sediment module. We repeat this procedure until the solution converges using the following stop criterion:

$$\frac{|w^0_s(x, z) - w^0_{s,\text{old}}(x, z)|}{w^0_{s,\text{old}}(x, z)} < 10^{-3} ,$$

in which $w^0_{s,\text{old}}$ is the leading-order settling velocity of the previous iteration. The iterative procedure is schematized in Fig. 2.2.

### 2.3 The Scheldt: observations and calibration

We apply our coupled flocculation-sediment transport model to a winter case (i.e., Jan.-Mar.) in the Scheldt estuary. To do so, we calibrate the model to a unique, long-term turbidity (cf. SPM) dataset for 2015 until 2017 measured within the environmental monitoring program called Onderzoek Milieu Effecten Sigmaplan (OMES). We divide the turbidity dataset in winter and summer because the turbidity data shows a strong seasonality in the Scheldt estuary (Maris and Meire, 2016). This section presents the Scheldt estuary, the long-term turbidity dataset, and the calibration method.
2.3.1 The Scheldt

The Scheldt estuary (Fig. 2.3) is a funnel-shaped estuary of approximately 160 km long, which flows through Belgium and debouches into the North Sea near Vlissingen (Netherlands). The Scheldt estuary has a relatively low averaged total freshwater discharge (172 m$^3$ s$^{-1}$ in our winter case) and can be considered a tide-dominated estuary (Meire et al., 2005; Waterinfo.be, 2019). The main tributaries are the Upper Sea Scheldt (upstream boundary; ~34% of total river discharge), the Rupel (at 95 km from the mouth; ~54% of total river discharge), and the Dender (at 123 km from the mouth; ~12% of total river discharge) (Waterinfo.be, 2019). Parameter values used in our winter case study in the Scheldt estuary are summarized in Table 2.2. These parameter values follow from Brouwer et al. (2018) if not mentioned explicitly in the text. The total river discharge $Q$, erosion parameter $M$, and floc break-up coefficient $k_B$ follow from calibration to the OMES SPM dataset presented in the following section.

2.3.2 OMES data

We calibrate our coupled flocculation-sediment transport model to a unique dataset representing a typical winter situation of SPM-distribution. In the frame of the ongoing OMES monitoring, vertical turbidity profiles were obtained using conductivity-temperature-depth (CTD) and turbidity casts from aboard a ship at 16 fixed locations.
Figure 2.3: The Scheldt estuary and its two main tributaries (Dender and Rupel). The red dots represent locations where monthly and biweekly CTD and turbidity casts were performed in the frame of the environmental monitoring program (OMES).
Table 2.2: Parameter values used in our winter case study in the Scheldt estuary.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A^0$</td>
<td>M$_2$ water level amplitude at x=0</td>
<td>1.77 m</td>
</tr>
<tr>
<td>$A^1$</td>
<td>M$_4$ water level amplitude at x=0</td>
<td>0.14 m</td>
</tr>
<tr>
<td>$\phi^1$</td>
<td>M$_4$ water level phase relative to M$_2$ tide at x=0</td>
<td>-1.3 deg</td>
</tr>
<tr>
<td>$Q$</td>
<td>Total river discharge in winter (calibrated)</td>
<td>233 m$^3$ s$^{-1}$</td>
</tr>
<tr>
<td>$c_{sea}$</td>
<td>Depth-averaged subtidal suspended sediment concentration at x=0</td>
<td>0.06 kg m$^{-3}$</td>
</tr>
<tr>
<td>$K_h$</td>
<td>Horizontal eddy diffusivity coefficient</td>
<td>100 m$^2$ s$^{-1}$</td>
</tr>
<tr>
<td>$K_\nu^0$</td>
<td>Vertical eddy diffusivity coefficient</td>
<td>$3.1 \times 10^{-2}$ m$^2$ s$^{-1}$</td>
</tr>
<tr>
<td>$M$</td>
<td>Erosion parameter in winter (calibrated)</td>
<td>$3.36 \times 10^{-3}$ s m$^{-1}$</td>
</tr>
<tr>
<td>$\sigma_\rho$</td>
<td>Prandtl-Smidth number ($=A_\nu/K_\nu^0$ with $A_\nu$ the vertical eddy viscosity)</td>
<td>1</td>
</tr>
<tr>
<td>$s_i^0$</td>
<td>Bed roughness coefficient (calibrated)</td>
<td>4.22 mm s$^{-1}$</td>
</tr>
<tr>
<td>$k^\text{min}_A$</td>
<td>Nondimensional minimal aggregation coefficient</td>
<td>0.29</td>
</tr>
<tr>
<td>$f_s$</td>
<td>Shape factor</td>
<td>$\pi/6$</td>
</tr>
<tr>
<td>$D_p$</td>
<td>Diameter primary mud particles</td>
<td>$4 \times 10^{-6}$ m</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Dynamic viscosity</td>
<td>0.0010518 Pa s</td>
</tr>
<tr>
<td>$\rho_s$</td>
<td>Density of sediment primary particles</td>
<td>2650 kg m$^{-3}$</td>
</tr>
<tr>
<td>$\rho_w$</td>
<td>Reference density of water</td>
<td>1000 kg m$^{-3}$</td>
</tr>
<tr>
<td>$k_B$</td>
<td>Floc break-up coefficient in winter (calibrated)</td>
<td>5600 s$^{1/2}$ m$^{-2}$</td>
</tr>
<tr>
<td>$S_1$</td>
<td>Flocculation salinity sensitivity calibration parameter</td>
<td>0.078</td>
</tr>
<tr>
<td>$S_2$</td>
<td>Flocculation salinity sensitivity calibration parameter</td>
<td>4.085 %</td>
</tr>
<tr>
<td>$s_{sea}$</td>
<td>Salinity boundary condition at the mouth</td>
<td>28.9 %</td>
</tr>
<tr>
<td>$x_{\text{sal}^1}$</td>
<td>Calibration parameter in postulated tanh salinity distribution in winter</td>
<td>37.8 km</td>
</tr>
<tr>
<td>$x_{\text{sal}^L}$</td>
<td>Calibration parameter in postulated tanh salinity distribution in winter</td>
<td>25.3 km</td>
</tr>
</tbody>
</table>
spread over the Belgian part of the Scheldt estuary during monthly or biweekly campaigns (Fig. 2.3). The campaigns were conducted independently of the tidal phase and spring neap tide. To obtain a typical winter distribution, we temporally averaged the Jan.-Mar. data from 2015-2017.

Turbidity was measured at various depths using an optical backscatter point sensor (OBS) of RBR type XR420 CTD+. During each campaign, the sensor was calibrated using a Formazine solution standard (Maris and Meire, 2016). The water body was vertically profiled using a minimal sampling frequency of $10^{-1}$ s$^{-1}$.

Simultaneously, two water samples were collected at each location using a Niskin bottle at approximately half the water depth and at the water surface. SPM concentrations were gravimetrically determined after filtration in the laboratory. During each campaign, 16 × 2 SPM water samples were collected, resulting in 32 SPM estimates. To translate turbidity to SPM, we used these 32 samples to apply a linear data fit. We assumed that the relation between turbidity and SPM is location (and time) independent, that is, equal for every location within one campaign. Calibration of turbidity to SPM resulted in a depth profile of SPM at 16 fixed locations in the Flemish part of the Scheldt estuary.

The winter case covers 11 measuring campaigns. We excluded data at the water surface and river bed because here, we typically have distortions due to, for example, air bubbles and high turbidity, respectively. To do so, we manually excluded depths at which we did not measure 8 of 11 measuring campaigns. We averaged the SPM concentration observations of the 11 measuring campaigns at each depth. We assumed this averaged value approximates the residual SPM concentration following, for example, Dijkstra et al. (2019a) and Cox et al. (2019). This assumption is valid when the number of estimates is sufficiently high, so averaging of the periodic temporal variability of the SPM concentration is negligible compared to the magnitude of the residual SPM concentration. Here, we assumed that the campaigns were randomly distributed within the tidal phase and spring neap tide, which was shown to be valid for OMES SPM sampling between 1995 and 2015 (Vandenbruwaene et al., 2016). Furthermore, we assumed that the SPM distribution within the time frame of our winter case is fixed on an estuarine scale (i.e., the scale of the iFlow model in the longitudinal direction). So, local effects due to, for example, temporal variations in river discharge can be neglected. Given a system-averaged standard deviation relative to the time-averaged SPM concentration of 0.43, we conclude that these assumptions are acceptable and that the number of estimates is sufficiently high.

On average, each turbidity profile consists of 20 individual observations, with an average distance between two consecutive observations of 0.36 m. We vertically interpolated the data to compare the subtidal SPM model output at the vertical model grid cells in our calibration. This results in a total number of 606 $(x,z)$ locations at which we compared data and model output. This number results from the fact that we measured at 16 stations and have 50 vertical model grid cells (a total of 800 grid cells) but excluded data near the water surface and river bed. The average number of data points $n$ for each location $(x,z)$ equals 9.4.
2.3.3 Calibration

We calibrated the linearized bed roughness coefficient \( s_{f0} \) such that the modeled M2 water levels match observations following \cite{Dijkstra et al. (2017)}. In our coupled flocculation-sediment transport model, we have three additional calibration parameters:

1. floc break-up parameter \( k_B \),
2. erosion parameter \( M \), and
3. river discharge \( Q \).

The iFlow model approximates the freshwater inflow by a subtidal discharge. In reality, the discharge shows a significant temporal variation, and observed average discharge is not always representative of average sediment transport. For example, the standard deviation of the averaged river discharge corresponding to our winter case is equal to \( 92 \) m\(^3\) s\(^{-1}\) \cite{Waterinfo.be, 2019}. To improve the correspondence between model output and observed sediment distribution, we included \( Q \) as a calibration parameter.

We run our model for various values of the calibration parameters \( k_B \in [2400, ..., 8000] \) s\(^{1/2}\) m\(^{-2}\), \( M \in [0.1, ..., 4] \times 10^{-3} \) s m\(^{-1}\), and \( Q \in [172, ..., 340] \) m\(^3\) s\(^{-1}\) and compare the model result to the measurements for each setting to find the best parameter values. The choice of the range in \( k_B \) and \( M \) is based on scaling (see Table 2.C.1) and observations \cite{Zhu et al., 2017}, respectively. The range of \( Q \) represents from the 60th until 90th percentile of the observations corresponding to our winter case \cite{Waterinfo.be, 2019}.
To objectively compare the model output to the measurements, we construct a cost function, which is based on a statistical two-tailed $t$ test. For each SPM data point location $(x,z)$, we compute whether the following test statistic:

$$t_{score} = \frac{< c_{meas.}(x, z) > - < c_{model}(x, z) >}{\text{std} / \sqrt{n}},$$

has a $t$ distribution with $n - 1$ degrees of freedom. If found to be true, we accept that the model output and data are statistically equal at location $(x,z)$. Here, $< c_{model}(x, z) >$ and $< c_{meas.}(x, z) >$ are the model subtidal suspended sediment concentration output and time-averaged measured SPM concentration at location $(x,z)$ respectively, std is the standard deviation of the measured SPM concentration at location $(x,z)$ and $n$ is the total number of data points at location $(x,z)$. We choose a significance level $\alpha = 0.05$. Last, we define a cost function value as

$$\text{cost} = 1 - \frac{N_{equal}}{N_{loc}},$$

where $N_{equal}$ is the number of locations at which the suspended sediment concentration model output and data are statistically equal and $N_{loc}$ is the total number of locations for which we have data. Consequently, the cost function output is a value between 1 (mismatch of model output and data) and 0 (perfect match of model output and data).

Evaluated by our cost function, we obtain an optimal parameter set $k_B = 5600 \, s^{1/2} \, m^{-2}$, $M = 0.00336 \, s \, m^{-1}$, and $Q = 233 \, m^3 \, s^{-1}$. The cost function output for various $k_B$, $M$, and $Q$ is presented in Fig. 2.4. We acquire a higher discharge of $233 \, m^3 \, s^{-1}$ in winter when compared with the time-averaged value of $172 \, m^3 \, s^{-1}$. This larger discharge is expected because of the asymmetry in flushing and buildup of an ETM: the impact of high river discharges on suspended sediment concentration is larger than the impact of low river discharges. Indeed, Brouwer et al. (2018) showed that the time scale of building up an ETM is two orders of magnitude larger ($\sim 10^2$ days) than the flushing of an ETM ($\sim 10^0$ days).

\section*{2.4 Results}

To analyze the relative importance of flocculation on large-scale sediment transport in the Scheldt estuary, we compare the modeled suspended sediment distribution and the different terms contributing to net sediment transport with and without flocculation. Next, we apply a sensitivity analysis of the calibration parameters $Q$ and $k_B$ when including flocculation and compare the results to those without flocculation.
2.4.1 Impact of flocculation on large-scale suspended sediment distribution

Application of the model to the winter case in the Scheldt estuary and model calibration results in the model output presented in Fig. 2.5.

Figure 2.5a shows the result of the subtidal suspended sediment concentration with flocculation. Figure 2.5b shows the long-term, time-averaged suspended sediment concentration dataset for winter in 2015-2017. These observations agree with SPM samples simultaneously taken at the water surface, which are presented in Cox et al. (2019). When we include flocculation (Fig. 2.5a), the model output complies both quantitatively and qualitatively with observations (Fig. 2.5b). The location and intrusion of the ETM are well captured. The model shows typical suspended sediment concentrations of 100-300 mg L$^{-1}$, which agrees with observations.

Figure 2.5c shows the model output of the leading-order settling velocity corresponding to the case with flocculation. Qualitatively this figure corresponds to the suspended sediment concentration in Fig. 2.5a. We expected this because we showed that the leading-order settling velocity scales linearly to the suspended sediment concentration [see Eq. (2.14)]. The distribution of $w_0$ is only (slightly) altered by the imposed longitudinal variation in shear rate $G$ and aggregation parameter $k_A$ [see Eqs. (2.7) and (2.8)]. Hence, the dependence of $w_0$ on $c_0$ is the most important factor affecting $w_0$.

Figure 2.5d shows the subtidal suspended sediment concentration without including flocculation, using a constant $w_s$ and the same river discharge as used in Fig. 2.5a. In the simplest model case in which we excluded flocculation, a constant $w_s$ of 3 mm s$^{-1}$ yields the best comparison to observations (Fig. 2.5b). This value corresponds to the value of the settling velocity found near the ETM (Fig. 2.5c). When we exclude flocculation (Fig. 2.5d), the model output qualitatively agrees with observations; we acquire an ETM at 60-80 km from the mouth and obtain more sediment near the bed than at the water surface. However, quantitatively the suspended sediment concentrations are up to a factor three too low. Furthermore, the intrusion of the ETM is too small. Note that our results are different from Dijkstra et al. (2019a) and Brouwer et al. (2018), since they only considered much lower discharges than what is considered here. Because we are interested in the individual impact of flocculation on the suspended sediment distribution, we did not recalibrate parameters $M$ and $Q$ for the case excluding flocculation. When these parameters are recalibrated, $M = 0.004$ s m$^{-1}$ and $Q = 172$ m$^3$ s$^{-1}$, resulting in a cost function value of 0.87. The modeled suspended sediment concentrations are still too low, although slightly higher than the previous case due to a decrease of $Q$. A further increase of $M$ does not increase the suspended sediment concentrations because we are in a suspended sediment supply-limited condition [see Brouwer et al. (2018)], that is, all sediment is eroded from the bed during a tidal cycle. This implies that the amount of sediment trapped, and not the erosion from the bed, is the limiting factor for the observed suspended sediment concentrations. Sediment trapping is related to net sediment transport processes, which are clearly less efficient in importing sediment when flocculation is not taken into account (see next section).
Figure 2.5: (a) Model output of the subtidal suspended sediment concentration (mg L\(^{-1}\)) with flocculation included. The optimal calibration parameters are \(k_B = 5600 \text{ s}^{1/2} \text{ m}^{-2}\), \(M = 0.00336 \text{ s m}^{-1}\), and \(Q = 233 \text{ m}^3 \text{ s}^{-1}\). The vertical dashed line depicts the intrusion limit of the observed suspended sediment concentration (i.e., \(c \lesssim 50 \text{ mg L}^{-1}\)).

(b) The long-term, time-averaged suspended sediment concentration dataset (mg L\(^{-1}\)).

(c) Leading-order, subtidal settling velocity \(w_{s0}\) (mm s\(^{-1}\)) with flocculation included.

(d) Model output of the subtidal suspended sediment concentration (mg L\(^{-1}\)) without including flocculation.
2.4.2 Impact of flocculation on net sediment transport

In this section, we focus on the underlying sediment-transport mechanisms that result in sediment transport and the buildup of an ETM presented in Fig. 2.5. For this, we introduce the advective net transport capacity \( T \), which equals the net redistribution of a (longitudinally) uniform layer of sediment on the bed (Dijkstra et al., 2019b). The net transport capacity only depends on hydrodynamic and sediment features and not on the location of the ETM. Therefore, it is a suitable measure to analyze the impact of various mechanisms on the total net sediment transport (Dijkstra et al., 2019a,b).

Figure 2.6 shows the six most important transport mechanisms contributing to net transport capacity with (Fig. 2.6a) and without (Fig. 2.6b) flocculation together with the total net transport capacity \( T_{\text{total}} \). This total net transport capacity \( T_{\text{total}} \) is the sum of all separate net transport capacities. When \( T_{\text{total}} = 0 \) and \( T_{\text{total}} \) changes sign from positive to negative (in land-inward direction), we have a convergence point. Here, we expect sediment to accumulate and thus the formation of an ETM. When comparing Figs. 2.6a and 2.6b we observe that the convergence point at approximately 80 km in the case with flocculation (Fig. 2.6a) shifted seaward to approximately 70 km when flocculation is not considered (Fig. 2.6b). In both cases, a significant decrease in total net transport capacity is present at 95 km, which is due to the Rupel tributary, resulting in downstream net sediment transport. Figure 2.6a also shows that the additional mechanism related to flocculation, \( w_1^s c_0^0 \) in Eq. (2.21), is always positive and consequently results in land-inward net sediment transport. Furthermore, it is the dominant mechanism at the mouth.

By comparing Figs. 2.6a and 2.6b it is clear that the contributions of the other transport mechanisms changed when flocculation is included. This is due to the spatial and temporal (M4) variations in the leading-order settling velocity \( w_0^s \), which were absent in the case without flocculation. As mentioned earlier, this alteration of the subtidal and M4 tidal signal in \( w_0^s \) impacts all contributions to the net sediment transport \( T \) [see Eqs. (2.19) and (2.20)].

2.4.3 Sensitivity analysis of floc break-up parameter \( k_B \) and total freshwater discharge \( Q \)

A sensitivity analysis of the floc break-up parameter \( k_B \) and freshwater discharge \( Q \) allows us to compute the impact of the magnitude of \( k_B \) and \( Q \) on the formation of ETM. To stress the importance of spatially varying \( w_s \), we show the sensitivity of ETM formation to \( k_B \) with flocculation only varying in space, not in time. We also compare the case with and without flocculation to calculate the impact of space and time dependence in \( w_s \) on the emergence of ETM. Finally, we vary the constant settling velocity \( w_0^s \) in the case without flocculation to assess the sensitivity of the ETM characteristics for this parameter.

Figure 2.7 shows the sensitivity of varying \( k_B \) (\( Q = 233 \) m\(^3\) s\(^{-1}\) and \( M = 0.00336 \) s m\(^{-1}\)) to depth-averaged subtidal suspended sediment concentration \( < \tilde{c} > \). In Fig.
Figure 2.6: Depth-averaged subtidal suspended sediment concentration $< \bar{c} >$, total advective net transport capacity $T_{\text{total}}$, and the six main hydrodynamical contributions to the total net transport capacity as a function of distance from the mouth (km) of our case study in the Scheldt estuary, showing results (a) with and (b) without flocculation included.
Figure 2.7: Depth-averaged subtidal suspended sediment concentration \(< \bar{c} > (g L^{-1})\) for varying floc break-up parameter \(k_B\) \((Q = 233 m^3 s^{-1} and M = 0.00336 s m^{-1})\) in the case with flocculation. The black dashed lines depict the locations of the ETM. (a) Result with time dependence in \(w_s\). (b) Result in which we excluded the time dependence in \(w_s\).

Figure 2.7 shows that the magnitude of \(k_B\) not only determines the location and intensity of ETM, but also the number of ETM. In Fig. 2.7a for \(k_B\) values larger than a critical value of 8000 s\(^{1/2}\) m\(^{-2}\), no ETM is found. For values between 4000 s\(^{1/2}\) m\(^{-2}\) and 8000 s\(^{1/2}\) m\(^{-2}\), one ETM is found at a fixed location. For values between 2400 s\(^{1/2}\) m\(^{-2}\) en 4000 s\(^{1/2}\) m\(^{-2}\), two ETM are found, with the most downstream ETM moving downstream with decreasing \(k_B\). This behavior is strongly linked to the temporal variability of settling velocities introduced by the flocculation process. Indeed, Fig. 2.7b shows the impact on ETM formation when only spatial variability in \(w_s\) is incorporated. In that case, ETM formation only occurs at significantly smaller \(k_B\) values \((\approx 3000 s^{1/2} m^{-2})\), while multiple ETM are not observed in the domain. Moreover, modeled suspended sediment concentrations are much lower when ignoring temporal variations in \(w_s\).

Figure 2.8 shows the impact of varying river discharge \(Q\) \((M = 0.00336 s m^{-1})\) on the depth-averaged subtidal suspended sediment concentration \(< \bar{c} >\) with flocculation (Fig. 2.8a, \(k_B = 5600 s^{1/2} m^{-2}\)) and without flocculation (Fig. 2.8b, \(w_s^0 = 3 mm s^{-1}\)). We keep \(Q\) at typical winter values. Similar to the sensitivity in \(k_B\), the magnitude of \(Q\) not only determines the location and intensity of ETM, but also the number of ETM. When flocculation is incorporated (Fig. 2.8a), we have one ETM for values between approximately 200 and 340 m\(^3\) s\(^{-1}\), with suspended sediment concentrations that compare
well with observations (Fig. 2.5b). For values between approximately 170 and 200 m$^3$ s$^{-1}$, we find an additional ETM at approximately 100-120 km. This second ETM has been observed in the Scheldt estuary when discharges are low (Cox et al., 2019). An ETM appearing in the tidal freshwater region during lower discharges was observed in various estuaries (Uncles et al., 2006; Allen et al., 1980). Both ETM move downstream with increasing $Q$. In the model with fixed settling velocity, only a single ETM is found, which shifts again downstream with increasing $Q$ (Fig. 2.8b). Moreover, the suspended sediment concentrations obtained are low. Consequently, flocculation is important to obtain quantitatively realistic results for relatively large and observed freshwater discharges $Q$.

Figure 2.9 shows the impact of varying constant settling velocity $w_s$ on $<\tau>$ in case effects of flocculation are not considered ($\dot{Q} = 233$ m$^3$ s$^{-1}$ and $\dot{M} = 0.00336$ s m$^{-1}$). The magnitude of the settling velocity determines the existence of an ETM. We need a minimal constant settling velocity of approximately 1.8 mm s$^{-1}$ to obtain one ETM. The optimal choice of the constant settling velocity is approximately 3 mm s$^{-1}$, resulting in the best comparison to observations. A settling velocity $w^0_s = 3$ mm s$^{-1}$ corresponds to a typical value near the ETM (Fig. 2.5c). Moreover, the depth-averaged subtidal suspended sediment concentrations are relatively low compared to the data, for all $w^0_s$. At $w^0_s = 3$ mm s$^{-1}$, although the locations of the ETM are equal, the suspended sediment concentrations are still approximately one-third of those found with time-independent flocculation (Fig. 2.7b), stressing the importance of spatially varying $w_s$. We cannot resolve the modeled low concentrations (cf. Fig. 2.8b) by altering $w^0_s$. Consequently, flocculation is required to acquire quantitatively realistic concentrations for large freshwater discharges in winter.
2.5 Discussion

Our results show that flocculation promotes land-inward net sediment transport in the Scheldt estuary. Relative to the iFlow model without flocculation, the model with flocculation is less sensitive to freshwater discharge. As a result, we still obtain an ETM for high freshwater discharges, which complies with observations (Cox et al., 2019). In addition, the iFlow model without flocculation results in depth-averaged suspended sediment concentrations that are typically too low for large freshwater discharge, stressing the importance of a temporal and spatial dependency of the settling velocity $w_s$ caused by flocculation.

2.5.1 Indications of spatial and temporal variations of the settling velocity in the Scheldt estuary

Observations in the Scheldt estuary confirm both this temporal dependence in $w_s$, which generates net sediment transport and the modeled magnitude of $w_s$. Although direct settling velocity measurements are scarce, Manning et al. (2007) measured an averaged settling velocity of macroflocs of 3.9 mm s$^{-1}$ in the ETM of the Scheldt at the entrance of Deurganckdok, approximately 62 km from Vlissingen. When we compare these measured settling velocities to the model output, we find that they more or less comply (Fig. 2.5c). Besides direct settling velocity measurements in the Scheldt estuary, Fettweis and Baeye (2015) observed a strong M2 and M4 tidal signal in floc size and settling velocity in the Southern North Sea, which complies with our model results [Eqs. (2.14) and (2.17)]. Furthermore, Schwarz et al. (2017) measured in situ settling velocities over one tidal cycle in the main channel of the Sieperda March (near the Dutch-Belgian border in Fig. 2.3) using both the Stokes formulation and the Reynolds-flux method. They also found
a distinct M4 tidal signal in settling velocity and floc size, which again agrees with our model results.

2.5.2 Flocculation resulting in land-inward net sediment transport

We showed that flocculation alters the sediment transport mechanisms and generates an additional and important sediment transport mechanism (Figs. 2.6a and 2.6b). Our method enabled a systematic analysis showing that flocculation, and more specifically, the M2 tidal signal in the settling velocity, results in land-inward net sediment transport (solid black line in Fig. 2.6a). Our results comply with Winterwerp (2011) who showed that including flocculation results in land-inward net sediment transport in the Ems River using a 1D vertical model. Xu et al. (2010) also concluded that flocculation results in land-inward sediment transport and promotes accumulation of suspended sediment by performing an idealized 2D model study in the Upper Chesapeake Bay. In general, the direction of net sediment transport depends on the tidal phase difference between flow velocity, suspended sediment concentration, and settling velocity, which might differ for various estuaries and is affected by both the spatial and temporal variations of the settling velocity caused by flocculation.

To estimate a condition for which flocculation results in land-inward net sediment transport, we only consider the M2 tidal signal in the settling velocity $w_{s12}^2$, which we showed can have a significant impact on net sediment transport. When the longitudinal water velocity $u$ (in land-inward direction) is in phase with the suspended sediment concentration $c_{floc}^{12}$ generated by $w_{s12}^2$, we have the condition for maximum land-inward sediment transport related to the flocculation contribution (see Fig. 2.10).

To meet this condition, it is found that $w_{s12}^2$ has to be shifted by a $-\pi/2$ phase relative to $u$, meaning that the M2 tidal signal in the settling velocity peaks at slack tide from flood to ebb (see Fig. 2.10). This condition holds assuming that the water column is fairly well mixed and $c_{floc}^{12}$ and $c_{04}^{04}$ are relatively small in comparison with $c_{12}^{12}$ and $c_{00}^{00}$. This condition is modified in other cases. Our reasoning is generalized in Appendix 2.E for $c_{04}^{04}$ not relatively small when compared with $c_{00}^{00}$ and general (i.e., not restricted to maximal) land-inward sediment transport.

2.5.3 The potential impact of flocculation on sediment transport in the Scheldt estuary

Our model results show that the floc break-up parameter $k_B$ determines the position, intensity, and existence of ETM (Fig. 2.7), which complies with Xu et al. (2010) who found that particle stickiness (cf. $k_B$) can have a significant influence on sediment trapping, especially when the stickiness is small (cf. $k_B$ large). As mentioned earlier, various factors such as salinity (Edzwald et al., 1974) and biotic sticky substances (Fettweis...
Figure 2.10: Illustration of the optimal condition for land-inward sediment transport with regard to the phase differences between the longitudinal water velocity $u$, the M2 tidal signal in the settling velocity $w_{12}^{s}$, and the suspended sediment concentration due to flocculation $c_{floc}^{12}$ (generated by $w_{12}^{s}$). The phase difference between $u$ and $w_{s}^{12}$ has to be $-\pi/2$, meaning $w_{s}^{12}$ has to peak at slack tide from flood (white zone) to ebb (blue zone). Because of inertia effects and vertical suspended sediment stratification, this phase condition results in $u$ and $c_{floc}^{12}$ being in phase, resulting in maximal land-inward sediment transport due to flocculation.

In our case study, the direct impact of salinity on flocculation and resulting sediment transport is relatively small. This complies with [Einstein and Krone (1962)] who concluded that variations of salinity in most of an estuary have only a small effect on the bond strength. Moreover, the experiments of [Edzwald et al. (1974)] show a relatively low impact in comparison with the minimal aggregation coefficient (cf. $k_B$) and saturation at relatively low salinity. Salinity might also have an indirect impact on flocculation through, for example, the production of biotic sticky substances [Alldredge et al., 1993; Bar-Zeev et al., 2015], which is only implicitly included through calibration of $k_B$.

Over the last two decades, the water quality has significantly improved in the Scheldt estuary because of the implementation of wastewater treatment in Brussels in 2006 [Brion et al., 2015; Cox et al., 2019]. Cox et al. (2019) suggested that changes in water quality might result in a large-scale impact on suspended sediment distribution through its influence on flocculation. Our flocculation model can be used to quantify the impact of water quality improvement and verify the hypothesis of Cox et al. (2019) and supplies a tool to systematically assess the mechanisms that explain the changes to the sediment transport. The main unknown for this is the relation between water quality and $k_B$, and more research is needed to establish practical relations for this.
2.5.4 Parameter variations in the Lagrangian flocculation model

To determine the individual impact of flocculation on the large-scale suspended sediment distribution, we compared a sediment transport model using a constant settling velocity and using the complex Lagrangian flocculation model of Winterwerp (2002). As a first step, we kept most parameters fixed in the Lagrangian flocculation model following Winterwerp (2002).

Over the last decade, various authors extended this flocculation model. For example, Maggi (2009) extended the model by separating the floc volume in a mineral and biomass fraction. More recently, Xu and Dong (2017) were able to model floc size dynamics more accurately when assuming that the fractal dimension \( n_f \) follows a normal distribution. As a final example, Kuprenas et al. (2018) implemented a dependence of floc size \( D_f \) and shear rate \( G \) in the calibration parameter \( q \) to correct for an overestimation of floc size at large shear rates (order of 50 s\(^{-1}\)).

In further research, the same scaling and perturbation procedure presented in this paper can be applied to extended versions of the Lagrangian model of Winterwerp. For example, by assuming constant parameter values (e.g., \( n_f = 2 \) and \( q = 0.5 \)), we found a linear relationship between the settling velocity and suspended sediment concentration. Different parameter values might result in a nonlinear relationship. An example of such a relation is given by van Leussen (1994), who proposed an at equilibrium relation between settling velocity, suspended sediment concentration, and shear rate: \( w_s = K_v L c_a \left( \frac{1 + a_v L G}{1 + b_v L G^2} \right) \), in which \( a_v L, b_v L, K_v L, \) and \( m_v L \) are empirically determined parameters. Such a formulation is similar to ours in that it depends on \( c \) and \( G \) but with a somewhat different relation, which may lead to a modification of the results. But, further discussion of this is out of the scope of the present paper.

2.6 Conclusions

In this paper, we coupled a flocculation model and a sediment transport model to acquire insight into the impact of flocculation on large-scale sediment transport in a tide-dominated estuary. The combination of this flocculation model and the iFlow model allowed us to identify the relative importance of individual processes for water flow and sediment transport (e.g., river, tide, density forcing, advection). We employed a perturbation approach to gain insight into the highly complex coupled equations that describe flocculation. The result reveals a simple linear relationship between \( w_s \) and \( c \), both at leading and first order.

We applied our framework to a winter case in the Scheldt estuary. We showed that the spatial and temporal variations of \( w_s \) due to flocculation are essential to reproduce observed suspended sediment concentrations. We were able to identify the impact of flocculation on individual transport mechanisms. We showed that flocculation alters most of the dominant transport mechanisms (e.g., river and tidal return flow) by introducing
both a spatial and temporal dependence in $w_s$.

To further investigate the impact of flocculation on large-scale sediment transport, we carried out a sensitivity analysis in which we showed that the magnitude of the floc break-up parameter $k_B$ and total freshwater discharge $Q$ determine the existence, intensity, and number of ETM.

Although we found examples in the literature of several other estuaries where flocculation is thought to promote sediment import, we were able to show that flocculation can also theoretically promote sediment export, depending on the phase difference between the tidal flow and suspended sediment concentration. Therefore, the effect of flocculation on the large-scale sediment transport in other estuaries needs to be assessed carefully based on local conditions.

List of symbols

**Latin**

$A^0$ Water level amplitude at $x = 0$ corresponding to the M2 tide

$A^1$ Water level amplitude at $x = 0$ corresponding to the M4 tide

$a_{vL}$ Parameter in the empirical formulation of van Leussen (1994) for the relation between $w_s$ and $c$

$B$ Width of the estuary

$b_{vL}$ Parameter in the empirical formulation of van Leussen (1994) for the relation between $w_s$ and $c$

$c$ Suspended sediment concentration

$c^0$ Leading-order suspended sediment concentration

$c^{00}$ Subtidal component of the leading-order $c$

$c_{floc}^{00}$ Flocculation-induced $c^{00}$

$c_{floc}^{04}$ Flocculation-induced $c^{04}$

$\hat{c}_{floc}^{00}$ Complex amplitude of $c^{00}$

$\hat{c}_{floc}^{04}$ Complex amplitude of $c^{04}$

$c^{10}$ Subtidal component of the first-order $c$

$c_{floc}^{10}$ Flocculation-induced $c^{10}$

$c_{floc}^{14}$ Flocculation-induced $c^{14}$

$\hat{c}_{floc}^{10}$ Complex amplitude of $c^{10}$

$c^{12}$ Component of the first-order $c$ corresponding to the M2 tide

$c_{floc}^{12}$ Flocculation-induced $c^{12}$

$c_{floc}^{14}$ Flocculation-induced $c^{14}$

$\hat{c}_{floc}^{12}$ Complex amplitude of $c^{12}$

$\hat{c}_{floc}^{14}$ Complex amplitude of $c^{14}$
\( \hat{c}^{14} \) Complex amplitude of \( c^{14} \)
\( \langle c_{\text{meas.}} \rangle \) Time-averaged measured SPM concentration
\( \langle c_{\text{model}} \rangle \) Modeled subtidal suspended sediment concentration
\( \langle c \rangle \) Depth-averaged subtidal suspended sediment concentration
\( c_{\text{sea}} \) Depth-averaged subtidal suspended sediment concentration at \( x = 0 \)
\( D_e \) Equilibrium floc size
\( D_f \) Floc size
\( D_p \) Primary particle size
\( f_s \) Shape factor
\( G \) Shear rate
\( g \) Gravitational acceleration
\( H \) Magnitude of the z-coordinate of the river bed
\( K_h \) Horizontal diffusivity coefficient
\( K_{vL} \) Parameter in the empirical formulation of van Leussen (1994) for the relation between \( w_s \) and \( c \)
\( K_v \) Vertical eddy diffusivity coefficient
\( K_0^v \) Leading order vertical eddy diffusivity coefficient
\( k_A' \) Aggregation parameter
\( k_{A\min} \) Nondimensional minimal aggregation coefficient
\( k_B \) Break-up parameter
\( M \) Erosion parameter
\( m_{vL} \) Parameter in the empirical formulation of van Leussen (1994) for the relation between \( w_s \) and \( c \)
\( N \) Number of flocs per unit volume
\( N_{\text{equal}} \) Number of locations at which the suspended sediment concentration model output and observations are statistically equal
\( N_{\text{loc}} \) Total number of locations at which we have SPM observations
\( n \) Number of measured SPM concentrations
\( n_f \) Fractal dimension
\( O \) Order of
\( p \) Calibration parameter in the flocculation model
\( q \) Calibration parameter in the flocculation model
\( Q \) Total freshwater discharge
\( R \) Reference level of the water surface elevation
\( \text{Re}_f \) Floc Reynolds number (i.e., \( w_s D_f / \nu \))
\( S \) Salinity
\( S_1 \) Flocculation salinity sensitivity calibration parameter
\( S_2 \) Flocculation salinity sensitivity calibration parameter
\( s_{f_0} \) Bed roughness coefficient
\( s_{\text{sea}} \) Salinity boundary condition at the mouth
\( \text{std} \) Standard deviation of the measured SPM concentration
\( T \) Net sediment transport
\( T_{\text{floc}} \) Flocculation-induced net sediment transport
\( T_{\text{total}} \) Total net transport capacity
\( t \) Time
\( t_{\text{score}} \) Student’s \( t \) statistic
\( u \) Water flow velocity in the longitudinal dimension
\( u_s \) \hspace{1cm} \text{Shear velocity}

\( u^0 \) \hspace{1cm} \text{Leading-order water flow velocity in the longitudinal dimension}

\( u^{00} \) \hspace{1cm} \text{Subtidal component of the leading-order } u

\( \hat{u}^{00} \) \hspace{1cm} \text{Complex amplitude of } u^{00}

\( u^{02} \) \hspace{1cm} \text{Component of the leading-order } u \text{ corresponding to the M2 tide}

\( \hat{u}^{02} \) \hspace{1cm} \text{Complex amplitude of } u^{02}

\( u^{04} \) \hspace{1cm} \text{Component of the leading-order } u \text{ corresponding to the M4 tide}

\( \hat{u}^{04} \) \hspace{1cm} \text{Complex amplitude of } u^{04}

\( u^1 \) \hspace{1cm} \text{First-order velocity in the longitudinal dimension}

\( u^{10} \) \hspace{1cm} \text{Subtidal component of the first-order } u

\( \hat{u}^{10} \) \hspace{1cm} \text{Complex amplitude of the subtidal component of the first-order } u

\( u^{12} \) \hspace{1cm} \text{Component of the first-order } u \text{ corresponding to the M2 tide}

\( \hat{u}^{12} \) \hspace{1cm} \text{Complex amplitude of } u^{12}

\( u^{14} \) \hspace{1cm} \text{Component of the first-order } u \text{ corresponding to the M4 tide}

\( \hat{u}^{14} \) \hspace{1cm} \text{Complex amplitude of } u^{14}

\( w \) \hspace{1cm} \text{Water flow velocity in the vertical dimension}

\( w^0 \) \hspace{1cm} \text{Leading-order water flow velocity in the vertical dimension}

\( w^1 \) \hspace{1cm} \text{First-order water flow velocity in the vertical dimension}

\( w_s \) \hspace{1cm} \text{Settling velocity of suspended sediment}

\( w^0_s \) \hspace{1cm} \text{Leading-order settling velocity of suspended sediment}

\( w^{02}_s \) \hspace{1cm} \text{Component of the leading-order } w_s \text{ corresponding to the M2 tide}

\( w^{04}_s \) \hspace{1cm} \text{Component of the leading-order } w_s \text{ corresponding to the M4 tide}

\( w^{0, \text{init.}}_s \) \hspace{1cm} \text{Initial leading-order } w_s \text{ in the iteration procedure}

\( w^{0, \text{old}}_s \) \hspace{1cm} \text{Previous value of the leading-order } w_s \text{ in the iteration procedure}

\( w^1_s \) \hspace{1cm} \text{First-order settling velocity of suspended sediment}

\( w^{10}_s \) \hspace{1cm} \text{Subtidal component of the first-order } w_s

\( w^{12}_s \) \hspace{1cm} \text{Component of the first-order } w_s \text{ corresponding to the M2 tide}

\( w^{14}_s \) \hspace{1cm} \text{Component of the first-order } w_s \text{ corresponding to the M4 tide}

\( w^{1, \text{init.}}_s \) \hspace{1cm} \text{Initial first-order } w_s \text{ in the iteration procedure}

\( w^{\text{min.}}_s \) \hspace{1cm} \text{Minimum settling velocity of suspended sediment}

\( x \) \hspace{1cm} \text{Spatial coordinate in the longitudinal dimension}

\( x_{\text{sal}} \) \hspace{1cm} \text{Calibration parameter in postulated tanh salinity distribution in winter}

\( x_{\text{sal}}^L \) \hspace{1cm} \text{Calibration parameter in postulated tanh salinity distribution in winter}

\( z \) \hspace{1cm} \text{Spatial coordinate in the vertical dimension}

\( z_1 \) \hspace{1cm} \text{Distance in the logarithmic layer above the river bed}

\( \tilde{z}_1 \) \hspace{1cm} \text{distance in the logarithmic layer above the river bed}

**Greek**

\( \beta \) \hspace{1cm} \text{Parameter defined in Table 2.1}

\( \gamma \) \hspace{1cm} \text{Parameter defined in Table 2.1}

\( \partial_t \) \hspace{1cm} \text{Partial derivative to } t

\( \partial_x \) \hspace{1cm} \text{Partial derivative to } x

\( \partial_z \) \hspace{1cm} \text{Partial derivative to } z

\( \epsilon \) \hspace{1cm} \text{Small perturbation number (i.e., } \xi/H \ll 1)
Appendix 2.A: salinity dependence

Following Warner et al. (2005), we fit the observed salinity data to the following postulated salinity distribution of our winter case in the Scheldt estuary:

$$\frac{s_{\text{sea}}}{2} \left(1 - \tanh \frac{x - x_{c}^{\text{Sal.}}}{x_{L}^{\text{Sal.}}} \right),$$

with $s_{\text{sea}}$ being the salinity boundary condition at the mouth and $x_{c}^{\text{Sal.}}$ and $x_{L}^{\text{Sal.}}$ being further undefined calibration parameters. Figure 2.A.1a shows the data fit.

Figure 2.A.1: (a) Measured salinity corresponding to our case study in the Scheldt estuary and corresponding data fit using Eq. (2.25). (b) Stability value $\alpha'$ as a function of salinity based on Edzwald et al. (1974).
et al. (1974) calculated the salinity dependence for stability value $\alpha'$ which is defined as

$$\dot{\alpha}'_{\text{aggregation}} = \frac{-4\alpha'\phi GN}{\pi},$$

(2.26)

where $\phi$ is the volumetric concentration, $G$ is the shear rate, and $N$ is the number concentration of flocs. Winterwerp and van Kesteren (2004) showed that

$$\phi = f_s ND^2_f,$$

(2.27)

with $f_s$ being the floc shape factor and $D_f$ being the floc size. Consequently, in identification with Eq. (2.6) and assuming the floc shape is spherical ($f_s = \pi/6$), we have

$$k_A' = \frac{2\alpha'}{3}.$$  

(2.28)

The latter relation allows us to use the salinity dependence of $\alpha'$ for $k_A'$. We postulate the following salinity $S$ dependence:

$$\alpha' = \left\{ \alpha_{\text{min}}' + S_3 [1 + \tanh (S(x) - S_2)] \right\},$$  

(2.29)

and we fit the latter function to the averaged data presented in Edzwald et al. (1974), which is shown in Fig. 2A.1b. We acquire $\alpha_{\text{min}}' = 0.0904$, $S_2 = 4.085\%$, and $S_3 = 0.03401$. We showed that net sediment transport only depends on the ratio of $k_A'$ and $k_B$ and not on the individual parameters. We can thus decide to make $k_A'$ solely a function of salinity $S$ and keep $k_B$ fixed without restricting the generality of our results:

$$k_A'(x) = k_{A\text{min}}' \left\{ 1 + S_1 [1 + \tanh(S(x) - S_2)] \right\},$$  

(2.30)

with $k_{A\text{min}}'$ being the minimal aggregation parameter. Combination of Eqs. (2.28) and (2.30) yields

$$S_1 = \frac{2}{3} \frac{S_3}{k_{A\text{min}}'}.$$  

(2.31)

### Appendix 2.B: sediment equations with ordering

The width-averaged sediment mass balance equation following Chernetsky et al. (2010) and Dijkstra et al. (2017) reads as

$$\partial_t c + u \partial_x c + w \partial_z c - \partial_x (w_s c + K_v \partial_x c) - \partial_z (K_h \partial_z c) = 0,$$

(2.32)

where $c$ is the suspended sediment concentration, $u$ and $w$ are the water velocity in the $x$ and $z$ direction, $w_s$ is the settling velocity, $K_v$ is the constant vertical eddy diffusivity.
coefficient, and $K_h$ is the constant horizontal diffusivity coefficient. At the water surface, we require that no sediment particles enter or leave the domain:

$$w_s c + \partial_z (K_{\nu} c) = 0 \quad \text{at} \quad z = R + \xi,$$

(2.33)

with $R$ being the reference level and $\xi$ being the water surface elevation. At the bed, we require

$$w_s c + K_{\nu} \partial_z c = D - E \quad \text{at} \quad z = -H,$$

(2.34)

where $D$ and $E$ are the deposition and erosion of sediment defined as

$$D = w_s c \quad \text{and} \quad E = M |\tau_b| f(a),$$

(2.35) (2.36)

where $M$ is the erosion parameter, $\tau_b$ is the shear stress at the bed, and $f(a)$ is the erodibility.

To apply ordering and perturbation theory, we first have to compute the order of each term in the differential equation and boundary conditions. For this, we use typical scales for each variables following Chernetsky et al. (2010):

$$\begin{align*}
\partial_t \tilde{c} + \frac{U}{\sigma L_{Ems}} \tilde{u} \partial_x \tilde{c} + \frac{U}{\sigma L_{Ems}} \tilde{w} \partial_z \tilde{c} \\
- \partial_z \left( \frac{W_s}{\sigma H_{0,Ems}} \tilde{w}_s c + \frac{K_{\nu}}{\sigma H_{0,Ems}} \tilde{K}_\nu \partial_z \tilde{c} \right) - \partial_z \left( \frac{K_h}{\sigma L_{Ems}^2} \partial_z \tilde{c} \right) = 0,
\end{align*}$$

(2.37)

where the tilde denotes dimensionless variables, $\mathcal{O}$ is the order of magnitude, $\sigma$ is the $M2$ tidal angular frequency, $U$ is the typical scale of the horizontal velocity of the $M2$ tide, $A_{M2}$ is the $M2$ tidal amplitude at seaward side, $H_{0,Ems}$ is the water depth at the mouth, $W_s$ is the typical settling velocity scale, $K_{\nu}$ is the typical vertical eddy diffusivity coefficient scale, and $L_{Ems}$ is the length of the Ems estuary. The typical scales following Chernetsky et al. (2010) are summarized in Table 2.B.1. The boundary conditions in dimensionless form read as

$$\begin{align*}
\frac{W_s}{\sigma H_0} \tilde{w}_s c + \frac{K_{\nu}}{\sigma H_0^2} \tilde{K}_\nu \partial_z \tilde{c} = 0 \quad \text{at} \quad \tilde{z} = \frac{R}{H_0} + \frac{A_{M2}}{H_0} \tilde{\xi} \quad \text{and} \\
\frac{K_{\nu}}{\sigma H_0^2} \tilde{K}_\nu \partial_z \tilde{c} = - \frac{\mathcal{E}/C}{\sigma H_0} \tilde{E} \quad \text{at} \quad \tilde{z} = -\tilde{H},
\end{align*}$$

(2.38) (2.39)

where $R$ is the water surface reference level, $\tilde{\xi}$ is the dimensionless water surface elevation, $\mathcal{E}$ is the erosion scale, and $C$ is the suspended sediment concentration scale.
Table 2.B.1: Typical scales of the variables used in the ordering and perturbation analysis by Chernetsky et al. (2010). The variables are deducted from both theoretical equations and measurements in the Ems estuary.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Scale</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>$H_{0,\text{Ems}}$</td>
<td>Averaged depth at seaward side</td>
<td>12.2 m</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$A_{M2}$</td>
<td>M2 tidal amplitude at seaward side</td>
<td>1.35 m</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$L_{\text{Ems}}$</td>
<td>Length Ems estuary</td>
<td>$63.7 \times 10^3$ m</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>Semidiurnal angular tidal frequency scale</td>
<td>$1.4 \times 10^{-4}$ s$^{-1}$</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$W_s$</td>
<td>Settling velocity scale</td>
<td>$2 \times 10^{-3}$ m s$^{-1}$</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$K_\nu K_\nu = K_\nu$</td>
<td>Vertical eddy diffusivity coefficient scale</td>
<td>$1.7 \times 10^{-2}$ m$^2$ s$^{-1}$</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$K_h$</td>
<td>Horizontal diffusivity coefficient scale</td>
<td>100 m$^2$ s$^{-1}$</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$U = \frac{\sigma A_{M2} E_{\text{Ems}}}{M}$</td>
<td>Typical horizontal velocity of the M2 tide</td>
<td>1 m s$^{-1}$</td>
<td>Balance of continuity equation</td>
</tr>
<tr>
<td>$W = \frac{H_{0,\text{Ems}} U}{L}$</td>
<td>Typical vertical velocity of the M2 tide</td>
<td>$1.9 \times 10^{-4}$ m s$^{-1}$</td>
<td>Balance of continuity equation</td>
</tr>
<tr>
<td>$J/C = \frac{\xi}{M}$</td>
<td>Ratio of typical erosion and suspended suspended sediment concentration scale</td>
<td>$1.4 \times 10^{-3}$ m s$^{-1}$</td>
<td>Balance of Eq. (2.44)</td>
</tr>
</tbody>
</table>

To apply perturbation theory, we write the solutions for $u$, $w$, $\xi$, and $c$ as a power series of a small parameter $\epsilon$ with $O(1)$:

\[
\begin{align*}
  u & = u^0 + u^1 + u^2 + O(\epsilon^3), \\
  w & = w^0 + w^1 + w^2 + O(\epsilon^3), \\
  \xi & = \xi^0 + \xi^1 + \xi^2 + O(\epsilon^3), \quad \text{and} \\
  c & = c^0 + c^1 + c^2 + O(\epsilon^3),
\end{align*}
\]  

(2.40)\-(2.43)

where $u^0$, $w^0$, $\xi^0$, and $c^0$ are assumed to be of leading order; $u^1$, $w^1$, $\xi^1$, and $c^1$ are of order $\epsilon$, and so on. Using the scaling, we acquire at leading order

\[
\begin{align*}
  \partial_t c^0 - \partial_z (w_s^0 c^0 + K_\nu^0 \partial_z c^0) = 0, \\
  w_s^0 c^0 + K_\nu^0 \partial_z c^0 = 0 \quad \text{at} \quad z = R, \quad \text{and} \\
  K_\nu^0 \partial_z c^0 = -E^0 \quad \text{at} \quad z = -H
\end{align*}
\]  

(2.44)\-(2.46)

and at first order

\[
\begin{align*}
  \partial_t c^1 - \partial_z (w_s^0 c^1 + K_\nu^0 \partial_z c^1) = \\
  -w_s^0 \partial_z c^0 - w_s^0 \partial_z c^0 + \partial_z (w_s^0 c^0) + \partial_z (K_\nu^0 \partial_z c^0), \\
  \underbrace{w_s^1 c^0 + K_\nu^0 \partial_z c^1_{\nu}}_{\partial_z c^0} = \\
  -w_s^1 c^0 - K_\nu^1 \partial_z c^0 - \underbrace{w_s^0 c^0_{\nu} + K_\nu^0 \partial_z c^0}_{\partial_z c^0} \xi^0 \quad \text{at} \quad z = R, \quad \text{and} \\
  K_\nu^0 \partial_z c^1 = -E^1 - K_\nu^1 \partial_z c^0 \quad \text{at} \quad z = -H,
\end{align*}
\]  

(2.47)\-(2.49)

in which we linearized around the reference level $R$ (i.e., we applied a Taylor expansion). Chernetsky et al. (2010) and Dijkstra et al. (2017) showed that $c^0$ has a subtidal and M4 tidal signal and $c^1$ has an M2 tidal signal given the assumptions listed above.
Appendix 2.C: scaling analysis and perturbation theory

Similarly, we scale the Winterwerp flocculation model shown in Eq. (2.12):

\[
\frac{1}{\gamma G_k B G^{1/2}(\gamma D_p^2)} W_s \tilde{w}_s \sigma C W_s^{-2} \left\{ \frac{U}{\sigma L} \partial_z \tilde{c} \tilde{w}_s^{-2} + \frac{U}{\sigma L} \partial_x [\tilde{u} \tilde{c} \tilde{w}_s^{-2}] \right\}
+ \partial_z \left\{ \frac{U}{\sigma L} \tilde{w} - \frac{W_s}{\sigma H_0} \tilde{w}_s \right\} - \frac{1}{L^2 \sigma} \partial_x [K_h \partial_x \tilde{c} \tilde{w}_s^{-2}] \right\}
\]

\[
= -k'_A D_p^{-2} L^{-1} \frac{\rho_s^{-1}}{\sigma} C \tilde{c} + W_s \tilde{w}_s - \frac{1}{\gamma D_p^2},
\]

\[
x \frac{1}{W_s} \tau W_s^{-2} \tilde{w}_s \frac{\sigma}{\sigma} \left\{ \frac{U}{\sigma L} \partial_z \tilde{c} \tilde{w}_s^{-2} + \frac{U}{\sigma L} \partial_x [\tilde{u} \tilde{c} \tilde{w}_s^{-2}] \right\}
+ \partial_z \left\{ \frac{U}{\sigma L} \tilde{w} - \frac{W_s}{\sigma H_0} \tilde{w}_s \right\} - \frac{1}{L^2 \sigma} \partial_x [K_h \partial_x \tilde{c} \tilde{w}_s^{-2}] \right\}
\]

\[
= \beta \frac{C}{W_s} \tilde{c} + \tilde{w}_s - \frac{1}{\gamma D_p^2 W_s},
\]

(2.50)

in which we used typical scales for the Scheldt and Ems estuary presented in Table 2.C.1 and \( \beta, \gamma, \) and \( \tau \) are defined in Table 2.1. To apply perturbation theory, we write the solution of \( w_s \) as a power series of a small parameter \( \epsilon \) with \( \mathcal{O}(1) \) following Chernetsky et al. (2010):

\[
w_s = w_s^0 + w_s^1 + w_s^2 + \mathcal{O}(\epsilon^3),
\]

(2.51)

(2.52)

where \( w_s^0 \) is assumed to be of leading order, \( w_s^1 \) is of order \( \epsilon \), and so on. Using the scaling from Eq. (2.50), we acquire at leading order in dimensional form the balance between
Table 2.C.1: Typical scales of the variables used in the ordering and perturbation analysis. The variables are deducted from both theoretical equations and measurements in the Scheldt and Ems estuary.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Scale</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>$H_0 \tilde{z} = z$</td>
<td>Averaged depth at seaward side</td>
<td>$1 \times 10^2$ m</td>
<td>Fig. 2.5b Scheldt estuary</td>
</tr>
<tr>
<td>$A_{M2}$</td>
<td>M2 tidal amplitude scale at seaward side</td>
<td>$1 \times 10^0$ m</td>
<td>Brouwer et al. (2018) Scheldt estuary</td>
</tr>
<tr>
<td>$D_p$</td>
<td>Primary particle size scale</td>
<td>$1 \times 10^{-5}$ m</td>
<td>Winterwerp (2002) Ems estuary</td>
</tr>
<tr>
<td>$k_A'$</td>
<td>Dimensionless floc aggregation parameter scale</td>
<td>$1 \times 10^{-1}$</td>
<td>Winterwerp (2000) Ems estuary</td>
</tr>
<tr>
<td>$L$</td>
<td>Length Scheldt estuary</td>
<td>$160 \times 10^3$ m</td>
<td>Brouwer et al. (2018) Scheldt estuary</td>
</tr>
<tr>
<td>$\sigma^{-1}f = t$</td>
<td>Semidiurnal angular tidal frequency scale</td>
<td>$1.4 \times 10^{-4}$ s$^{-1}$</td>
<td>Chernetsky et al. (2010)</td>
</tr>
<tr>
<td>$W_s \tilde{w}_s = w_s$</td>
<td>Settling velocity scale</td>
<td>$2 \times 10^{-3}$ m s$^{-1}$</td>
<td>Brouwer et al. (2018) and Manning et al. (2007) Scheldt estuary</td>
</tr>
<tr>
<td>$K_v \tilde{K}_v = K_v$</td>
<td>Vertical eddy diffusivity coefficient scale</td>
<td>$1.7 \times 10^{-2}$ m$^2$s$^{-1}$</td>
<td>Chernetsky et al. (2010) Ems estuary</td>
</tr>
<tr>
<td>$\rho_s$</td>
<td>Density of primary particles</td>
<td>$2.6 \times 10^{3}$ kg m$^{-3}$</td>
<td>Chernetsky et al. (2010) Ems estuary</td>
</tr>
<tr>
<td>$C \tilde{c} = c$</td>
<td>Suspended sediment concentration scale</td>
<td>$2 \times 10^{-1}$ kg m$^{-3}$</td>
<td>Fig. 2.5b Scheldt estuary</td>
</tr>
<tr>
<td>$G$</td>
<td>Shear rate scale</td>
<td>$2 \times 10^0$ s$^{-1}$</td>
<td>Eq. (2.53) Scheldt estuary</td>
</tr>
<tr>
<td>$k_B$</td>
<td>Dimensional floc break-up parameter scale</td>
<td>$5 \times 10^{5}$ s$^{-1/2}$ m$^{-2}$</td>
<td>Balance floc aggregation and break-up term</td>
</tr>
<tr>
<td>$U = \frac{\sigma A_{M2} L}{H_0}$</td>
<td>Typical horizontal velocity of the M2 tide</td>
<td>$2.3$ m s$^{-1}$</td>
<td>Balance depth-averaged continuity equation</td>
</tr>
</tbody>
</table>

Floc aggregation and floc break-up:

$$0 = -k_A' c^0 D_p^{-2} f_s^{-1} \rho_s^{-1} + k_B G^{1/2} (\gamma D_p^{-2} w_s^0 - 1),$$

$$\Rightarrow w_s^0 = \frac{k_A'}{k_B \sqrt{G}} f_s^{-1} \rho_s^{-1} \gamma^{-1} c^0 + \gamma^{-1} D_p^2,$$

$$\Rightarrow w_s^0 = \beta (c^0 + \kappa), \quad (2.53)$$

with $\kappa$ defined in Table 2.1. We added the second-order term $-k_B G^{1/2}$ from the break-up term to ensure that, in the limit for $c^0 \to 0$, the floc size is equal to the primary particle size: $D_f \to D_p$, which corresponds to a settling velocity of massive ($n_f = 3$) primary particles:

$$w_{s,\text{min.}} = \frac{(\rho_s - \rho_w) g D_p^2}{18 \mu} = \beta \kappa. \quad (2.54)$$

We can add this (small) term to our leading-order result and leave out the $\beta \kappa$ term in our higher-order calculations without restricting the generality of our results. The first-order
equation yields

\[
\frac{1}{\gamma(w_0^s)_{-1}^{1/\beta}} G \left\{ \partial_t c^0(w_0^s)^{-2} - \partial_z [c^0(w_0^s)^{-1}] - \partial_z [K^0 \partial_z c^0(w_0^s)^{-2}] \right\}
\]

\begin{equation}
= -k_A D_p^{-2} f_s^{-1} \rho_s^{-1} c^1 + k_B G^{1/2} (\gamma D_p^{-2} w_0^s),
\end{equation}

aggregation

break-up

\[ w_s^1 - \beta c^1 = \frac{1}{\gamma G k_B G^{1/2} (\gamma D_p^{-2})} \beta \times \]

\[
\left\{ \frac{\partial_t c^0(w_0^s)^{-2} - \partial_z [c^0(w_0^s)^{-1}]}{1/\beta} - \frac{\partial_z [K^0 \partial_z c^0(w_0^s)^{-2}]}{1/(\beta^2 c^0)} \right\},
\]

\[
= \frac{\partial_z (1/\beta) = 0}{\tau} w_s^1 - \beta c^1 = \frac{\tau}{\beta} \left\{- \frac{1}{(c^0)^2} \partial_t c^0 + K^0 \left[ \frac{-2}{(c^0)^3} (\partial_z c^0)^2 + \frac{1}{(c^0)^2} \partial_z c^0 \right] \right\},
\]

\[ \Rightarrow \]

\[
= \frac{1}{\beta} \left\{ \frac{1}{(c^0)^2} \left[ -\partial_t c^0 + K^0 \partial_z c^0 \right] - 2 K^0 \left( \partial_z c^0 \right)^2 \right\},
\]

\[ \Rightarrow w_s^1 = \beta c^1 - 2 \tau \frac{\partial_z c^0}{c^0} \left[ 1 + \frac{K^0}{w_0^s} \partial_z c^0 \right].\]

\[ (2.56) \]

**Appendix 2.D: calculation of tidal components of \( w_s \)**

In iFlow, we work with tidal components instead of time series. Therefore, we project the solutions of \( w_0^s \) and \( w_1^s \) on the subtidal, \( M_2 \), and \( M_4 \) tidal components. The amplitudes of the terms that are linear in \( c^0 \) and \( c^1 \) are trivial, for example:

\[ \hat{w}_s^{00} = \beta \hat{c}^{00} \quad \text{and} \quad \hat{w}_s^{04} = \beta \hat{c}^{04}, \]

\[ (2.57) \quad (2.58) \]

with \( \hat{c}^{00} \) and \( \hat{c}^{04} \) being (sub)tidal amplitudes. The projection of the second term in Eq. (2.56) is nontrivial. In general, an amplitudes \( A^n \) of a periodic function \( \text{func}(t) \) can be obtained by

\[ A^n = \frac{1}{T} \int_{-T/2}^{+T/2} \text{func}(t) \frac{e^{+i \frac{2 \pi n}{T} t} + e^{-i \frac{2 \pi n}{T} t}}{2} dt, \]

\[ (2.59) \]
with \( T \) being the period of function \( \text{func}(t) \) and in which \( n \) denotes the \((\text{sub-})\text{tidal constituent} \) (i.e., subtidal, M2, and M4). Substitution of \( \tilde{t} = M2t = (2\pi/T)t \) yields

\[
A^n = \frac{1}{2\pi} \int_{-\pi}^{+\pi} \text{func}(\tilde{t}) \frac{e^{+i\tilde{t}t} + e^{-i\tilde{t}t}}{2} d\tilde{t}.
\] (2.60)

Application to the second term of the right-hand side in Eq. (2.56), without the \(-2\tau\) factor, and substitution of \( \zeta = e^{it} \) yields

\[
A^n = \frac{1}{2\pi i} \oint \left\{ \frac{1}{\zeta} \left[ \frac{\partial_z \tilde{c}^{00} + \partial_z \tilde{c}^{00*} + (\partial_z \tilde{c}^{04})\zeta^{2} + (\partial_z \tilde{c}^{04*})\zeta^{-2}}{\tilde{c}^{00} + \tilde{c}^{00*} + \tilde{c}^{04} + \tilde{c}^{04*}} \right] \cdot \left[ 1 + 2 \frac{K_0^0}{\beta} \frac{\partial_z \tilde{c}^{00} + \partial_z \tilde{c}^{00*} + (\partial_z \tilde{c}^{04})\zeta^{2} + (\partial_z \tilde{c}^{04*})\zeta^{-2}}{\tilde{c}^{00} + \tilde{c}^{00*} + \tilde{c}^{04} + \tilde{c}^{04*}} \right] \right\} \frac{\zeta^{-n} + \zeta^{+n}}{2} d\zeta.
\] (2.61)

We use the residue theorem:

\[
\oint \text{func}(\zeta) d\zeta = 2\pi i \sum_k \frac{1}{(m-1)!} \lim_{\zeta \to \zeta_k} \frac{d^{m-1}}{d\zeta^{m-1}} \left[ (\zeta - \zeta_k)^m \text{func}(\zeta) \right],
\] (2.62)

in which \( \zeta_k \) is a pole of integrand \( \text{func} \) and \( m \) is the order of pole \( \zeta_k \). The integrand in Eq. (2.61) has five poles:

\[
\pm \zeta_{k_1} = \pm \sqrt{-\left(\tilde{c}^{00} + \tilde{c}^{00*}\right) + \sqrt{\left(\tilde{c}^{00} + \tilde{c}^{00*}\right)^2 - 4\tilde{c}^{04} \tilde{c}^{04*}}} \frac{2\tilde{c}^{04}}{2\tilde{c}^{04}},
\] (2.63)

\[
\pm \zeta_{k_2} = \pm \sqrt{-\left(\tilde{c}^{00} + \tilde{c}^{00*}\right) - \sqrt{\left(\tilde{c}^{00} + \tilde{c}^{00*}\right)^2 - 4\tilde{c}^{04} \tilde{c}^{04*}}} \frac{2\tilde{c}^{04}}{2\tilde{c}^{04}},
\] (2.64)

\[
\zeta = 0.
\] (2.65)

Assuming \( |\tilde{c}^{00}| > |\tilde{c}^{04}| \), only the three poles \( \zeta = \pm \zeta_{k_1} \) and \( \zeta = 0 \) lay inside the curvature of the integral and thus results in a nonzero contribution (this assumption is not necessary to prove that the M2 constituent is equal to 0). Consequently, we have to calculate the residue for three poles in order to compute the integral in Eq. (2.61). In the following, we show that the M2 tidal component is equal to 0 by computing the residue for pole \( \zeta = 0 \) and using the symmetry of the residue for the other two poles:

\[
\text{Res}_{\zeta_{k_1}} = -\text{Res}_{-\zeta_{k_1}} \quad \text{for n is odd \quad and \quad (2.66)}
\]

\[
\text{Res}_{\zeta_{k_1}} = \text{Res}_{-\zeta_{k_1}} \quad \text{for n is even. \quad (2.67)}
\]

56
2.D.1 Pole $\zeta = 0$

Calculating the residual for pole $\zeta = 0$ yields

for M0,\[ \frac{\partial_z e^{04*}}{e^{04*}}, \] (2.68)

for M2, 0, and

for M4,\[ \left\{ 2K_\nu(\partial_z e^{04*})^2 + \beta \left( (e^{04*})^2 \partial_z (e^{00} + c^{00*}) \right. \\
- \left. (c^{00} + c^{00*}) e^{04*} \partial_z e^{04*} \right) \right\} \frac{1}{2\beta(e^{04*})^3}, \] (2.70)

Consequently, the residue corresponding to the M2 constituent for pole $\zeta = 0$ is equal to 0.

2.D.2 Poles $\zeta = \pm \zeta_{k_1}$

We compute the residue for pole $\zeta = +\zeta_{k_1}$:

\[ \frac{1}{(m-1)!} \lim_{\zeta \to \zeta_{k_1}} \frac{d^{m-1}}{d\zeta^{m-1}} \left\{ \frac{1}{\zeta} \left[ \frac{Z_1}{Z_3} \cdot \left( 1 + 2\zeta^2 \frac{K_\nu Z_1}{Z_3^2} \right)^2 \right] \right\}, \] (2.71)

with

\[ Z_1 = \partial_z e^{04}(\zeta^2 - \zeta_{\partial_z k_1}^2)(\zeta^2 - \zeta_{\partial_z k_2}^2) \text{ symmetric and} \] (2.72)

\[ Z_3 = e^{04}(\zeta^2 - \zeta_{k_1}^2)(\zeta^2 - \zeta_{k_2}^2), \] (2.73)

and $\zeta_{\partial_z k_{1/2}}$ being the roots of the numerator:

\[ \zeta_{\partial_z k_{1/2}} = \left\{ \left[ - (\partial_z e^{00} + \partial_z e^{00*}) \right. \right.
\pm \sqrt{\left(\partial_z e^{00} + \partial_z e^{00*}\right)^2 - 4(\partial_z e^{04})(\partial_z e^{04*})} \left. \right] \frac{1}{2 \partial_z e^{04}} \right\}^{1/2}. \] (2.74)
The order \( m \) of the pole \( \zeta = \zeta_{k1} \) in the first term in Eq. (2.71) is equal to 1. In the second term, the order equals 3. Consequently, the residues yield

\[
\lim_{\xi \to \zeta_{k1}} \frac{\zeta^{-n-1} + \zeta^{+n-1}}{2} F_1
\]

\[
+ \frac{K_1}{2\beta} \lim_{\xi \to \zeta_{k1}} \left\{ (n+1)(n)\zeta^{n-1} F_2 + 2(n+1)\zeta^n \partial_\zeta F_2 + \zeta^{n+1} \partial_\zeta \zeta F_2 \right. \\
\left. + (-n+1)(-n)\zeta^{-n-1} F_2 + 2(-n+1)\zeta^{-n} \partial_\zeta F_2 + \zeta^{-n+1} \partial_\zeta \zeta F_2 \right\},
\]

(2.75)

with

\[
Z_1 = (\partial_\zeta c^{04})(\zeta^2 - \zeta_{\partial_z k1}^2)(\zeta^2 - \zeta_{\partial_z k2}^2) \quad \text{symmetric},
\]

(2.76)

\[
Z_2 = c^{04}(\zeta + \zeta_{k1})(\zeta^2 - \zeta_{\partial_z k1}^2) \quad \text{asymmetric},
\]

(2.77)

\[
\partial_\zeta Z_1 = 2\zeta(\zeta^2 - \zeta_{\partial_z k1}^2) \partial_\zeta c^{04} \\
\quad + 2\zeta(\zeta^2 - \zeta_{\partial_z k1}^2) \partial_\zeta c^{04} \quad \text{asymmetric},
\]

(2.78)

\[
\partial_\zeta Z_2 = (\zeta^2 - \zeta_{\partial_z k1}^2) c^{04} + 2\zeta(\zeta + \zeta_{k1}) c^{04} \quad \text{symmetric},
\]

(2.79)

\[
\partial_\zeta \zeta Z_1 = 2(\zeta^2 - \zeta_{\partial_z k1}^2) \partial_\zeta c^{04} + 2(\zeta^2 - \zeta_{\partial_z k2}^2) \partial_\zeta c^{04} \\
\quad + 8\zeta^2 \partial_\zeta c^{04} \quad \text{symmetric},
\]

(2.80)

\[
\partial_\zeta \zeta Z_2 = 4\zeta c^{04} + 2(\zeta + \zeta_{k1}) c^{04} \quad \text{asymmetric},
\]

(2.81)

\[
F_1 = \frac{Z_1}{Z_2} \quad \text{asymmetric},
\]

(2.82)

\[
F_2 = \frac{(Z_1)^2}{(Z_2)^3} \quad \text{asymmetric},
\]

(2.83)

\[
\partial_\zeta F_1 = 2 \frac{Z_1 \partial_\zeta Z_1}{(Z_2)^3} \quad \text{symmetric, and}
\]

(2.84)

\[
\partial_\zeta F_2 = 2 \frac{\partial_\zeta Z_1}{(Z_2)^3} + 2 \frac{Z_1 \partial_\zeta Z_1}{(Z_2)^3} - 6 \frac{Z_1 (\partial_\zeta Z_1)(\partial_\zeta Z_2)}{(Z_2)^4} \\
\quad - 3 \left\{ 2 \frac{Z_1 (\partial_\zeta Z_1)^2}{(Z_2)^4} + \frac{(Z_1)^2 \partial_\zeta \zeta Z_2}{(Z_2)^4} \\
\quad \quad - 4 \frac{(Z_1)^2 (\partial_\zeta Z_2)^2}{(Z_2)^5} \right\} \quad \text{asymmetric},
\]

(2.85)

in which we used that for pole \( \zeta = -\zeta_{k1} \) the factor \((\zeta + \zeta_{k1})\) is replaced by \((\zeta - \zeta_{k1})\) to determine the symmetry of \( Z_2 \). For \( n \) is odd (cf. M2, M6, M10, etc.), the solution in Eq. (2.75) is asymmetric and consequently the sum of residuals is equal to 0. For \( n \) is even (cf. M0, M4, etc.) the solution is symmetric. Consequently, adding the solution for the
pole $\zeta = 0$, the final solution yields

$$A^1 = 0$$
$$A^{0/2} = \lim_{\zeta \to \zeta_k} (\zeta^{-n-1} + \zeta^{+n-1}) F_1$$

$$+ \frac{K_0^0}{\beta} \lim_{\zeta \to \zeta_k} \left\{ (n+1)(n)\zeta^{n-1} F_2 + 2(n+1)\zeta^n \partial_\zeta F_2 \right.$$}

$$\left. + \partial_\zeta \zeta^{n+1} F_2 + (-n+1)(-n)\zeta^{-n-1} F_2 \right.$$}

$$\left. + 2(-n+1)\zeta^{-n} \partial_\zeta F_2 + \zeta^{-n+1} \partial_{\zeta\zeta} F_2 \right\}$$

$$(\zeta - \zeta_n - 1) \left( F_1 + K_0^0 \nu \beta \lim_{\zeta \to \zeta_k} (\zeta^{n+1} \partial_\zeta F_2 + 2(n+1)\zeta^n \partial_\zeta F_2) \right.$$}

$$\left. + 2(-n+1)\zeta^{-n} \partial_\zeta F_2 + \zeta^{-n+1} \partial_{\zeta\zeta} F_2 \right\}$$

which combined with the other trivial terms results in

$$\hat{\omega}^{10}_s = \beta c^{10} - 2\tau A^0,$$  \hspace{1cm} (2.88)

$$\hat{\omega}^{12}_s = \beta c^{12},$$  \hspace{1cm} (2.89)

$$\hat{\omega}^{14}_s = \beta c^{14} - 2\tau A^2.$$  \hspace{1cm} (2.90)

### Appendix 2.E: phase requirements for land-inward sediment transport due to flocculation

In this section, we demonstrate the phase conditions for which the M2 contribution of flocculation (solid black line in Fig. 2.6a) results in land-inward net sediment transport:

$$T^{12}_{\text{loc}} = \left\langle B \int_{-R}^{R} u^{02} c^{12}_{\text{loc}} dz \right\rangle > 0,$$  \hspace{1cm} (2.91)

in which $c^{12}_{\text{loc}}$ follows from Eq. (2.47):

$$\partial_t c^{1}_{\text{loc}} - \partial_z (u^{02}_{\text{loc}} c^{12}_{\text{loc}} + \partial_z K_B^{12} c^{1}_{\text{loc}}) = \beta \partial_\zeta (c^1 c^0).$$  \hspace{1cm} (2.92)

We assume that the water column is fairly well mixed, such that the vertical phase differences between $u^{02}$ and $c^{12}$ and $c^{04}$ are negligible. We define the phases in polar coordinates as

$$\dot{\hat{u}}^{02} = |\hat{u}^{02}|,$$  \hspace{1cm} (2.93)

$$\dot{\hat{c}}^{12} = |\hat{c}^{12}| e^{i\psi},$$  \hspace{1cm} (2.94)

$$\dot{\hat{c}}^{04} = |\hat{c}^{04}| e^{i\phi},$$  \hspace{1cm} (2.95)

$$\dot{\hat{c}}^{12}_{\text{loc}} = |\hat{c}^{12}_{\text{loc}}| e^{i\zeta}.$$  \hspace{1cm} (2.96)
Using these definitions and Eqs. (2.1) and (2.4), we rewrite Eq. (2.91) as

\[
\mathcal{T}_{floc}^{12} = B \int_{-H}^{R} \frac{1}{4} \left( \hat{u}^{02} \hat{c}_{floc}^{12} + \hat{u}^{02} \hat{c}_{floc}^{12} \right) dz
\]

\[
= \frac{B}{2} \int_{-H}^{R} |\hat{u}^{02}| |\hat{c}_{floc}^{12}| \Re \left[ e^{i\zeta} \right] dz.
\] (2.97)

So, \( \mathcal{T}_{floc}^{12} > 0 \) if \( \Re \left[ e^{i\zeta} \right] > 0 \), that is, if \(-\pi/2 < \zeta < \pi/2\). In the following, we link \( \zeta \) to \( \phi \) and \( \psi \) by integrating Eq. (2.92):

\[
\int_{-H}^{R} \partial_t \hat{c}_{floc}^{12} dz = \beta \Delta (c^1 c^0),
\] (2.98)

in which we used the fact that only the right-hand side of Eq. (2.92) generates an M2 tidal signal and

\[
\Delta (c^1 c^0) = (c^1 c^0) \bigg|_{z=R} - (c^1 c^0) \bigg|_{z=-H},
\] (2.99)

which results in

\[
i \omega M_2 \int_{-H}^{R} \hat{c}_{floc}^{12} dz = \beta \Delta \left( c_{floc}^{12} c^{00} + \frac{1}{2} c_{floc}^{12} c^{04} \right),
\]

\[
\implies \omega M_2 \int_{-H}^{R} |\hat{c}_{floc}^{12}| e^{i\zeta} dz = \beta \Delta \left( |c_{floc}^{12}| |c^{00} e^{i(\psi - \pi/2)}
\right.
\]

\[
+ \frac{1}{2} |c_{floc}^{12}| |c^{04} e^{i(\phi - \psi - \pi/2)} \right),
\] (2.100)

where we used Eqs. (2.2) and (2.4) and identified the M2 tidal phases in the right-hand side. We split the problem as

\[
\omega M_2 \int_{-H}^{R} |\hat{c}_{floc,1}^{12}| e^{i\zeta_1} dz = \beta \Delta \left( |\hat{c}_{floc,1}^{12}| |\hat{c}^{00} e^{i(\psi - \pi/2 + \pi)} \right),
\]

\[
\implies \zeta_1 = \psi + \pi/2
to and
\]

\[
\omega M_2 \int_{-H}^{R} |\hat{c}_{floc,2}^{12}| e^{i\zeta_2} dz = \beta \Delta \left( |\hat{c}_{floc,2}^{12}| |\hat{c}^{04} e^{i(\phi - \psi - \pi/2 + \pi)} \right),
\]

\[
\implies \zeta_2 = \phi - \psi + \pi/2,
\] (2.102)
in which we used the fact that a -1 factor results in a $\pi$ phase. Therefore, as a strong requirement for land-inwards sediment transport due to the M2 contribution of flocculation, we require that

$$-\pi < \psi < 0 \quad \text{and} \quad -\pi < \phi - \psi < 0.$$  \hspace{1cm} (2.103)

Likely, $|\hat{c}_{\text{floc,1}}^{12}| \gg |\hat{c}_{\text{floc,2}}^{12}|$, so the first requirement may be leading.
Bibliography


Chapter 3

The impact of temporal variability in phytoplankton on SPM dynamics

Abstract

Many estuaries exhibit seasonality in the estuary-scale distribution of suspended particulate matter (SPM). This SPM distribution depends on various factors, including freshwater discharge, salinity intrusion, erodibility, and the ability of cohesive SPM to flocculate into larger aggregates. Various authors indicate that biotic factors, such as the presence of algae and their excretion of sticky transparent exopolymer particles (TEP), affect the flocculation and erosion processes. Consequently, seasonality in these biotic factors may play a role in the observed seasonality in SPM. Whereas the impact of abiotic factors on seasonality in SPM is well studied, the relative contribution of biotically induced seasonality is largely unknown. In this study, we employ two approaches to assess the aggregated importance of biotically induced seasonality in flocculation and erosion on seasonality in SPM in the Scheldt estuary. In the first approach, we focus on seasonality of in situ observations in the Scheldt estuary of turbidity, floc size, Chlorophyll-a, and TEP, showing that the abiotic parameters show seasonality, while seasonality in TEP is ambiguous. The second approach concerns a reverse engineering method to calibrate biotically affected parameters of a coupled sediment transport-flocculation model to turbidity observations, allowing us to compare the modeled SPM concentrations to the observations. Driven by seasonality in freshwater discharge, the model captures both the observed seasonality in SPM without requiring biotically induced seasonality in flocculation and erosion, which is supported by the absence of seasonality in TEP.

3.1 Introduction

The suspended particulate matter (SPM) concentrations in estuaries often show regions in which SPM accumulates, resulting in zones with locally elevated SPM concentrations. These regions are called estuarine turbidity maxima (ETM), of which both the magnitude and location can show a significant seasonality. Such seasonality was observed in, for example, the Scheldt estuary (Cox et al., 2019), the Chesapeake Bay (Stanford et al., 2001), the Weser (Kappenberg and Grabemann, 2001) and the Hudson River estuary (Ralston et al., 2012). This seasonality in ETM is driven by seasonality in factors affecting transport of SPM, such as freshwater discharge (Allen et al., 1980; Uncles et al., 2006), salinity (MacCready and Geyer, 2010), erosion properties of the sediment bed (Stal, 2010), and settling velocity (Kranenburg, 1994; Winterwerp and van Kesteren, 2004; Verney et al., 2009; Fettweis et al., 2014). The latter two are related to both abiotic seasonal variations, such as variations in turbulence, salinity, and SPM (Dyer, 1989; van Leussen, 1994; Lai et al., 2018) and biotic variations (Frostick and McCave, 1979; Alldredge et al., 1993; Passow et al., 2001).

Biotic seasonality may impact the erosion properties of the sediment bed in multiple ways. On the one hand, various authors found that algal activity, which typically peaks in spring and summer, often stabilizes the sediment bed (Frostick and McCave, 1979) and induces the formation of bedforms (Malarkey et al., 2015) by, for example, the excretion of
extracellular polymeric substances (EPS), of which sticky transparent exopolymer particles (TEP) are a particulate form (Passow, 2002), and which bind the sediment together (Stal, 2010). On the other hand, the subsequent grazing of the algae by bioturbatory macroheterotrophs may generate an increase in erodibility (Paterson and Black, 1999).

Besides erosion properties of the sediment bed, biotic factors may also impact flocculation, which is defined as the aggregation and break-up of cohesive SPM, resulting in seasonal variations of the settling velocity of SPM flocs. Biotic characteristics, such as organic content, directly influence the differential density and structure of flocs (Kranenburg, 1994; van Leussen, 1994; Fall et al., 2021). Furthermore, organic content impacts, for example, the floc strength and collision efficiency (Winterwerp and van Kesteren, 2004), averaged floc size (Mietta et al., 2009), and floc break-up (Alldredge et al., 1990). Finally, in situ observations show a correlation between Chlorophyll-a (Chl-a) and flocculation efficiency (Verney et al., 2009), and TEP and floc strength (Fettweis et al., 2014).

These biotic seasonal variations in the settling velocity and erosion properties of the sediment bed are hypothesized to affect seasonality in ETM characteristics in estuaries. However, this hypothesis has not been systematically investigated. Therefore, in this contribution, we focus on quantifying the importance of such biotic seasonal variations to the seasonality in ETM, applied to the Scheldt estuary (Belgium and the Netherlands).

The Scheldt estuary is a particularly good candidate to study the influence of biotic seasonal variations on the seasonality in ETM for two reasons. Firstly, the settling velocity and erosion properties have been found to be important to the ETM formation (Brouwer et al., 2018; Horemans et al., 2020) and the corresponding floc size dynamics have been related to biotic activity (Wartel and Francken, 1998; Chen et al., 2005, 2018). Secondly, a recent study by Cox et al. (2019) reported a change in the seasonal ETM patterns and suggested that a change in biotically induced flocculation dynamics may be responsible for this change.

Assessing the relative importance of the abiotic and biotic factors that impact seasonality in SPM is challenging because of their strong interdependence. In addition, while reasonably accurate models exist to parameterize the effect of abiotic flocculation and erosion, explicit parameterizations of biotic effects relating, for example, Chl-a and TEP to flocculation and erodibility, rely on many assumptions (see Lai et al. (2018) for a recent review). This makes it difficult to dynamically solve the impact of biota on the large-scale ETM dynamics through flocculation and erosion, using explicitly defined bio-physical-chemical interactions.

In view of this, we employ a reverse engineering approach, combined with a detailed analysis of observations to assess the influence of biotically induced seasonality in flocculation and erosion on the large-scale SPM dynamics. Concerning the observations, we quantitatively study the seasonality in observed turbidity (cf. SPM), floc size, Chl-a, and TEP. This provides information on the seasonality of SPM dynamics, flocculation, and biota. Taking a reverse engineering model approach, following Chen et al. (2018), the model does not rely on explicit model parameterizations for biotic flocculation and erosion. Instead, the accumulated biotic impact is accounted for parametrically, using two.
empirical parameters; one in the flocculation model of Horemans et al. (2020) and one in the erosion model (Partheniades, 1965; Kandiah, 1974; Dijkstra et al., 2019). Abiotic seasonal variations, including variations of the river discharge, salinity, and abiotic effects on flocculation and erosion are explicitly taken into account. The two biotically affected parameters are calibrated to long-term SPM observations of both summer and winter conditions in the Scheldt estuary. We hypothesize that, if a seasonality in biological factors were to have a significant impact on the large-scale SPM distribution, we would obtain a significantly different value of these biotically affected parameters to represent summer and winter conditions in the Scheldt estuary.

3.2 Material and methods

In this section, we first discuss some of the characteristics of the Scheldt estuary. Next, the measurement methods used to obtain the in situ observations are introduced and the model approach is presented.

3.2.1 Study area

The Scheldt estuary is an approximately 160 km long, funnel-shaped estuary (Fig. 3.1). It flows through Belgium into the North Sea near Vlissingen (Netherlands). Because of its relatively small freshwater discharge, the Scheldt estuary can be considered a tide-dominated and well-mixed estuary (Meire et al., 2005). The total time-averaged freshwater discharge $Q$ over the years 2015-2018 equals $40 \text{ m}^3 \text{s}^{-1}$ and $174 \text{ m}^3 \text{s}^{-1}$ in summer (Jun.-Aug.) and winter (Jan.-Mar.), respectively (Waterinfo.be, 2019). The main tributaries of the Scheldt estuary are the Rupel and the Dender. The Upper Sea Scheldt boundary (i.e., the upstream boundary of the Scheldt estuary), the Rupel, and the Dender are responsible for 16.4, 77.2, and 6.4 % of the total river discharge in summer and 35.4, 53.1, and 11.5 % in winter, respectively.

The maintenance of the navigation channel to the port of Antwerp (first four red dots from the Dutch-Belgian border in Fig. 3.1) requires intensive dredging activities. To minimize the risk of flooding, the dredged material is dumped back into the Scheldt estuary, which comes with significantly higher SPM loads (~$10^6$ ton year$^{-1}$) than the fluvial input (Dijkstra et al., 2019). At the main dumping locations at km 73 and 78 from the mouth, the time-averaged dumped material between 2001-2015 corresponds to a high SPM input of approximately 60 and 100 kg s$^{-1}$, respectively. Consequently, these dumping activities impact the SPM concentration locally, which was confirmed by a multivariate regression analysis of SPM observations between 1996-2016. This analysis showed that the SPM concentrations at the dumping sites are dominantly correlated to dumping activities (IMDC, 2016). The yearly fluvial input of SPM at the upstream boundary and tributaries is of the order of $10^4$-$10^5$ ton year$^{-1}$ and is assumed to scale linearly with the freshwater discharge (Plancke et al., 2017; Dijkstra et al., 2019).
Figure 3.1: The Scheldt estuary and its two main tributaries (Rupel and Dender). The dots represent the locations where monthly and biweekly turbidity and floc size profiles were measured in the frame of the OMES (Dutch: “Onderzoek Milieu Effecten Sigma-plan”) environmental monitoring program. The transparent exopolymer particles (TEP) concentration was only measured at 11 instead of the 16 stations, depicted in red.
3.2.2 In situ observations

The Belgian part of the Scheldt estuary has been monitored by the long-term OMES (Dutch: “Onderzoek Milieu Effecten Sigmaplan”) monitoring campaign (Maris and Meire, 2017). Within this campaign, various variables, including turbidity, SPM, salinity, primary particle size, floc size, Chl-a, and TEP concentration, have been measured biweekly or monthly and independently of the tidal phase and spring-neap tide at 16 fixed stations (see Fig. 3.1). In the following, we discuss the measurement methods to obtain the observations of the four variables we focus on: turbidity (Maris et al., 2021b), floc size (Maris et al., 2021a), Chl-a (Maris and Meire, 2021), and TEP (Vyverman et al., 2021).

Turbidity depth profiles were measured over the years 2015-2018 using an optical backscatter point sensor (OBS) of RBR type XR420 CTD+. To translate turbidity to SPM concentration, we simultaneously collected two SPM samples at approximately the water surface and half the water depth and applied a linear data fit (Horemans et al., 2020). The observations cover 18 and 14 campaigns in summer (Jun.-Aug.) and winter (Jan.-Mar.), respectively, resulting in 18 and 14 SPM profiles at each of the 16 locations.

From Sep. 2017 until Jan. 2019, floc size profiles were measured mainly in the upper half of the water column, simultaneously with the turbidity profiles. Floc size was measured using an optical laser diffraction instrument, a Sequoia laser in situ scattering and transmissometry (LISST) 200x instrument. The LISST 200x instrument measures the floc size volume distribution for 36, nonequidistant size classes with a floc size ranging between 1-500 µm. This particle size distribution was used to determine the geometric averaged floc size following the averaging method of Sequoia Scientific (2019) to correct for the nonequidistant size classes. Because the estimated floc size is highly sensitive to the averaging method, we also use two additional averaging methods in Appendix 3.A.

Simultaneously with measuring the turbidity and floc size profiles, bucket samples at the water surface were taken to estimate the Chl-a and TEP concentration. The TEP measurements only started in 2018. We estimated the Chl-a concentration, following the spectrophotometric method described in Rice et al. (2017) that corrects for turbidity, Chlorophyll-b, Chlorophyll-c, and Pheophytin pigments. We used 50 ml water samples, a 1 cm pathway cuvette, and a Shimadzu UV-1700 spectrophotometer. The detection threshold of the method is approximately 10 µg L⁻¹. When Chl-a values are below this threshold value, they are put to 10 µg L⁻¹. The total number of observed Chl-a concentrations per station equals 24 in summer and ranges from 11 up to 13 in winter, depending on the station. TEP concentrations were estimated according to the colorimetric method described in Claquin et al. (2008) adapted from the original method described in Pascow and Alldredge (1995). TEP concentrations were only measured once a month at 11 stations (see Fig. 3.1, red dots), resulting in 66 estimates in both summer and winter.
3.2.3 Model and model experiments

**iFlow model** The model used is the process-based, width-averaged, idealized model called ‘iFlow’ that solves for water motion and SPM trapping in tide-dominated estuaries using the width-averaged shallow water and SPM mass balance equations in equilibrium condition ([Dijkstra et al., 2017](#)), including the effects of flocculation ([Winterwerp, 2002](#) [Horemans et al., 2020](#)). The width and bathymetry of the estuary are approximated by smooth profiles, focusing on the estuary-scale hydro- and SPM dynamics only. The model resolves the tidal and subtidal water motion and SPM concentration using a scaling and perturbation approach. This approach drastically reduces computation time, allowing us to carry out an extensive sensitivity analysis. Moreover, it simplifies the interpretation of the results as it allows us to study physical processes separately.

The water motion is forced at the mouth by a tidal signal and at the Upper Sea Scheldt boundary and two main tributaries by a constant water inflow that depends on the season. SPM dynamics are forced by a constant concentration of SPM at the mouth and an inflow of SPM at the Upper Sea Scheldt boundary, which is obtained from observations and equals the product of the subtidal SPM concentration and freshwater discharge. Although clearly detectable in the SPM observations at the dumping locations, we do not include dumping of dredged material because it acts locally on a much smaller time scale (~hours) and does not show seasonality. Therefore, it is out of the scope of this study, which focuses on large-scale SPM dynamics on the seasonal scale. Moreover, we are interested in biotically affected, and not anthropogenically induced, erosion and flocculation. The longitudinal salinity profile is prescribed as a depth- and tide-independent sigmoid profile ([Warner et al., 2005](#)) that depends on the season (see Appendix 3.B). This assumption is reasonable as the Scheldt is a well-mixed estuary in all seasons. Erosion of SPM is assumed to scale with the magnitude of the bed shear stress. The biotic impact on erosion is implicitly included in the corresponding scaling factor called the erosion parameter $M$. This parameter is an intrinsic property of the sediment bed, including biotic effects, and determines the erosion flux $E$ for a fixed bed shear stress $\tau_b$ and erodibility $f$:

$$E = M |\tau_b| f(a)$$

(3.1)

where $a$ is the availability of easily erodible fine sediment and $M$ is the erosion parameter.

The flocculation model describes the spatial and temporal evolution of the settling velocity of a single class of flocs. The effects of abiotic conditions (e.g., salinity, SPM concentration, and turbulence) are contained explicitly, whereas biotic effects (such as that of TEP) are implicitly included in the flocculation model. The parameter controlling the importance of biotic effects is denoted by $\lambda$, and is defined as the ratio of the minimal floc aggregation $k_A^{\text{min}}$ and floc break-up parameter $k_B$. These two parameters characterize the flocculation properties given a fixed SPM concentration, salinity $S$, and shear rate $G$. For the model implementation of the shear rate $G$ in summer and winter, we refer the reader to Appendix 3.C. A careful scaling and perturbation analysis shows that $\lambda$ impacts
the settling velocity \( w_s \) (upto fist order) (Horemans et al., 2020) as

\[
w_s = \frac{1}{18 \mu f_s} \frac{\rho_s - \rho_w}{\rho_s} g \frac{w_{s,\min}^A}{k_B} f^{\text{sal}}(S) \cdot c + w_{s,\min},
\]

(3.2)
in which we assumed a fractal dimension \( n_f = 2 \) (see Appendix 3.D for \( n_f \neq 2 \)), spherical flocs, and that the floc Reynolds Number \( Re_f = w_s D_f / \nu \ll 1 \), where \( \nu \) is the kinematic viscosity of water, and in which \( w_{s,\min} \) is the settling velocity of the primary particles, \( \mu \) is the dynamic viscosity of water, \( f_s \) is the shape factor of the flocs, \( \rho_s \) is the density of the primary particles, \( \rho_w \) is the density of water, \( g \) is the gravitational acceleration, \( c \) is the SPM concentration, and \( f^{\text{sal}}(S) \) describes salinity driven flocculation by increasing the colloid stability for increasing salinity (Edzwald et al., 1974; Horemans et al., 2020). The impact of salinity driven flocculation is relatively low, which agrees with Einstein and Krone (1962) who showed that system-wide variations in salinity have only a minor influence on the bond strength. Salinity may also impact flocculation indirectly through, for example, its impact on TEP production (Alldredge et al., 1993; Bar-Zeev et al., 2015). This biotic impact is parametrically included in \( \lambda \).

**Reverse engineering model approach** To detect a biotically induced seasonality that impacts the seasonality in estuary-scale SPM dynamics, we follow Chen et al. (2018) by applying a reverse engineering model approach. This means that the biotically induced parameters \( \lambda \) and \( M \) are calibrated to observed SPM concentrations in summer and winter and that we statistically evaluate if the calibrated \( \lambda \) and \( M \) are the same in both seasons. We vary \( \lambda \) and \( M \) within a realistic range of values. The range in \( \lambda \) is \( 36-121 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2 \) and in \( M \) is \( 0.1-6 \times 10^{-3} \text{ s}^{-1} \text{ m}^{-1} \), which is based on scaling (Horemans et al., 2020) and observations (Zhu et al., 2017), respectively.

As a first step in the reverse engineering approach, we average the observed SPM concentrations of the various measuring campaigns at each depth. We assume that this averaged value approximates the residual SPM concentration similar to Dijkstra et al. (2019), Cox et al. (2019) and Horemans et al. (2020). This assumption is reasonable when the number of estimates is sufficiently large (\( \gtrsim 10 \)), so by averaging the periodic temporal variability of the SPM concentration vanishes (Horemans et al., 2020). Here, we assume that the campaigns are homogeneously distributed within the spring-neap cycle and tidal phase, which was shown to be valid for the SPM sampling within the OMES framework between 1995 and 2015 (after 2015, the setup of the campaigns has not changed) (Vandenbruwaene et al., 2016). Moreover, we assume that each year between 2015-2018 corresponds to a similar SPM distribution on the estuary scale. To comply with the latter requirement, we leave out the year 2016 because of its exceptionally high amount of rainfall. Next, to compare the modeled residual SPM concentration to the time-averaged observed SPM concentration, we interpolate the time-averaged SPM observations to a grid with 16 cells in the along-channel direction (corresponding to the observation stations) and 50 cells in the vertical direction (independent of the water depth). The average SPM concentration in each cell is thus based on the average of 18 and 14 observations in summer and winter, respectively. In the remainder of this study, we do not consider the SPM concentration in the vertical cells closest to the sediment bed.
and water surface, as there are insufficient measurements there (due to distortions caused by air bubbles and high turbidity at the water surface and sediment bed, respectively). As a second step, we quantify the accuracy of the modeled SPM concentration compared to the observations using a cost function that is based on a statistical two-tailed $t$ test. The cost function varies between 0 (perfect match between modeled SPM concentration and observations) and 1 (complete mismatch between modeled SPM concentration and observations). We refer the reader to Horemans et al. (2020) for a detailed description of the cost function. We exclude the SPM observations at the dumping locations (total of two stations) because here we expect high variability in SPM due to anthropogenic dumping of sediment, which is out of the scope of this study and not included in the model (see Section 3.2.3).

Finally, uncertainty in the SPM observations results in uncertainty in the optimal (cf. lowest cost function value) $(\lambda, M)$ pairs. To quantify this uncertainty, we assume that the average SPM concentration in summer and winter is normally distributed with mean value equal to the mean of the OMES observations and standard deviation equal to the standard deviation in the OMES observations divided by the square root of the number of observations (i.e., 18 and 14 in summer and winter, respectively). To equate this to a probability distribution for the optimal $(\lambda, M)$ pair, we randomly sample 500 averaged SPM concentrations from this normal distribution. Because the SPM concentrations in the vertical direction are not independent, we require that the relative deviation from the mean is equal at every depth at a given station, thus keeping the shape of the vertical time-averaged SPM profile fixed. Next, we calibrate $\lambda$ and $M$ for each sample. The resulting calibration is used to compute the standard deviation $\sigma_{\lambda,M}$ of the 500 $(\lambda, M)$ pairs for summer and winter. Note that we do not assume that $\lambda$ and $M$ are necessarily normally distributed. Finally, to quantitatively compare the optimal $(\lambda, M)$ pairs in summer and winter conditions, we use this standard deviation $\sigma_{\lambda,M}$ to compute the difference of the optimal $\lambda$ and $M$ values in summer and winter conditions, relative to the corresponding standard deviation:

$$\Delta_{\lambda} = \frac{|\lambda_{\text{summer}} - \lambda_{\text{winter}}|}{\sqrt{(\sigma_{\lambda,\text{summer}})^2 + (\sigma_{\lambda,\text{winter}})^2}}$$

$$\Delta_{M} = \frac{|M_{\text{summer}} - M_{\text{winter}}|}{\sqrt{(\sigma_{M,\text{summer}})^2 + (\sigma_{M,\text{winter}})^2}}$$

(3.3)

If $\Delta_{\lambda,M} < 1$, the average difference between summer and winter is less than one standard deviation and we will conclude that the difference of $\lambda$ or $M$ in winter and summer conditions is within the uncertainty of the SPM observations. The model parameter values for both the summer and winter conditions in the Scheldt estuary are given in Table 3.1. These values are either based on observations or calibration. Previous work indicated that the average summer discharge is representative, while the representative winter discharge is slightly above the average of 174 m$^3$ s$^{-1}$ to correct for the large variability in discharges during winter (Horemans et al., 2020). The parameters corresponding to the depth- and tide-independent salinity profile (i.e., $x_c$ and $x_L$) were fitted to salinity observations (see Appendix 3.B). The system-averaged primary particle size $D_p$ was estimated using observations (see Appendix 3.E). If not mentioned explicitly, other used parameter values are presented in Horemans et al. (2020). For completeness, we repeated these parameter values in Table 3.F.1.

To test the robustness of the model, we apply a sensitivity analysis of two crucial
parameters which are assumed to be fixed in the model: the primary particle size $D_p$ and fractal dimension $n_f$. To apply a sensitivity analysis to the latter parameter, we have generalized the results presented in Horemans et al. (2020) for $n_f \neq 2$ (see Appendix 3.D). Based on the experiments presented in Appendix 3.E and the literature (Winterwerp and van Kesteren, 2004), we vary $D_p$ and $n_f$ between 4-25 µm and 1.7-3, respectively.

### 3.3 Results

This section first introduces the seasonality of the in situ observations of SPM, floc size, Chl-a, and TEP concentrations in the Scheldt estuary. Next, we show the results of our model experiments, focusing on the seasonality in biotically affected calibration parameters $\lambda$ and $M$ and modeled SPM concentration. Finally, we present the sensitivity analysis of the modeled SPM concentration to $D_p$ and $n_f$.

#### 3.3.1 Seasonality in the in situ observations

##### 3.3.1.1 Seasonality in SPM

Figure 3.2 shows the observed time-averaged SPM concentration in summer and winter. Both in summer and winter, the concentrations range up to approximately 300 mg L$^{-1}$ and typically increase with depth. In summer (Fig. 3.2a), two ETM are observed, located near km 80 (depicted by I) and 120 (depicted by II). The ETM have a length of approximately 10 and 30 km, respectively. The ETM near km 80 is potentially due to anthropogenic dumping activities in this region (depicted by the vertical dashed lines), as mentioned above. In winter (Fig. 3.2b), the ETM at km 80 is also present, while the ETM near km 120 moves downstream, and shows significantly lower SPM concentrations (depicted by II). Moreover, we detect an additional ETM near km 60 (depicted by III). We thus conclude that the SPM distribution in the Scheldt estuary shows strong seasonality, which complies with the results of Cox et al. (2019) and has been observed in many other estuaries (see Burchard et al. (2018) for a recent review).

##### 3.3.1.2 Seasonality in floc size

Figure 3.3 shows the observed depth- and time-averaged floc size measured in summer and winter 2018. The error bars depict the standard error of the depth-averages corresponding to the various campaigns. In summer, we have an approximately constant depth-averaged floc size in the longitudinal direction. The floc sizes are in the order of 100 µm. In winter, the floc sizes decrease in the Upper Sea Scheldt beyond km 120 and are statistically smaller compared to the floc sizes in summer (one-tailed $t$ test, p-value $< 10^{-4}$). The averaged floc sizes in the Upper Sea Scheldt beyond km 120 are 115 and 77 µm in summer.
Table 3.1: Parameter values used in our model study in the Scheldt estuary in summer and winter conditions.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Q_{\text{summer}}$</td>
<td>Total river discharge in summer (calibrated)</td>
<td>$40 \text{ m}^3 \text{s}^{-1}$</td>
</tr>
<tr>
<td>$Q_{\text{winter}}$</td>
<td>Total river discharge in winter (calibrated)</td>
<td>$233 \text{ m}^3 \text{s}^{-1}$</td>
</tr>
<tr>
<td>$c_{\text{upper Sea Scheldt}}$</td>
<td>Subtidal SPM concentration at the Upper Sea Scheldt boundary (source)</td>
<td>$0.035 \text{ kg m}^{-3}$</td>
</tr>
<tr>
<td>$M_{\text{summer}}$</td>
<td>Erosion parameter in summer (calibrated)</td>
<td>$2.7 \times 10^{-3} \text{ s m}^{-1}$</td>
</tr>
<tr>
<td>$M_{\text{winter}}$</td>
<td>Erosion parameter in winter (calibrated)</td>
<td>$3.0 \times 10^{-3} \text{ s m}^{-1}$</td>
</tr>
<tr>
<td>$\sigma_p$</td>
<td>Prandtl Schmidt number</td>
<td>$0.5$</td>
</tr>
<tr>
<td>$s_{f_0}$</td>
<td>Bed roughness coefficient (calibrated)</td>
<td>$4.22 \text{ mm s}^{-1}$</td>
</tr>
<tr>
<td>$k_{\text{min.}}$</td>
<td>Nondimensional minimal aggregation coefficient</td>
<td>$0.29$</td>
</tr>
<tr>
<td>$f_s$</td>
<td>Shape factor</td>
<td>$\pi/6$</td>
</tr>
<tr>
<td>$n_f$</td>
<td>Fractal dimension of flocs</td>
<td>$2$</td>
</tr>
<tr>
<td>$D_p$</td>
<td>Diameter primary mud particles (source)</td>
<td>$14 \times 10^{-6} \text{ m}$</td>
</tr>
<tr>
<td>$\rho_s$</td>
<td>Density of SPM primary particles</td>
<td>$2650 \text{ kg m}^{-3}$</td>
</tr>
<tr>
<td>$\rho_w$</td>
<td>Reference density of water</td>
<td>$1000 \text{ kg m}^{-3}$</td>
</tr>
<tr>
<td>$\lambda_{\text{summer}}$</td>
<td>Ratio of minimal aggregation and floc break-up parameter in summer (calibrated)</td>
<td>$72.5 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$</td>
</tr>
<tr>
<td>$\lambda_{\text{winter}}$</td>
<td>Ratio of minimal aggregation and floc break-up parameter in winter (calibrated)</td>
<td>$65.9 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$</td>
</tr>
<tr>
<td>$s_{\text{sea}}$</td>
<td>Salinity boundary condition at the mouth</td>
<td>$28.9 \text{ psu}$</td>
</tr>
<tr>
<td>$x_{c_{\text{sal, summer}}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in summer</td>
<td>$51.9 \text{ km}$</td>
</tr>
<tr>
<td>$x_{L_{\text{sal, summer}}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in summer</td>
<td>$31.2 \text{ km}$</td>
</tr>
<tr>
<td>$x_{c_{\text{sal, winter}}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in winter</td>
<td>$38.3 \text{ km}$</td>
</tr>
<tr>
<td>$x_{L_{\text{sal, winter}}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in winter</td>
<td>$24.8 \text{ km}$</td>
</tr>
</tbody>
</table>
(a) Observations of the SPM concentration (mg L\(^{-1}\)) in summer

(b) Observations of the SPM concentration (mg L\(^{-1}\)) in winter

Figure 3.2: Measured long-term time-averaged suspended particulate matter (SPM) concentration (mg L\(^{-1}\)) over the years 2015-2018 in (a) summer, showing two ETM at locations I and II, and (b) winter showing three ETM at locations I, II, and III. The vertical dashed lines depict the dumping locations.

and winter, respectively. We thus conclude that, on the estuary scale (we observe a local anomaly near km 60 and km 80), the floc sizes only show seasonality in the upstream part of the Scheldt estuary (> 120 km). This is also valid using other averaging methods (see Appendix 3.A).

3.3.1.3 Seasonality in Chl-a and TEP

The measured time averages of the observations of Chl-a and TEP concentrations in summer and winter are shown in Fig. 3.4. The error bars represent the standard error of the mean corresponding to the concentrations of the various campaigns. The Chl-a observations (Fig. 3.4a) show a strong seasonality in the Upper Sea Scheldt (> 100 km), where the Chl-a concentration in summer is an order of magnitude larger than in winter. In contrast, although the TEP observations (Fig. 3.4b) show local seasonality at km 58, 121, and 138, they do not show a significant system-wide seasonality in the Upper Sea Scheldt (one-tailed t test, p-value > 0.38). The error bars corresponding to the TEP measurements are relatively large, especially in the Lower Sea Scheldt (< 100 km). This is due to the relatively small number of estimates and the large temporal variability in TEP, a variability also observed in other estuarine and marine systems, such as the Neuse River Estuary (Wetz et al., 2009), the Pearl River estuary (Sun et al., 2012), and the Chesapeake bay (Malpezzi et al., 2013). What is most important within the scope of this study is that the Chl-a observations show seasonality, whereas the TEP observations do not.
Figure 3.3: Measured time- and depth-averaged floc size in summer (red) and winter (blue). The error bars depict the standard error of the estimates corresponding to the various campaigns.

Figure 3.4: (a) Measured time-averaged Chlorophyll-a (Chl-a) concentration in summer and winter over the years 2015-2018 and (b) corresponding measured transparent exopolymer particles (TEP) concentration of 2018. The error bars depict the standard error of the estimates corresponding to the various campaigns.
We conclude that we observe a clear seasonality in SPM and floc size dynamics beyond km 120, whereas some biotic properties show seasonality, and others do not; the contrast between the strong seasonality in Chl-a and the absence of seasonality in TEP on the estuary scale, both biotic factors reported to (indirectly) influence SPM dynamics, makes it difficult to use these observations as indicators for biotically induced seasonality in SPM. Therefore, the observations are complemented with model experiments which are shown in the following section.

3.3.2 Seasonality in the model experiments

3.3.2.1 Model parameters $\lambda$ and $M$

Minimizing the cost function separately for summer and winter results in an optimal parameter set $\lambda = 72.5 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 2.7 \times 10^{-3} \text{ s m}^{-1}$ in summer (cost value equals 0.22) and $\lambda = 65.9 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 3.0 \times 10^{-3} \text{ s m}^{-1}$ in winter (cost value equals 0.17). These optimal $(\lambda,M)$ pairs thus show a weak ($\sim 10\%$) seasonality in both $\lambda$ and $M$.

To explore the sensitivity of the model results to variations of the calibration parameters $\lambda$ and $M$, Fig. 3.5 shows the cost function as a function of $\lambda$ (horizontal axis) and $M$ (vertical axis), with the value of the cost function indicated by color. Optimal values are indicated by $\bigcirc$. The dark blue colors indicate the range of $(\lambda,M)$ pairs with a good performance: the white dashed contours depict the cost function values which are 25% larger than the optimal cost function value. In summer (Fig. 3.5a), this contour covers a range with $\lambda$ close to $70 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$ and $2 \times 10^{-3} < M < 5 \times 10^{-3} \text{ s m}^{-1}$. So, the optimal $\lambda$ value is found in a relatively narrow parameter range and is quite insensitive to the value of $M$. In winter (Fig. 3.5b), the range of optimal values is around $\lambda \approx 60 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$ and $M > 2 \times 10^{-3} \text{ s m}^{-1}$. Note our winter result slightly differs from the winter result presented in Horemans et al. (2020) because we added fluvial inflow of SPM at the Upper Sea Scheldt boundary (see Table 3.1), used a Prandtl Schmidt number of 0.5 instead of 1, and excluded the observations at the dumping locations in our calibration. We also included turbidity observations of 2018 in our calibration.

The red dots in Fig. 3.5 depict the probability density of $(\lambda,M)$ given the uncertainty in the residual SPM data set to which the model is calibrated. Bigger dots signify a higher probability density, while smaller dots signify a lower probability density. We use these distributions of $(\lambda,M)$ in summer and winter to quantitatively compare the optimal $(\lambda,M)$ pairs using Eq. (3.3) and detect a potential seasonality given the uncertainty in the residual SPM observations. The standard deviation of the $\lambda$ and $M$ distributions in the summer and winter conditions are $\sigma_\lambda^\text{summer} = 12.9 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $\sigma_\lambda^\text{winter} = 10.5 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $\sigma_M^\text{summer} = 0.9 \times 10^{-3} \text{ s m}^{-1}$, and $\sigma_M^\text{winter} = 1.4 \times 10^{-3} \text{ s m}^{-1}$. Consequently, the relative difference of the optimum values in summer and winter $\Delta$ [see Eq. (3.3)] for $\lambda$ and $M$ are $\Delta_\lambda = 0.4$ and $\Delta_M = 0.18$, respectively. We thus conclude that we do not detect seasonality in $\lambda$ and $M$ given the uncertainty of the residual SPM observations.
Figure 3.5: Cost function values for various $\lambda$ and $M$ calibrated to the observed residual SPM data set in (a) summer and (b) winter. The optimal calibration parameter set corresponds to $\lambda = 72.5 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 2.7 \times 10^{-3} \text{ s m}^{-1}$ in summer and $\lambda = 65.9 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 3 \times 10^{-3} \text{ s m}^{-1}$ in winter. The white dashed contours depict the cost function values which are 25% larger than the optimal cost function value. The red dots depict the probability density of the 500 $(\lambda,M)$ pairs given the uncertainty in the residual SPM data set to which the model is calibrated. Bigger dots signify a higher probability density, while smaller dots signify a lower probability density. We choose two additional $(\lambda,M)$ pairs, depicted by (2) and (3), to study the impact of the choice in $(\lambda,M)$ located in the zone of high probability density on the corresponding model output. The corresponding $\lambda$ and $M$ values are $\lambda = 72.5 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 2.7 \times 10^{-3} \text{ s m}^{-1}$ (i.e., optimal summer conditions) and $\lambda = 45.3 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 1.1 \times 10^{-3} \text{ s m}^{-1}$, respectively.
3.3.2.2 Seasonality in modeled SPM concentration

To further investigate the optimal calibration values of $\lambda$ and $M$, we analyze the corresponding modeled SPM concentration distribution for the optimal $(\lambda, M)$ pair (see (1) in Fig. 3.5) for both summer and winter conditions. Given the estuary-scale of the model and measurement uncertainty (cf. dark blue regions and distribution of $(\lambda, M)$ pairs in Fig. 3.5), we also show the SPM distribution corresponding to two additional $(\lambda, M)$ pairs (see (2) and (3) in Fig. 3.5b). We require that these two pairs result in an acceptable model performance and are within the zone of high probability density in winter (red dots in Fig. 3.5b). We choose the first pair to be equal to the optimal $(\lambda, M)$ pair in summer and the second pair to be located in the second zone of $(\lambda, M)$ pairs in winter where we detect a high probability density (depicted by the red dots in Fig. 3.5b). The corresponding $\lambda$ and $M$ values are $\lambda = 72.5 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 2.7 \times 10^{-3} \text{ s m}^{-1}$ and $\lambda = 45.3 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$, $M = 1.1 \times 10^{-3} \text{ s m}^{-1}$, respectively. The corresponding cost function values equal 0.26 and 0.30, respectively.

![Model output of the SPM concentration](image1)

![Observations of the SPM concentration](image2)

Figure 3.6: (a) Model output of the suspended particulate matter (SPM) concentration in summer. The optimal calibration parameters are (1) $\lambda = 72.5 \times 10^{-6} \text{ s}^{-1/2} \text{ m}^2$ and $M = 3 \times 10^{-3} \text{ s m}^{-1}$. The vertical dashed lines depict the dumping locations and the symbols (I), (II) the locations of the observed ETM. (b) The long-term, time-averaged SPM concentration observations (mg L$^{-1}$) in summer.

Figures 3.6-3.7 show the modeled SPM concentration in summer and winter using these values of $\lambda$ and $M$ and the corresponding observations. We present the model result and observations in summer in Fig. 3.6a and Fig. 3.6b, respectively. Figure 3.6a shows two modeled ETM at the correct locations, that is, a weak ETM at the dumping location near km 80 and an ETM at km 120 (depicted by (I) and (II), respectively). The length of the ETM is approximately 10 and 30 km, respectively. Both model results and observations show concentrations up to approximately 300 mg L$^{-1}$. The model seems to overestimate the vertical gradient in SPM.
Figure 3.7: (a) Model output of the suspended particulate matter (SPM) concentration in winter. The optimal calibration parameters are $\lambda = 65.9 \times 10^{-6}$ s$^{-1/2}$ m$^2$ and $M = 3.0 \times 10^{-3}$ s m$^{-1}$. The vertical dashed lines depict the dumping locations and the symbols I, II, and III the locations of the observed ETM. (b) The long-term, time-averaged SPM concentration observations (mg L$^{-1}$) in winter. (c)-(d) Model output of the SPM concentration in winter with $\lambda = 72.5 \times 10^{-6}$ s$^{-1/2}$ m$^2$, $M = 2.7 \times 10^{-3}$ s m$^{-1}$ and $\lambda = 45.3 \times 10^{-6}$ s$^{-1/2}$ m$^2$, $M = 1.1 \times 10^{-3}$ s m$^{-1}$, respectively.

In winter, the model also captures the locations of the ETM for the optimal $(\lambda,M)$ pair (1). The model result (Fig. 3.7a) and observations (Fig. 3.7b) both show an ETM near km 80, approximately 10 km long (depicted by I). The ETM in the model result potentially merged into the ETM caused by dumping activities, which are not considered in the model setup. The model also correctly captures the accumulation of SPM at km 100 (depicted by II) and km 60 (depicted by III). The concentrations are in the order of 100-300 mg L$^{-1}$. What is most important within the scope of this study is that the
ETM near km 120 in summer shifts downstream to km 100 and an additional ETM near km 60 arises in winter. In other words, the model captures this seasonality in the ETM.

Figures 3.7c and 3.7d show the modeled SPM concentration corresponding to the two alternative \((\lambda, M)\) pairs (see \(\circ\) and \(\triangle\) in Fig. 3.5d). Compared to our previous result (Fig. 3.7b), no major change in case of parameter values associated with \(\circ\) is observed (Fig. 3.7c). We observe a slight decrease in the SPM concentration, especially near the sediment bed, but the locations of the ETM are unchanged. The modeled SPM concentration corresponding to pair \(\triangle\) (Fig. 3.7d) is different when compared to the optimal SPM concentration (Fig. 3.7a): the ETM near km 60 (depicted by \(\diamond\)) vanished, the ETM near km 80 and 100 are still present (depicted by \(\mathbb{1}\) and \(\mathbb{2}\), respectively). Although the cost function showed that \(\triangledown\) is a reasonable calibration, visual inspection favors \(\circ\) and \(\diamond\). Accepting \(\circ\) as a good fit further supports our conclusions that we can explain the observed seasonality in the estuary-scale SPM distribution without requiring seasonality in biotically induced flocculation and erosion parameters. The seasonality in SPM can be captured by only accounting for seasonality in freshwater discharge and turbulence.

### 3.3.3 Sensitivity analysis of the modeled SPM concentration to \(D_p\) and \(n_f\)

Figure 3.8 shows the sensitivity of the depth-averaged SPM concentration to \(D_p\) and \(n_f\). The results of the SPM concentration in summer and winter are neither quantitatively nor qualitatively changed significantly when varying \(D_p\) (Figs. 3.8a-b) and \(n_f\) (Figs. 3.8c-d). To conclude, our sensitivity analysis of the modeled SPM concentration to \(D_p\) and \(n_f\) shows that our results and conclusions do not change when choosing different parameter values.

### 3.4 Discussion

#### 3.4.1 Model performance and reverse engineering approach

In this contribution, we aim to answer the question whether biotically induced seasonality has a dominant impact on the observed seasonality in SPM on the estuary scale in the Scheldt estuary. We take estuary scale to mean the location and approximate SPM concentration of the ETM, which are captured in our model. On this scale, seasonal differences in parameters representing biotic effects are not necessary. To be specific, any seasonal differences in optimal calibration of \((\lambda, M)\) is within the uncertainty in the observations. Furthermore, choosing between optimally calibrated \((\lambda, M)\) in summer and winter or equal \((\lambda, M)\) in summer and winter only has an \(O(10^{-2})\) effect on the cost function representing goodness-of-fit. Moreover, by lumping all biotic effects of different scale and nature in only two parameters, our conclusions are only valid for the aggregated
Figure 3.8: Sensitivity of the depth-averaged suspended particulate matter (SPM) concentration to the primary particle size $D_p$ in (a) summer and (b) winter, and to the fractal dimension $n_f$ in (c) summer and (d) winter.
biotic impact on a large scale. This may still imply that the effect of individual biotic processes approximately cancel out against each other or are of significant importance locally, for example, in shallow areas.

### 3.4.2 Biotic seasonality in floc size in the Scheldt estuary

Various studies on the seasonality in floc size dynamics in the Scheldt estuary have been reported in the literature. These studies have linked these dynamics to the presence of biota but read different conclusions. We explain the seasonality in floc size by seasonality in freshwater discharge and corresponding SPM concentrations: a larger discharge results in flushing of the SPM concentrations and a larger shear rate locally at the upstream boundary [see Eq. (3.5) and Fig. 3.C.1]. A decrease in SPM concentration and an increase in shear rate result in a decrease in floc size. Hence, it is worth discussing this.

Wartel and Francken (1998) measured in situ floc size between km 40 and 100 using a Benthos plankton camera, which measures in situ flocs ranging in size from 30- to 1625 µm (the LISST 200x instrument we used measures floc sizes ranging from 1 to 500 µm). They found slightly lower averaged floc sizes of approximately 70 µm, but did not measure a longitudinal trend in floc size between km 60-100. Their averaged floc size of approximately 90 µm at km 110 in winter and the absence of a significant seasonality (autumn versus spring) of the floc size at km 100 compare well to our observations. They attributed this slightly larger floc size at km 110 to an increase in organic content and the presence of micro-organisms, which excrete EPS (of which TEP is a subgroup).

In contrast, Chen et al. (2005) measured floc size locally at both a freshwater and brackish tidal flat in the Scheldt estuary. They used the so called ‘pipette-method’ to determine floc sizes. Eisma et al. (1991) showed that observations using this method did not correlate to observations carried out using the in situ Benthos plankton camera. At the freshwater tidal flat (at approximately km 100), the mean floc size increased from approximately 40 µm in winter upto 80 µm in spring. In the brackish tidal flat (at approximately km 20), it increased from approximately 30 µm in winter upto 90 µm in summer. Interestingly, the measured floc sizes in Chen et al. (2005) are significantly smaller in winter compared to our observations and the ones of Wartel and Francken (1998). Chen et al. (2005) also found a significant seasonality in floc size, which they linked to the presence of biota (i.e., organic content).

More recently, Chen et al. (2018) measured in situ floc sizes, again not in the estuary as a whole, but locally in the coastal region at the mouth (i.e., southern North Sea). By using a reverse engineering model approach, they found a significant seasonality in floc strength, which was inversely related to floc break-up. More specifically, they showed that by decreasing the floc strength in winter (Feb. and Mar.) from \(3 \times 10^{-11}\) N to \(1 \times 10^{-11}\) N, their model simulations of floc size significantly improved when compared to the in situ observations. They attributed this seasonality in floc strength to biological effects (i.e., seasonality in TEP), which we did not observe in the Scheldt estuary as a whole (Fig. 3.4b).
To conclude, floc size observations in the Scheldt estuary show different trends in the literature. These differences may be attributed to the usage of different methodologies (e.g., pipette method, in situ Benthos plankton camera) and the high dependency on the system of interest (e.g., main channel, tidal flat, coastal region). Our observations comply with the main trends found by [Wartel and Francken (1998)], who also used an in situ method to measure floc size in the main channel.

### 3.5 Conclusions

In this contribution, we studied the importance of biotically induced seasonality in flocculation and erosion to seasonality in ETM formation applied to the Scheldt estuary. We presented in situ observations of SPM, floc size, Chl-a, and TEP. The SPM and floc size observations show seasonality. Chl-a and TEP observations show no correlation: Chl-a observations show a significant seasonality, whereas we do not detect seasonality in TEP on the estuary scale. These unrelated results make it difficult to use observations of Chl-a and TEP alone as indicators for a potential seasonal impact of biota on the seasonality in ETM formation. Therefore, we combined the observations with a reverse engineering model approach to assess the aggregated impact of biotically induced seasonality in flocculation and erosion on the seasonality in large-scale ETM formation. In the model, this aggregated biotic impact through flocculation and erosion is implicitly parameterized by the ratio of the minimal floc aggregation and floc break-up parameter $\lambda$ and the erosion parameter $M$.

Our model results suggest a weak (~10%) seasonality in both $\lambda$ and $M$, which is within the uncertainty of the SPM observations. Hence, to explain the observed seasonality in large-scale ETM formation, our results suggest that seasonality in $\lambda$ and $M$ is not required on the estuary scale. We capture the seasonality in SPM mainly by seasonality in freshwater discharge and resulting seasonality in SPM concentration and shear rate.

### List of symbols

1. Denotes the region where the first ETM is located
2. Denotes the region where the second ETM is located
3. Denotes the region where the third ETM is located
4. Denotes the optimal $(\lambda, M)$ pair in winter
5. Denotes the first alternative $(\lambda, M)$ pair in winter
6. Denotes the second alternative $(\lambda, M)$ pair in winter

### Latin

- $a$: Availability of easily erodible fine sediment at the sediment bed
Suspended sediment concentration
Subtidal SPM concentration at the Upper Sea Scheldt boundary
Floc size
Primary particle size
Erosion flux of suspended sediment from the sediment bed
Erodibility
Shape factor
Function that describes salinity driven flocculation
Shear rate
Gravitational acceleration
Nondimensional minimal aggregation coefficient
Break-up parameter
Erosion parameter
Erosion parameter in summer
Erosion parameter in winter
Fractal dimension
Order of
Total freshwater discharge
Total freshwater discharge in summer
Total freshwater discharge in winter
Floc Reynolds number (i.e., \(w_s D_f / \nu\))
Salinity
Bed roughness coefficient
Salinity boundary condition at the mouth
Settling velocity of suspended sediment
Minimum settling velocity of suspended sediment
Calibration parameter in postulated tanh salinity distribution
Calibration parameter in postulated tanh salinity distribution in summer
Calibration parameter in postulated tanh salinity distribution in winter
Calibration parameter in postulated tanh salinity distribution
Calibration parameter in postulated tanh salinity distribution in summer
Calibration parameter in postulated tanh salinity distribution in winter
Calibration parameter in postulated tanh salinity distribution
Difference of the optimal \(\lambda\)-values in summer and winter, relative to the corresponding standard deviation (see Eq. 3.3)
Difference of the optimal \(M\)-values in summer and winter, relative to the corresponding standard deviation (see Eq. 3.3)
Ratio of the minimal aggregation and floc break-up parameter
Ratio of the minimal aggregation and floc break-up parameter in summer
Ratio of the minimal aggregation and floc break-up parameter in winter
Dynamic viscosity of water
Kinematic viscosity of water
Density of sediment primary particles
Reference density of water
Appendix 3.A: floc size using various averaging methods

To show that the estimated floc size is highly sensitive to the averaging method, we use two additional averaging methods: 1. the same averaging method as proposed by Sequoia Scientific (2019) but weighting the midpoint instead of the class number and 2. the median floc size by mass instead of volume. The difference between weighting by mass instead of volume depends on the structure of the flocs. Using our reference floc model assumption (fractal dimension is 2), averaging by mass is equivalent to averaging by area (Bowers et al., 2017). So, we divided the volume concentration of each size class by the corresponding midpoint.

Figure 3.A.1 shows the observed depth- and time-averaged floc size measured in summer and winter 2018 using the averaging method proposed by Sequoia Scientific (2019) (circles), weighting the midpoint instead of the size class number (triangles), and the time- and depth-averaged median floc size by mass (rectangles). The error bars depict the standard error of the depth-averages corresponding to the various campaigns. Using other averaging methods may increase (weighting the midpoint by volume) or decrease (weighting by mass) the overall floc size. However, using all three averaging methods, we observe seasonality in the upstream region of the Scheldt estuary. We thus conclude that, on the estuary scale (we have a local anomaly near km 60 and km 80, depending on the averaging method), the floc sizes only show seasonality in the upstream part of the Scheldt estuary (> 120 km).

Appendix 3.B: salinity profile

Following Warner et al. (2005), we fit the salinity data measured in summer and winter in the Scheldt estuary to the following postulated salinity distribution:

\[
\frac{s_{\text{sea}}}{2} \left( 1 - \tanh \frac{x - x_{\text{Sal.}}^{c}}{x_{\text{Sal.}}^{L}} \right),
\]

where \(s_{\text{sea}}\) is the salinity boundary condition at the mouth and \(x_{\text{Sal.}}^{c}\) and \(x_{\text{Sal.}}^{L}\) are further undefined calibration parameters. Figure 3.B.1 shows the salinity data and corresponding

\[\text{Figure 3.A.1}\]

\[\text{Figure 3.B.1}\]
Figure 3.A.1: Measured time- and depth-averaged floc size in summer (red) and winter (blue) using the averaging method proposed by Sequoia Scientific (2019) (circles), weighting the midpoint instead of the size class number (triangles), and the time- and depth-averaged median floc size by mass (rectangles). The error bars depict the standard error of the estimates corresponding to the various campaigns.

data fit in summer and winter. Table 3.1 lists the corresponding parameter values.

Appendix 3.C: shear rate in the flocculation model

Following Pejrup and Mikkelsen (2010) the shear rate $G$ reads

$$G(x, \tilde{z}_1) = \sqrt{\frac{\langle u_s \rangle^3 (1 - \tilde{z}_1)}{\nu H \kappa' \tilde{z}_1}},$$

in which $x$ is the coordinate in the longitudinal direction, $\kappa'$ is the Von Karman constant, $\nu$ is the kinematic viscosity of water, $H$ is the water depth, $\tilde{z}_1 = z_1 / H$ is the relative water depth, and $\langle u_s \rangle$ is the subtidal friction velocity. To ensure that the model captures the impact of seasonality in river discharge on $G$, we define the subtidal friction velocity following Horemans et al. (2020) as

$$\langle u_s \rangle(x) = \frac{\kappa'}{\ln \frac{z_1(x)}{s_{f_0}}} \left( |u^0(x, z_1) + u_{riv.}(x, z_1)| \right),$$

where $z_1$ is the distance in the logarithmic layer above the bed, $s_{f_0}$ the bed roughness coefficient, $u^0$ the leading-order, longitudinal flow velocity, which consist of an M2 tidal signal (Chernetsky et al., 2010; Dijkstra et al., 2017), and $u_{riv.}$ the (subtidal) longitudinal, river-induced flow velocity. Consequently, an increase in freshwater discharge (cf. $u_{riv.}$) results in an increase in shear rate $G$. Following Horemans et al. (2020) we approximate $G$ by its value at $z_1 = H/2$ and use this value $G(x, \tilde{z}_1 = 1/2)$ as a proxy for the whole water column, which is acceptable because vertical variations in $G$ are relatively small.
in the logarithmic layer (typically < 10% in our case study). Moreover, we neglect tidal variation in \( G \) which is, within the scope of this study, acceptable because the seasonality in tidal variability in our case study is small; the main difference in longitudinal water velocity between summer and winter is the riverine influence, locally at the upstream boundary, which is thus captured by the model. Figure 3.C.1 shows the shear rate in both summer and winter conditions.

The increase of freshwater discharge in winter increases the shear rate locally at the upstream boundary. Further downstream, between km 100-140, it decreases the shear rate slightly because of a decrease of the tidal energy in this region.

**Appendix 3.D: perturbation approach for \( n_f \neq 2 \)**

We extend the flocculation model for general \( n_f \). This requires a reanalysis of the results presented in Horemans et al. (2020). However, the same semianalytical perturbation approach can be applied. At leading order \( O(1) \), a careful analysis shows that only the second term in the result of the leading order settling velocity \( w_s^0 \) presented in Horemans et al. (2020) changes:

\[
   w_s^0 = \beta c^0 + \frac{D_p}{D_f} (w_s^0)^{\frac{n_f-2}{2}},
\]

(3.7)

with

\[
   \tilde{D}_f = 18^{\frac{1}{n_f-1}} \left( \frac{\mu D_p^{n_f-3}}{g(\rho_s - \rho_w)} \right)^{\frac{1}{n_f-1}},
\]

(3.8)

and where \( \beta = (\rho_s - \rho_w)gk'_A/(18\mu_f \rho_s \sqrt{Gk_B}) \), with \( k'_A \) being the aggregation parameter, and \( c^0 \) is the leading order SPM concentration. For \( n_f = 2 \), the second term in Eq. (3.7)
Figure 3.C.1: Modeled shear rate in summer and winter. The increase of freshwater discharge in winter increases the shear rate locally at the upstream boundary. Further downstream between km 100-140, it decreases the shear rate slightly due to a decrease of tidal energy in this region.

is a constant value, being the settling velocity of the primary particles [Horemans et al., 2020]. For \( n_f \neq 2 \), this second term is a function of both time and space. Consequently, at first order \( O(\epsilon) \) (with \( \epsilon \) being the small perturbation parameter), the result becomes more complex but can still be easily solved to the first order settling velocity \( w_s^1 \):

\[
\begin{align*}
A & = \frac{Gk' A 18^{n_f-1} \left( \frac{1}{D_p} \right)^{6-2n_f} \left( 18^{n_f-1} \left( \frac{\mu D_p^{n_f-3}}{g(\rho_s-\rho_w)} \right)^{\frac{1}{n_f-1}} \right)^{-2n_f} \left( \frac{\mu D_p^{n_f-3}}{g(\rho_s-\rho_w)} \right)^{\frac{3}{n_f-1}}}{(fs\rho_s)^2} \quad \text{and} \\
B & = \frac{G^{1.5} k_B 18^{n_f-1} \left( \frac{1}{D_p} \right)^{3-n_f} \left( 18^{n_f-1} \left( \frac{\mu D_p^{n_f-3}}{g(\rho_s-\rho_w)} \right)^{\frac{1}{n_f-1}} \right)^{-n_f} \left( \frac{\mu D_p^{n_f-3}}{g(\rho_s-\rho_w)} \right)^{\frac{2}{n_f-1}}}{fs\rho_s}.
\end{align*}
\]
Appendix 3.E: primary particle size and density

Within the OMES campaign, we took 1 L samples from a bucket sample conducted at the water surface to estimate primary particle size (Maris et al., 2021c). Samples were collected once a month at each station, which resulted in three samples per station for both summer and winter. At km 58, we exceptionally have only two samples per season.

Primary particle size distribution was measured using a Mastersizer2000. The detection range of the Mastersizer2000 is 0.02 to 2000 µm. After shaking the 1 L samples intensively, the samples were decanted in the dispersion unit of the Mastersizer2000 until the obscuration reached a value of approximately 10%. We used the default absorption and refractive index (i.e., 0.1 and 1.52, respectively) and standard Lorenz-Mie method for spherical particles. The samples were stirred and pumped with a frequency of 750 and 2000 rpm, respectively, using tap water as a dispersant. Using this primary particle size distribution, we calculate the primary particle size corresponding to summer and winter using two averaging procedures. Figure 3.E.1 shows the primary particle size using a standard geometric average method often referred to as $D[1,0]$. The corresponding system averaged value is 14.2 and 14.9 µm in summer and winter, respectively. The error bars represent the (Gaussian) standard error. Based on these estimates, we chose a primary particle size of 14 µm in our model.

The organic fractions of SPM, which have a density of $\sim 1000$ kg m$^{-3}$, can reach up to 30% in Scheldt estuary (Maris and Meire, 2017), and thus the organic fraction may have a relative impact via primary particle density (i.e., 2650 kg m$^{-3}$) according to Fall et al. (2021) of

\[
\Delta \rho_s = 1 - \frac{0.3 \cdot 1000 + 0.7 \cdot 2650}{2650} \sim 20\%
\]  

(3.12)

This sensitivity was tested, and for such a decrease in $\rho_s$, we found a slight land-inward shift of the ETM in both summer and winter. We also observed this land-inward shift for
decreasing $\lambda$ (Fig. 3.7). This is to be expected because a decrease in $\rho_s$ is equivalent to a decrease in $\lambda$, which follows from the factor $(\rho_s - \rho_w)/\rho_s$ in Eq. (3.2).

Table 3.F.1: Parameter values used in our model study in the Scheldt estuary in summer and winter conditions.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A^0$</td>
<td>$M_2$ water level amplitude at x=0</td>
<td>1.77 m</td>
</tr>
<tr>
<td>$A^1$</td>
<td>$M_4$ water level amplitude at x=0</td>
<td>0.14 m</td>
</tr>
<tr>
<td>$\phi^1$</td>
<td>$M_4$ water level phase relative to $M_2$ tide at x=0</td>
<td>-1.3 deg</td>
</tr>
<tr>
<td>$Q_{\text{summer}}$</td>
<td>Total river discharge in summer (calibrated)</td>
<td>40 m$^3$ s$^{-1}$</td>
</tr>
<tr>
<td>$Q_{\text{winter}}$</td>
<td>Total river discharge in winter (calibrated)</td>
<td>233 m$^3$ s$^{-1}$</td>
</tr>
<tr>
<td>$c_{\text{sea}}$</td>
<td>Depth-averaged subtidal SPM concentration at x=0</td>
<td>0.06 kg m$^{-3}$</td>
</tr>
<tr>
<td>$c_{\text{upper Sea Scheldt}}$</td>
<td>Subtidal SPM concentration at the Upper Sea Scheldt boundary (source)</td>
<td>0.035 kg m$^{-3}$</td>
</tr>
<tr>
<td>$K_h$</td>
<td>Horizontal eddy diffusivity coefficient</td>
<td>$100$ m$^2$ s$^{-1}$</td>
</tr>
<tr>
<td>$K_v$</td>
<td>Vertical eddy diffusivity coefficient</td>
<td>$3.1 \times 10^{-2}$ m$^2$ s$^{-1}$</td>
</tr>
<tr>
<td>$M_{\text{summer}}$</td>
<td>Erosion parameter in summer (calibrated)</td>
<td>$2.7 \times 10^{-3}$ s m$^{-1}$</td>
</tr>
<tr>
<td>$M_{\text{winter}}$</td>
<td>Erosion parameter in winter (calibrated)</td>
<td>$3.0 \times 10^{-3}$ s m$^{-1}$</td>
</tr>
<tr>
<td>$\sigma_\rho$</td>
<td>Prandtl Schmidt number</td>
<td>0.5</td>
</tr>
<tr>
<td>$s_{f_0}$</td>
<td>Bed roughness coefficient (calibrated)</td>
<td>$4.22$ mm s$^{-1}$</td>
</tr>
<tr>
<td>$k_{\text{min.}}$</td>
<td>Nondimensional minimal aggregation coefficient</td>
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</tr>
<tr>
<td>$f_s$</td>
<td>Shape factor</td>
<td>$\pi/6$</td>
</tr>
<tr>
<td>$n_f$</td>
<td>Fractal dimension of flocs</td>
<td>2</td>
</tr>
<tr>
<td>$q$</td>
<td>Calibration parameter in flocculation model</td>
<td>0.5</td>
</tr>
<tr>
<td>$p$</td>
<td>Calibration parameter in flocculation model</td>
<td>1</td>
</tr>
<tr>
<td>$D_p$</td>
<td>Diameter primary mud particles</td>
<td>$14 \times 10^{-6}$ m</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Dynamic viscosity</td>
<td>0.0010518 Pa s</td>
</tr>
<tr>
<td>$\rho_s$</td>
<td>Density of SPM primary particles</td>
<td>2650 kg m$^{-3}$</td>
</tr>
<tr>
<td>$\rho_w$</td>
<td>Reference density of water</td>
<td>1000 kg m$^{-3}$</td>
</tr>
<tr>
<td>$\lambda_{\text{summer}}$</td>
<td>Ratio of minimal aggregation and floc break-up parameter in summer (calibrated)</td>
<td>$72.5 \times 10^{-6}$ s$^{-1/2}$ m$^2$</td>
</tr>
<tr>
<td>$\lambda_{\text{winter}}$</td>
<td>Ratio of minimal aggregation and floc break-up parameter in winter (calibrated)</td>
<td>$65.9 \times 10^{-6}$ s$^{-1/2}$ m$^2$</td>
</tr>
<tr>
<td>$s_{\text{sea}}$</td>
<td>Salinity boundary condition at the mouth</td>
<td>28.9 psu</td>
</tr>
<tr>
<td>$x_{c, \text{sal, summer}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in summer</td>
<td>51.9 km</td>
</tr>
<tr>
<td>$x_{\text{sal, summer}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in summer</td>
<td>31.2 km</td>
</tr>
<tr>
<td>$x_{c, \text{sal, winter}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in winter</td>
<td>38.3 km</td>
</tr>
<tr>
<td>$x_{\text{sal, winter}}$</td>
<td>Calibration parameter in sigmoid salinity distribution in winter</td>
<td>24.8 km</td>
</tr>
</tbody>
</table>
Bibliography


Chapter 4

The impact of temporal variability in SPM on phytoplankton

Abstract

Phytoplankton primary production (PP) in turbid estuaries is often limited by light availability. Two important factors altering light climate are solar irradiance at the water surface and exponential light extinction coefficient within the water column. Additionally, the depth of the water body changes the light climate and corresponding PP by altering the ratio of the euphotic and mixing depth in a well-mixed estuary. These three parameters are highly variable yet are often assumed to be constant by both experimental scientists and modelers because of a lack of data or to reduce complexity. Because assuming constant parameters introduces an error, we utilize an idealized model of depth-integrated PP to analyze the (individual) impact of temporal variability in these three parameters. We only consider the main tidal and solar constituents in temporal variability of the forcings and apply a second-order moment approximation to analyze the bias introduced to time-averaged PP estimates by neglecting temporal fluctuations. We demonstrate that the sign and magnitude of this bias are system-specific and depend on two nondimensional parameters that characterize the system. The first is equivalent to the ratio of mixing and photic depth. The second accounts for typical incident irradiance and the photosynthetic parameters of the phytoplankton population present. To demonstrate the applicability of our approach, we apply the model to two cases in the Scheldt estuary (Belgium) in the brackish and freshwater part. In the first application, we study the impact of fluctuations on phytoplankton in dynamic equilibrium, where biomass is assumed to be constant. We show that variability in solar irradiance has the largest impact on time-averaged PP in dynamic equilibrium, resulting in a 30 % decrease compared to time-invariant forcing. By comparing with a numerical integrator, we show that a second-order moment approximation correctly predicts the order of magnitude of the impact of temporal variability of the individual parameters. In the second application, we study the impact of fluctuations on unbounded exponential phytoplankton growth. Also here, fluctuations in solar irradiance have the largest impact and lead to a significant decrease in exponential growth. In this case study, we show that temporal fluctuations delay the onset of the biomass by two weeks and decrease the biomass by a factor of 14 after two weeks compared to time-invariant forcing. Additionally, we show that the temporal fluctuations induce low-frequency variability in phytoplankton biomass with similar periodicity as the spring-neap cycle, making it challenging to observe these phenomena in real-world time series.

4.1 Introduction

Primary production (PP) in estuaries and the corresponding phytoplankton biomass dynamics result from a complex interaction of physical and biological processes (Alpine and Cloern, 1992; Dijkstra et al., 2019). Examples of factors that directly impact PP and phytoplankton biomass dynamics are physical-chemical aspects such as river flushing (Filardo and Dunstan, 1985; Liu and de Swart, 2015), temperature (Eppley, 1972), salinity (Lucas et al., 1998), nutrients (Tilman et al., 1982; Cira et al., 2016), and light
availability (Sverdrup, 1953; Desmit et al., 2005), or biological factors such as grazing (Lionard et al., 2005). Due to these complex interactions, we expect both strong high-frequency (∼hours⁻¹-days⁻¹) and low-frequency (∼weeks⁻¹-months⁻¹) dynamics generated by water flow and seasonality in, for example, water temperature, light availability, and freshwater discharge.

To acquire insight into the complex interacting processes affecting PP and phytoplankton biomass dynamics, both intensive monitoring campaigns (Maris and Meire, 2016) and a broad range of models have been set up, ranging from numerically-costly three-dimensional models (Chen and Mynett, 2006; Chao et al., 2010), idealized two-dimensional models (McSweeney et al., 2017), one-dimensional models (Brinkman, 1993; Soetaert and Herman, 1995; Vanderborght et al., 2002; Volta et al., 2016; Liu et al., 2018) to zero-dimensional models (Desmit et al., 2005; Cox et al., 2009).

Due to the high complexity of PP and phytoplankton biomass dynamics, idealized models are particularly well suited because they simplify the interpretation of the model results and reduce computational costs, allowing for long-term simulations and extensive sensitivity analyses. More specifically, often model parameters are considered constant to remove nonlinearities and allow for analytical solution procedures (Dijkstra et al., 2017) instead of computationally costly iterative solution methods that tidally resolve the phytoplankton dynamics (Arndt et al., 2011). By using constant model parameters, the impact of temporal variability, which is clearly present in factors affecting PP, is neglected. Only a few studies focus on the individual effect of this, sometimes neglected, high-frequency temporal variability in the factors affecting PP and phytoplankton biomass dynamics. The few studies that do exist are limited to specific examples (Desmit et al., 2005).

Therefore, this study aims to construct a generic, idealized model that allows us to study the impact of temporal fluctuations of individual tidal and solar variables of PP and the corresponding phytoplankton dynamics on an hourly-daily time scale.

We focus on well-mixed, turbid- and nutrient-rich estuarine systems, allowing to analytically compute depth-averaged PP and only consider light limitation, thereby excluding nutrient depletion and temperature dependence. The biological impact of grazing is implicitly included in a constant mortality rate and calibrated to Chlorophyll-a (Chl-a) observations. Moreover, we assume tide-dominated systems and only include periodic temporal variability due to the tides and solar cycle. We focus on the three important factors altering the light climate on the hourly-daily time scale: solar irradiance at the water surface, exponential light extinction coefficient within the water column, and depth of the water body, which changes the ratio of euphotic and mixing depth in a well-mixed estuary.

To model the impact of a changing light climate on phytoplankton biomass growth, we use the Platt light limitation function (Platt et al., 1980). By assuming a well-mixed estuary, we focus on depth-averaged Platt light limitation (DAP), allowing for an analytical solution approach. Moreover, to explicitly compute the impact of temporal variability of the individual parameters and correct for the usage of averaged parameter values, we
use the analytical, second-order up-scaling method often referred to as the moment approximation. This method is based on a second-order Taylor expansion. In the past, the moment approximation has been successfully applied to terrestrial PP (Bolker and Pacala, 1997; Wirtz, 2000). However, terrestrial PP significantly differs from estuarine PP because the light climate in estuarine systems also depends on the tidal variability in depth, suspended sediment concentration, and its corresponding complex temporal variability.

To show the applicability of our approach, we work out two applications in the Scheldt estuary (Belgium). Firstly, we apply our framework to a time-averaged production assuming dynamic equilibrium (application 1). This reflects the situation where phytoplankton biomass is top-down controlled by grazing organisms (e.g., zooplankton), which is often the case in real ecosystems, and remains relatively constant during a large part of the season when primary productivity rates are positive. Such dynamic equilibrium is commonly observed after an initial spring bloom. Secondly, we utilize our idealized model to study phytoplankton dynamics at the onset of a phytoplankton bloom with unbounded biomass growth (application 2).

In this paper, we first introduce the phytoplankton model and DAP function, the corresponding moment approximation, and the two applications in the Scheldt estuary in Section 4.2. In Section 4.3, we present the generic results of the moment approximation and apply these results to our two applications in the Scheldt estuary. Finally, we discuss our results in Section 4.4 and summarize our conclusions in Section 4.5.

4.2 Material and methods

This section presents the depth-integrated model for phytoplankton biomass dynamics and corresponding DAP function \( \Lambda \) that includes our three main parameters of interest: the solar irradiance at the water surface \( E_0 \), exponential light extinction coefficient \( k_d \), and water depth \( d \). Next, we apply a second-order moment approximation to the DAP function \( \Lambda \), which forms the basis of our subsequent analysis. Finally, we work out applications 1 and 2 in the Scheldt estuary using our framework and present the corresponding observations. To acquire an analytical solution in application 2, we only include sinusoidal (cf. periodic) temporal variability in \( E_0 \), \( k_d \), and \( d \) (see Section 4.2.4). A conceptual diagram of our approach is illustrated in Fig. 4.1.

4.2.1 Phytoplankton model

To describe PP in a tidal, well-mixed system, we start from the following one-dimensional, differential equation for depth-averaged phytoplankton biomass concentration growth rate
Figure 4.1: Conceptual diagram of our idealized model. In a well-mixed, light-limited tidal system, gross primary production (GPP) is limited by the DAP function $\Lambda(t)$, which is influenced by solar irradiance and tides. The tides also directly alter biomass dynamics through advection and diffusion in the longitudinal direction. We apply the moment approximation to $\Lambda(t)$ and idealize temporal variability in $E_0$, $k_d$, and $d$. This simplification results in an analytical expression that relates temporal variability of GPP and corresponding phytoplankton biomass dynamics to variability in $E_0$, $k_d$, and $d$. We apply the idealized model to time-averaged GPP in dynamic equilibrium (application 1) and a phytoplankton bloom (application 2) in the Scheldt estuary (Belgium).

(units kg m$^{-3}$ s$^{-1}$) (Langdon, 1993; Desmit et al., 2005):

$$\frac{\partial \tilde{B}(x,t)}{\partial t} + \frac{1}{d} \int_0^d \left[ \frac{\partial}{\partial x} (\tilde{u} \tilde{B}) + \frac{\partial}{\partial x} \left( K_h \frac{\partial}{\partial x} \tilde{B} \right) \right] dz$$

change in $\tilde{B}(t)$ due to advection/diffusion

$$= \tilde{B} \cdot P_{\text{max}} \cdot \left( \frac{1}{d} \int_0^d \tilde{\Lambda} dz \right) \cdot (1 - \rho_B - m)$$

where $\tilde{B}$ is the depth-averaged phytoplankton biomass concentration, $x$ is the distance from the mouth, $z$ is the distance from the water surface in the downward direction, $t$ is the time, $\rho_B$ is the respiration and excretion coefficient, $m$ is the mortality coefficient, $P_{\text{max}}$ is the maximum photosynthetic rate, $\tilde{\Lambda}$ is a GPP light limitation function, $\tilde{u}$ is the (vertically homogeneous) water velocities in the longitudinal direction, and $K_h$ is the longitudinal eddy diffusivity coefficient.
In Eq. (4.1), we assume vertically, well-mixed systems resulting in a depth-independent phytoplankton biomass concentration \( \tilde{B}(x, t) \). Furthermore, we exclude nutrient limitation since we consider PP in nutrient replete, light-limited systems. Moreover, we also exclude temperature dependence and implicitly included grazing of phytoplankton in the mortality coefficient \( m \). Crucially, we assume balanced-growth conditions: growth and photosynthesis are considered never to be decoupled on the time scale of our application (i.e., hours-days). All photosynthesis is used for biomass production \( \{\text{Berman-Frank and Dubinsky, 1999}\} \). Such decoupling is typical in nutrient limitation situations, which we assume not to occur in the turbid, nutrient replete estuaries under study. Furthermore, we assume that phytoplankton species surviving in turbid conditions are adapted to harvest every photon possible and are thus able to store excess photosynthesized material rather than excrete photosynthetically-derived dissolved organic carbon compounds (PDOC). Consequently, also during the short periods that cells reside in the upper part of the water column where light availability is maximal, growth is balanced. To model light limitation, we use the Platt light limitation function \( \tilde{\Lambda} \) (Platt et al., 1980):

\[
\tilde{\Lambda} = \left\{ 1 - \exp \left[ -\frac{\alpha P_{\text{max}}}{I} \right] \right\},
\]

in which \( \alpha \) is the photosynthetic efficiency and \( I \) is the light intensity. Using the Lambert-Beer expression for exponential light attenuation in a water body, Eq. (4.2) yields

\[
\tilde{\Lambda}(z, t) = \left\{ 1 - \exp \left[ -\tilde{\beta} \exp(-\tilde{\bar{z}}) \right] \right\},
\]

in which \( \tilde{\beta} \) and \( \tilde{\bar{z}} \) are dimensionless numbers defined as

\[
\tilde{\beta} = \frac{\alpha E_0}{P_{\text{max}}} \quad \text{and} \quad \tilde{\bar{z}} = k_d \cdot z.
\]

Because the system is assumed to be well-mixed, the total biomass production \( \text{GPP}(t) \) integrated over the water depth at a fixed location \( x \) reads as

\[
\text{GPP}(t) = B(t) \cdot P_{\text{max}} \cdot \Lambda(t),
\]

with \( \Lambda \) being the depth-averaged Platt expression:

\[
\Lambda(t) = \frac{1}{d} \int_0^d \tilde{\Lambda}(z, t) dz.
\]

For an analytical solution of the DAP, the reader is referred to Appendix 4.A. In the following section, we apply the moment approximation to \( \Lambda(t) \) to study the impact of temporal variability in the individual factors \( E_0, k_d, \) and \( d \) and their interactions on PP.

### 4.2.2 Moment approximation

Temporal variability in \( E_0, k_d, \) and \( d \) impacts phytoplankton dynamics through the time integral of the DAP. This is our starting point to assess the impact of tidal and solar
fluctuations on GPP using the moment approximation. As a first step, we add a (further unspecified) time dependence to the variables in the DAP:

\[ E_0(t) = \langle E_0 \rangle + E'_0(t), \quad (4.7) \]
\[ k_d(t) = \langle k_d \rangle + k'_d(t), \quad \text{and} \]
\[ d(t) = \langle d \rangle + d'(t), \quad (4.9) \]

in which the angle brackets denote ‘time-averaged value of’. This construction allows us to perform a Taylor expansion of the DAP. Consequently, we can estimate the impact of temporal variability of the individual parameters on phytoplankton dynamics and their interactions. Also, it is easy to revert to the time-independent result by making \( E'_0(t), k'_d(t), \) and \( d'(t) \) equal to zero. As a second step, we apply a Taylor expansion to the analytic expression of the DAP around the time-averaged values in Eqs. (4.8) and (4.9). For a detailed analysis, we refer the reader to Appendix 4.B. Upto second order, this Taylor expansion reads as

\[
\Lambda(t) \approx \text{mfun}_0(\beta, \lambda) \times \left\{ 1 + \right. \\
+ \text{mfun}_\kappa(\beta, \lambda) \cdot [\kappa(t) + \delta(t)] \\
+ \text{mfun}_\epsilon(\beta, \lambda) \cdot \epsilon(t) \\
+ \text{mfun}_\kappa^2(\beta, \lambda) \cdot [\kappa^2(t) + \delta^2(t)] \\
+ \text{mfun}_\epsilon^2(\beta, \lambda) \cdot \epsilon^2(t) \\
+ \text{mfun}_{\kappa\epsilon}(\beta, \lambda) \cdot [\epsilon(t) \cdot \kappa(t) + \epsilon(t) \cdot \delta(t)] \\
+ \text{mfun}_{\kappa\delta}(\beta, \lambda) \cdot [\delta(t) \cdot \kappa(t)] \left. \right\}, \tag{4.10}
\]

in which \( \kappa, \epsilon, \) and \( \delta \) are dimensionless relative fluctuations defined as

\[
\epsilon(t) = \frac{E'_0(t)}{\langle E_0 \rangle}, \quad \kappa(t) = \frac{k'_d(t)}{\langle k_d \rangle}, \quad \text{and} \\
\delta(t) = \frac{d'(t)}{\langle d \rangle} \quad (4.11)
\]

and \( \beta \) and \( \lambda \) are defined as

\[
\beta = \frac{\alpha \langle E_0 \rangle}{P_{\text{max}}} \quad \text{and} \quad \lambda = \langle k_d \rangle \cdot \langle d \rangle. \quad (4.12)
\]

The mfun functions determine the impact of temporal variability in \( E_0, k_d, \) and \( d \) on the DAP upto second order and are defined in Appendix 4.B. By definition, the function mfun\(_0\) is the DAP without temporal parameter fluctuations. The other terms in Eq. (4.10) quantify the impact of temporal variability of the individual forcings (terms \( \sim \) mfun\(_\kappa\), mfun\(_\epsilon\), mfun\(_\kappa^2\), and mfun\(_\epsilon^2\)) and their interactions (terms \( \sim \) mfun\(_{\kappa\epsilon}\) and mfun\(_{\kappa\delta}\)). By construction of \( \Lambda, \kappa \) and \( \delta \) come with identical mfun functions:
The mfun functions are determined by two dimensionless numbers $\lambda$ and $\beta$, which characterize the system under study and have a clear physical interpretation. $\lambda$ is a measure for the photic properties of the water column. Turbid and deep systems have a large $\lambda$, while transparent and shallow systems correspond with a small $\lambda$. For fully mixed systems, $\lambda$ is proportional to the ratio of the mixing depth and euphotic depth, that is, the depth at which the light intensity is 1% of the incident light (Sverdrup, 1953; Kromkamp and Peene, 1995; Desmit et al., 2005). This ratio determines the time phytoplankton spends in the dark and thus affects net PP. Often a critical depth is defined: net positive PP can only occur when the depth of the mixed layer is less than the critical value (Sverdrup, 1953). Our results confirm that this ratio is an important characteristic of the system, not only controlling the potential for net PP but also controlling the response of PP to temporal variability in light climate. The second dimensionless number $\beta$ increases with increasing solar irradiance and photosynthetic efficiency. Consequently, for a fixed $P_{\text{max}}/\alpha$ ratio, an equatorial system (cf. more solar irradiance) typically corresponds to a larger value of $\beta$ than a system in the northern hemisphere. In the following, we apply our approach to two case studies in the Scheldt estuary using real observations.

### 4.2.3 Application 1: time-averaged GPP

In the first application, we use our idealized model to analyze the impact of temporal fluctuations on time-averaged GPP in case phytoplankton biomass is top-down controlled by grazing organisms (e.g., zooplankton) and can be considered constant, although $\text{GPP} > 0$. Consequently, time-averaged GPP becomes proportional to time-averaged DAP:

$$\left\langle \text{GPP}(t) \right\rangle = B \cdot P_{\text{max}} \cdot \left\langle A(t) \right\rangle.$$

\[\text{(4.16)}\]
We compute $\langle \Lambda(t) \rangle$ using Eq. (4.10):

$$
\langle \Lambda(t) \rangle \approx \text{mfun}_0(\beta, \lambda) \left\{ 1 + 
+ \text{mfun}_{\kappa^2}(\beta, \lambda) \left[ \text{Var}[k_d(t)] + \text{Var}[d(t)] \right]
+ \text{mfun}_{\epsilon^2}(\beta, \lambda) \cdot \text{Var}[E_0(t)]
+ \text{mfun}_{\kappa \epsilon}(\beta, \lambda) \left[ \text{Cov}[E_0(t), k_d(t)] + \text{Cov}[E_0(t), d(t)] \right]
+ \text{mfun}_{\kappa^2 \epsilon}(\beta, \lambda) \cdot \text{Cov}[k_d(t), d(t)] \right\},
$$

(4.17)

in which $\text{Var}$ and $\text{Cov}$ are the relative statistical identities variance and covariance. For example, we have

$$
\text{Var}[k_d(t)] = \langle \kappa^2(t) \rangle \quad \text{and} \quad \langle \epsilon(t) \cdot \kappa(t) \rangle.
$$

(4.18, 4.19)

As a result of the definitions in Eqs. (4.8) and (4.9), the time averages of the $\sim \text{mfun}_{\kappa^2}$ and $\text{mfun}_{\epsilon^2}$-terms in Eq. (4.10) disappear in Eq. (4.17).

Eq. (4.17) reveals an important interpretation of the second-order mfun functions; variability in $k_d$, $d$, and $E_0$ results in a systematic upward or downward shift in its time-average $\langle \Lambda(t) \rangle$. The second-order mfun functions represent the factors by which the variances and covariances have to be multiplied to calculate the magnitude of this shift. Moreover, the mfun functions define the sign of the shift in $\langle \Lambda(t) \rangle$ and corresponding $\langle \text{GPP} \rangle$. The magnitude is both determined by the average system characteristics ($\lambda$, $\beta$) and the temporal variability in $E_0$, $k_d$, and $d$, which define the magnitude of the variance and covariance factors in Eq. (4.17), and is thus different when considering typical summer or winter conditions.

Therefore, we apply our idealized model to both a summer (10-06-2013 until 02-07-2013) and winter (20-02-2013 until 13-03-2013) case in the Scheldt estuary. Figures 4.2 and 4.3 show the high-frequency observations of $E_0$, turbidity, and $d$ we use in our first application. The observations were measured in the Scheldt estuary in the brackish region at Kruibeke (~85 km from the mouth). We choose Kruibeke because here our assumptions (e.g., well-mixed, light-limited) are valid and a long-term record of continuous time series of turbidity exists (Cox et al., 2015).

The turbidity time series were obtained using a YSI 6600 multiparameter probe, equipped with an optical turbidity sensor (YSI6136). Turbidity is expressed in NTU (Nephelometric Turbidity Units). Light from the emitter enters the sample and scatters off particles in the water. The light, scattered at 90 degrees, enters a detector fiber and
Table 4.1: Summarized values corresponding to the time series from Figs. 4.2 and 4.3. $N$ is the number of measurements of $E_0$, $k_d$, and $d$. Var and Cov are the relative variance and covariance as defined in Eqs. (4.18) and (4.19).

<table>
<thead>
<tr>
<th></th>
<th>Summer case</th>
<th>Winter case</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N$</td>
<td>3000</td>
<td>3000</td>
</tr>
<tr>
<td>$\langle E_0 \rangle$</td>
<td>171 J m$^{-2}$ s$^{-1}$</td>
<td>70.0 J m$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>$\text{Var}[E_0]$</td>
<td>1.633</td>
<td>2.704</td>
</tr>
<tr>
<td>$\text{Cov}[E_0, k_d]$</td>
<td>0.043</td>
<td>-0.021</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>92.6</td>
<td>95.2</td>
</tr>
<tr>
<td>$\langle k_d \rangle$</td>
<td>10.8 m$^{-1}$</td>
<td>11.1 m$^{-1}$</td>
</tr>
<tr>
<td>$\text{Var}[k_d]$</td>
<td>0.099</td>
<td>0.143</td>
</tr>
<tr>
<td>$\text{Cov}[k_d, d]$</td>
<td>0.036</td>
<td>0.037</td>
</tr>
<tr>
<td>$\beta$</td>
<td>3</td>
<td>1.05</td>
</tr>
<tr>
<td>$\langle d \rangle$</td>
<td>8.60 m</td>
<td>8.60 m</td>
</tr>
<tr>
<td>$\text{Var}[d]$</td>
<td>0.049</td>
<td>0.045</td>
</tr>
<tr>
<td>$\text{Cov}[d, k_d]$</td>
<td>0.036</td>
<td>0.037</td>
</tr>
</tbody>
</table>

Table 4.2: Overview of parameter values used in this paper. The values are based on the literature (Kromkamp and Peene, 1995, 2005; Cox et al., 2010; Maris and Meire, 2016).

<table>
<thead>
<tr>
<th>$P_{\text{max}}$</th>
<th>5 mg C (mg Chl-a)$^{-1}$ h$^{-1}$</th>
<th>C:Chl-a</th>
<th>25 mg C (mg Chl-a)$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha$</td>
<td>0.21 mg C (mg Chl-a)$^{-1}$ h$^{-1}$. (J m$^{-2}$ s$^{-1}$)$^{-1}$</td>
<td>$B_0$</td>
<td>775 µg L$^{-1}$</td>
</tr>
</tbody>
</table>

is measured by a photodiode (YSI, 2017a). The turbidity probe was calibrated in the laboratory using calibration liquid YSI6074 and demineralized water as a zero reference. To compute $k_d$ [m$^{-1}$] from turbidity [NTU], we used the following expression (Cox et al., 2017):

$$k_d = 0.39(2.1\text{NTU}^{-1} \cdot \text{turb.} - 1.82)^{0.64} \text{m}^{-1}. \quad (4.20)$$

Data sets of turbidity and depth at Kruibeke were obtained from waterinfo.be (Waterinfo.be, 2019). We used depth recordings from the tidal gauge at Hemiksem, located approximately 4 km upstream from Kruibeke. Incident irradiance data $E_0(t)$ was obtained from the Royal Meteorological Institute (RMI) of Belgium (RMI, 2013) in Stabroek (~25 km from Kruibeke and 50 km from Schellebelle, the other case study site, see below). The sampling interval of $E_0$, turbidity, and depth was 10, 15, and 15 min, respectively. A summary of the parameter values corresponding to the data sets is presented in Table 4.1. Other parameter values used in our calculations are summarized in Table 4.2. These parameters were based on the literature (Kromkamp and Peene, 1995, 2005; Cox et al., 2010; Maris and Meire, 2016). To assess the accuracy of the moment approximation up to second-order terms, we compare the results to the output of a numerical integrator of DAP, making use of the same time series.
Figure 4.2: (a)-(b) Turbidity and (c)-(d) water depth measured in the Scheldt estuary at Kruibeke station (Belgium) in summer and winter.
4.2.4 Application 2: phytoplankton bloom

During typically observed phytoplankton spring blooms, we cannot assume dynamic equilibrium of $\tilde{B}(t, x)$ in Eq. (4.1). In the following, we study the full solution of Eq. (4.1). This spatio-temporal differential equation cannot be solved analytically without further assumptions. Therefore, we again focus on a specific location $x = x^*$ and assume that $\tilde{B}(t, x)$ evolves linearly along the estuarine axis near $x = x^*$. Moreover, we suppose that its shape in the longitudinal direction is time-independent on the time scale of our application (≈days):

$$
\tilde{B}(x, t) = B(t)f(x),
$$

in which $f(x)$ is a linear function of $x$. Furthermore, we assume a phase of $-\pi/2$ between the longitudinal velocity $u(t)$ and the water elevation (cf. $d$) following van Rijn (2010). For a detailed derivation, the reader is referred to Appendix 4.C. The solution of $B(t)$ yields

$$
B(t) = \tilde{A}e^{-m^*\cdot t + (1-\rho_B)\cdot \Pi f \Lambda(t)dt} \times e^{\left[-\frac{\gamma}{\omega_{M2}} \cdot \cos(\omega_{M2} t + \Phi_{M2} + \theta)\right]},
$$

with

$$
\tilde{A} = B_0 e^{- (1 - \rho_B) \cdot \Pi f \Lambda(t)dt|_{t=0} - \frac{\gamma}{\omega_{M2}} \cdot \cos(\Phi_{M2} + \theta)} \quad \text{and} \quad (4.23)
$$

where $\omega_{M2}$ is the angular frequency corresponding to the semidiurnal lunar M2 tide and $\Phi_{M2}$ and $\theta$ are the phases corresponding to the idealized time dependence of the water depth $d$ [see below, Eq. (4.28)]. The coefficients $m^*$ and $\gamma$ are defined in Appendix 4.C. The integrated loss rate $m^*$ incorporates losses due to grazing, mortality, and residual
water currents. $\gamma$ is a measure of the amplitude of the sinusoidal behavior due to advective transport of phytoplankton biomass [see Eq. (4.41)]. Again, we apply the moment approximation to the solution of $B(t)$ in Eq. (4.22), which yields

$$\int Z\Lambda(t) dt \approx m_{\text{fun}}_0 \cdot t \times \left\{ 1 + \frac{m_{\text{fun}}.(\beta, \lambda)}{t} \cdot \int [\kappa(t) + \delta(t)] dt + \frac{m_{\text{fun}}.\kappa(\beta, \lambda)}{t} \cdot \int \epsilon(t) dt + \frac{m_{\text{fun}}.\kappa^2(\beta, \lambda)}{t} \cdot \int [\kappa^2(t) + \delta^2(t)] dt + \frac{m_{\text{fun}}.\kappa\epsilon(\beta, \lambda)}{t} \cdot \int \epsilon(t) \cdot \kappa(t) dt + \frac{m_{\text{fun}}.\kappa\delta(\beta, \lambda)}{t} \cdot \int [\delta(t) \cdot \kappa(t)] dt + \frac{m_{\text{fun}}.\delta(\beta, \lambda)}{t} \cdot \int [\delta(t) \cdot \delta(t)] dt \right\}.$$  (4.24)

To calculate the integrals in Eq. (4.24), we use approximate expressions for $\epsilon(t)$, $\kappa(t)$, and $\delta(t)$. We assume that solar irradiance $E_0$ is given by a truncated sinusoid following Cox et al. (2015):

$$E_0(t) = \max(0, a + b \cos(\omega_{S1} t)), \quad (4.25)$$

in which $a$ and $b$ are further unspecified parameters and $\omega_{S1}$ is the diurnal angular frequency. Consequently, we neglect aperiodic temporal variability in $E_0$ over more rapid time scales (seconds to minutes, due to, for example, cloud shading) and only include the dominant day-night pattern. We simplify Eq. (4.25) by only including the first three temporal terms of the Fourier series:

$$E_0(t) = \langle E_0 \rangle + \sum_{n=1}^{3} \hat{E}_n \cos(n\omega_{S1} t). \quad (4.26)$$

Similarly, we only consider the dominant and periodic $\omega_{M2}$ and $\omega_{M4} = 2\omega_{M2}$ temporal fluctuations in $k_d$ and $d$ and truncate frequencies larger than $M4$ (i.e., $M6$, $M8$, etc. are neglected) following Chernetsky et al. (2010) Dijkstra et al. (2017) and Horemans et al. (2020):

$$k'_d(t) = \hat{K}_2 \cos(\omega_{M2} t + \Phi_{M2}) + \hat{K}_4 \cos(\omega_{M4} t + \Phi_{M4}), \quad (4.27)$$

$$d'(t) = \hat{D}_2 \cos(\omega_{M2} t + \Phi_{M2} + \theta), \quad (4.28)$$

in which $\hat{K}_2$, $\hat{K}_4$, and $\hat{D}_2$ and $\Phi_{M2}$, $\Phi_{M4}$, and $\theta$ are (calibrated) amplitudes and phases, respectively. To estimate these amplitudes and phases, we fitted Eqs. (4.26)-(4.28) to
Figure 4.4: (a) Turbidity, (b) water depth, and (c) Chlorophyll-a (Chl-a) concentration measured in the Scheldt estuary at Schellebelle (Belgium) in spring during a phytoplankton bloom. (d) The solar irradiance $E_0$ is measured at Stabroek (Belgium), located approximately 50 km from Schellebelle.

observed time series (see Appendix 4.D). We use time series measured in the Scheldt estuary in the freshwater zone at Schellebelle ($\sim$140 km from the mouth) during a spring bloom (29-03-2017 until 19-04-2017, Fig. 4.4). The same methodology and sensors as in application 1 were used. Chl-a concentration was measured using a fluorescence sensor [YSI6025, wavelength 435-470 nm (YSI, 2017b)] with a sampling interval of 5 minutes. Parameters $m^*$, $\gamma$, and $\rho_B$ are obtained by fitting Eq. (4.22) to the observed Chl-a time series. We used a fixed ratio of 25 mg C (mg Chl-a)$^{-1}$ to transform Chl-a concentration to phytoplankton biomass concentration (Maris and Meire, 2016), assuming balanced-growth conditions. Table 4.3 summarizes the resulting estimated parameters. The order of magnitude of $\gamma$ and $m^*$ corresponds to the estimated order in Appendix 4.C, that is, $O(10^{-1})$ h$^{-1}$ and $O(10^{-2})$ h$^{-1}$, respectively. The order of the fitted value for $\rho_B$ agrees with values found by Desmit et al. (2005).
Table 4.3: Parameter values estimated by fitting the presumed idealized time dependencies for $E_0(t)$, $k_d(t)$, and $d(t)$ [Eqs. (4.26)-(4.28)] and analytical solution of $B(t)$ to the corresponding data sets of $k_d(t)$, $E_0(t)$, $d(t)$, and $B(t)$ measured in the Scheldt estuary at Schellebelle (Figs. 4.4a-c).

<table>
<thead>
<tr>
<th></th>
<th>$J$ m$^{-2}$ s$^{-1}$</th>
<th>$m^{-1}$</th>
<th>$h^{-1}$</th>
<th>$\omega$</th>
<th>$\Phi$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_1$</td>
<td>142</td>
<td>$\langle k_d \rangle$</td>
<td>4.56</td>
<td>$\langle d \rangle$</td>
<td>4.5 m</td>
</tr>
<tr>
<td>$E_2$</td>
<td>75.3</td>
<td>$K_2$</td>
<td>0.82</td>
<td>$D_2$</td>
<td>1.36 m</td>
</tr>
<tr>
<td>$E_3$</td>
<td>3.44</td>
<td>$K_4$</td>
<td>0.43</td>
<td>$\rho_B$</td>
<td>0.837</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

To summarize, the combination of the moment approximation in Eq. (4.24) and idealized time dependence in $E_0$, $k_d$, and $d$ [Eqs. (4.26)-(4.28)] allows us to analytically solve the time integral of $\Lambda(t)$ and corresponding analytical solution of $B(t)$ presented in Eq. (4.22). The main advantage of this approach is that it enables us to directly analyze the propagation of the dominant harmonics in forcings $E_0$, $k_d$, $d$, and advective water flow into the dynamics of $B(t)$. The analytical solution of $\int \Lambda(t) dt$ is listed in Appendix 4.E.

4.3 Results

We first show the generic results of the moment approximation by presenting the mfun functions [see Eq. (4.10)], which translate temporal variability in $k_d$, $E_0$, and $d$ to temporal variability in $\Lambda$, GPP, and phytoplankton biomass dynamics. Next, using these mfun functions, we present the results of our two applications in the Scheldt estuary, i.e., time-averaged GPP in dynamic equilibrium (application 1) and a phytoplankton bloom in dynamic non-equilibrium (application 2).

4.3.1 Moment approximation: the mfun functions

4.3.1.1 mfun functions of the zeroth- and first-order terms

The zeroth-order mfun$^0$ function is, by definition, equal to the time-averaged DAP factor $\langle \Lambda \rangle$ in the absence of temporal variability in $E_0$, $k_d$, and $d$. Figure 4.5 shows mfun$^0$ for various $\beta$ and $\lambda$. mfun$^0$ ranges from 0 (maximal light limitation; no GPP) to 1 (no light limitation; maximal GPP) and has a simple structure; it increases with increasing $\beta$ (cf. increasing solar irradiance/PP efficiency) and decreases with increasing $\lambda$ (cf. increasing turbidity/depth). The mfun$^\kappa$ and mfun$^\epsilon$ functions are plotted in Figs. 4.6a and 4.6b, respectively. These first-order functions represent the system-dependent multiplication factors that are required to compute the amplitude of different harmonics in light climate (cf. $\Lambda$), GPP, and corresponding biomass dynamics from the respective amplitudes in $E_0$, $k_d$, and $d$. mfun$^\kappa$ is negative for every value of $\lambda$ and $\beta$ (Fig. 4.6b), simply reflecting
Figure 4.5: The mfun⁰ function for various β and λ. mfun⁰ corresponds to the DAP function Λ in the absence of fluctuations. The averaged parameter set corresponding to the Scheldt estuary cases described in this paper is also illustrated (i.e., λ ≈ 60, β ≈ 3).

Figure 4.6: The first-order mfun functions (a) mfunₖ and (b) mfunₐ for various β and λ. The averaged parameter set corresponding to the Scheldt estuary cases described in this paper is also illustrated (i.e., λ ≈ 60, β ≈ 3).

4.3.1.2 mfun functions of the second-order terms

The second-order mfun functions represent the factors by which the variances and covariances have to be multiplied to calculate the magnitude of a systematic upward or downward shift in its time-average $\langle \Lambda(t) \rangle$ [Eq. (4.17)]. They have a more interesting structure (Fig. 4.7). Figure 4.7 shows that mfun₂ is always negative. In contrast, the
Figure 4.7: The second-order mfun functions (a) mfun_\kappa^2, (b) mfun_\epsilon^2, (c) mfun_\kappa\epsilon, and (d) mfun_\kappa\delta for various \beta and \lambda. The averaged parameter set corresponding to the Scheldt estuary cases described in this paper is also illustrated (i.e., \lambda \approx 60, \beta \approx 3). The contour where the mfun function equals zero and changes sign is depicted by a dashed curved line.
sign of the other second-order mfun functions depends on the specific values of $\lambda$ and $\beta$ (Figs. 4.7a, 4.7c, and 4.7d). This has a simple interpretation for the time-averaged $\langle \text{GPP} \rangle$ [Eqs. (4.16) and (4.17)]. Indeed, temporal variability in $E_0$ always results in a decrease in $\langle \text{GPP} \rangle$, while variability in light attenuation and depth can lead to both a decrease or an increase in $\langle \text{GPP} \rangle$, depending on the characteristics of the system under study. Here, we assumed that the covariance terms due to $E_0$ ($\sim \text{mfun}_\epsilon\kappa$) are negligible compared to the variance term ($\sim \text{mfun}_\epsilon^2$) in Eq. (4.17). Similarly, for phytoplankton in the exponential growth phase (cf. bloom), second-order temporal variability in $E_0$ always result in a decrease of the exponential growth, while variability in light attenuation and depth can lead to both a decrease or an increase of the exponential growth. It must be noted that tidal systems for which $\text{mfun}_\kappa^2$ is negative ($\lambda \lesssim 5$, Figs. 4.7a, 4.7c, and 4.7d), correspond to shallow and clear systems which can be found in, for example, the Dutch Wadden Sea or the Oosterschelde (Rijkswaterstaat, 2020). These systems often experience nutrient limitation, and thus the results presented here are valid only during periods without nutrient limitation.

To summarize, the results of the mfun functions show that the propagation of temporal variability in the parameters affecting light climate (i.e., $E_0$, $k_d$, and $d$) to temporal variability in GPP and phytoplankton biomass dynamics may show opposing trends for the different parameters. Moreover, this propagation depends on the time-independent characteristics of the system (i.e., $\lambda$ and $\beta$) and may thus differ for different systems and seasons. In the following sections, we use these results to study our two applications in the Scheldt estuary, covering different seasons.

### 4.3.2 Application 1: time-averaged GPP

In this section, we use the mfun function results from the moment approximation presented in the previous section to study the impact of temporal variability in light climate (cf. $E_0$, $k_d$, and $d$) on time-averaged $\langle \text{GPP} \rangle$ in dynamic equilibrium in the brackish part of the Scheldt estuary. Because we expect different results for different seasons, we analyze both a summer and winter case.

At our relatively deep and turbid case study location, $\text{mfun}_{\kappa^2}$ is always positive (Fig. 4.7b) and thus temporal variability in $k_d$ and $d$ lead to an increase in $\langle \text{GPP} \rangle$. In contrast, temporal variability in $E_0$ decreases $\langle \text{GPP} \rangle$ (Fig. 4.7b). Figure 4.8 shows the results of $\langle \text{GPP} \rangle$ when including temporal variability in all parameters (all var), normalized with $\langle \text{GPP} \rangle$ with time-averaged parameters (i.e., $\langle E_0 \rangle$, $\langle k_d \rangle$, and $\langle d \rangle$) for various integration times $T$ (8.5 h-21 days) using the moment approximation (left) and a numerical integrator (right), applied to a summer (top) and winter (bottom) case. When the presented value is $\approx 1$, the impact of temporal variability is negligible. To compute the separate impact of temporal variability in $E_0$, $k_d$, and $d$ on $\langle \text{GPP} \rangle$, we also show the results exclusively including temporal variability in the individual parameters $E_0$ ($E_0$ var), $k_d$ ($k_d$ var), and $d$ ($d$ var).
Figure 4.8: Time-averaged gross primary production \( \langle \text{GPP} \rangle \) in dynamic equilibrium, relative to \( \langle \text{GPP} \rangle \) using time-averaged parameters \( \langle E_0 \rangle, \langle k_d \rangle \), and \( \langle d \rangle \) for various integration times (8.5 h-21 days) using the moment approximation (left) and a numerical integrator (right) in (a)-(b) summer and (c)-(d) winter. Our framework allows us to compute the impact of temporal variability in an individual parameter on \( \langle \text{GPP} \rangle \) (cf. d var, \( k_d \) var, and \( E_0 \) var) or when temporal variability in all parameters is included (cf. all var). When the presented value is \( \approx 1 \), the impact of temporal variability is negligible.
4.3.2.1 Summer case

In summer, the relative impact of temporal variability on $\langle \text{GPP} \rangle$ is < 10 % for $k_d$ and $d$ and 10-30 % for $E_0$. Overall, Fig. 4.8 shows good correspondence between the moment approximation and the numerical integrator, even for integration times $T < 24$ h, that is, the period corresponding to the angular frequency of temporal variability in $E_0$. In this specific summer case study, excluding temporal fluctuations in $E_0$, $k_d$, and $d$ results in an error of approximately 20 %. Temporal variability in $E_0$ ($E_0 \text{ var}$) has the largest impact on $\langle \text{GPP} \rangle$.

4.3.2.2 Winter case

Also in the winter case, the moment approximation is in good correspondence with the numerically integrated impact of temporal variability in $E_0$, $k_d$, and $d$ on $\langle \text{GPP} \rangle$, respectively approximately 20-50, 5-20, and < 10 %. However, we have a slight decrease in accuracy, which is due to the relatively large temporal fluctuations in solar irradiance compared to the averaged solar irradiance in winter. This results in a less accurate approximation by sinusoids of the solar temporal fluctuations [Eq. (4.26)]. The main difference with the summer case is an increase in the relative importance of fluctuations in $k_d$. This difference follows from the relative magnitude of temporal variability in $k_d$ (cf. $\text{Var}[k_d(t)]$), which is in winter an order of magnitude larger compared to summer (see Table 4.1).

To summarize, our summer and winter cases illustrate that the impact of fluctuations in light climate on $\langle \text{GPP} \rangle$ in dynamic equilibrium may be important, season-dependent, and that a careful analysis based on the observed time series of $E_0$, $k_d$, and $d$ is required.

4.3.3 Application 2: phytoplankton bloom

In this section, we apply our idealized model to study $B(t)$ in the exponential growth phase. Using our analytical solution of $B(t)$ and corresponding second-order approximation and results of the mfun functions, our approach allows for an extensive analysis by separating the individual impact of temporal variability in $E_0$, $k_d$, and $d$ and their interactions on (exponential) phytoplankton biomass growth. We analyze the time series of the onset of a phytoplankton bloom in spring in the freshwater region of the Scheldt estuary.

Figure 4.9 shows both the observed $B(t)$ (cf. Chl-a) and model result using our idealized model [Eq. (4.22)] with fitted and idealized expressions of temporal fluctuations in $E_0$, $k_d$, and $d$ [Eqs. (4.26)-(4.28)]. Our model captures the exponential growth of $B(t)$: both the observed (data) and modeled (all var) $B(t)$ doubles after approximately two weeks (from $\sim 750$ $\mu$g L$^{-1}$ to $\sim 1500$ $\mu$g L$^{-1}$). To study the impact of individual temporal variability on $B(t)$, Fig. 4.9 shows the modeled $B(t)$ in which we only include
Figure 4.9: Observed phytoplankton biomass $B(t)$ (data) and model result by including all temporal fluctuations (all var) using our second-order approximation. Our approach allows us to easily identify and exclude individual forcings. The dashed green line (no fluct.) shows the model output for $B(t)$ which is only forced by time-invariant $\langle E_0 \rangle$, $\langle k_d \rangle$, and $\langle d \rangle$. We also show $B(t)$ when we exclusively include temporal fluctuations in $d$ (d var), $k_d$ (k_d var), $E_0$ (E_0 var), and temporal fluctuations due to advective transport of $B(t)$ (adv. var). The orange dashed graph (low freq.) shows $B(t)$ when, on top of the exponential growth, low-frequency fluctuations due to interactions between temporal variability in $E_0$ and $k_d$ and $d$ are included, which results in exponential coefficients $\sim \sin[(\omega M_2 - 2\omega S_1)t]$.

specific temporal variability. When $B(t)$ is forced by time-invariant $\langle E_0 \rangle$, $\langle k_d \rangle$, and $\langle d \rangle$ (Fig. 4.9 no fluct.), it grows much faster than observed. After two weeks, $B(t)$ when no temporal fluctuations are included is approximately a factor of 14 larger than observed. This illustrates the importance of the $\sim t$ terms generated by temporal variability in $E_0$, $k_d$, and $d$ (see Appendix 4.E for definitions of these terms). In particular, this overestimation is due to the absence of temporal variability in $E_0$, which again decreases phytoplankton growth. If we solely include temporal variability in $E_0$, the modeled $B(t)$ is lower than observed (Fig. 4.9 E_0 var). Including temporal fluctuations in $E_0$, but excluding fluctuations in $k_d$ and $d$, leads to underestimating $B(t)$ with a factor of 15 after two weeks. In contrast, exclusively including temporal fluctuations in $d$ (Fig. 4.9 d var) and $k_d$ (Fig. 4.9 k_d var) results in an overestimation of $B(t)$ with, respectively, a factor of 75 and 34 after two weeks. These results are consistent with application 1, but show that the accumulated impact is much larger than expected from the calculations assuming dynamic equilibrium. Finally, temporal fluctuations due to advective transport of $B(t)$ do not result in additional exponential growth (Fig. 4.9 adv. var) when compared to $B(t)$ forced by time-invariant parameters $\langle E_0 \rangle$, $\langle k_d \rangle$, and $\langle d \rangle$ (Fig. 4.9 no fluct.).

Temporal variability significantly reduces the exponential increase in our case study, which has important consequences, particularly at the onset of phytoplankton blooms. Indeed, during the onset of a phytoplankton bloom, a slight change in phytoplankton growth rate can change the balance with loss terms (e.g., respiration, grazing, and flushing), and result in net growth. Thus, other characteristics being equal, a system with fluctuating parameters has an earlier/later onset of the spring phytoplankton bloom than
a system without fluctuations. In our case study, temporal variability in $E_0$, $k_d$, and $d$ delay the onset of the bloom by approximately two weeks (see Appendix 4.F for a detailed derivation). Therefore, knowing the magnitude of fluctuations in the forcings $E_0$, $k_d$, and $d$ is at least as important as having a good and realistic estimate of respiration, grazing, and flushing, which in our analysis are included in $m^*$.

Temporal fluctuations in $E_0$, $k_d$, and $d$ do not only result in additional $\sim t$ terms in the exponential in Eq. (4.22), but also result in sinusoidal terms. Of particular interest are the nontrivial interaction terms leading to additional low-frequency variability in $B(t)$ (Fig. 4.9 low freq.). These low-frequency fluctuations have a frequency similar to a spring-neap cycle of the horizontal tide (see Appendix 4.E for a definition of these low-frequency fluctuations $\sim \sin[(\omega_{M2} - 2\omega_{S1})t]$). This makes it difficult to separate the effects of spring-neap cycles of the horizontal tide from the interaction terms. The model seems to overestimate the impact of interaction terms on $B(t)$. A potential explanation is that the idealized expressions for temporal variability in $k_d$ and $d$ do not include the spring-neap cycle. These results highlight the need for caution when interpreting low-frequency patterns in observed Chl-a time series. As shown, these may both result from the spring-neap cycle and interaction terms in the light limitation factor.

To summarize, our analysis of the phytoplankton bloom in spring illustrates that the impact of temporal variability in light climate and advective water flow on $B(t)$ in the exponential growth phase may have crucial consequences: it may result in a 14 times lower $B(t)$ after two weeks, delay the onset of the bloom by two weeks, and result in low-frequency temporal variability with similar periodicity as the spring-neap cycle. As in application 1, temporal variability in $k_d$ and $d$ increases, whereas temporal variability in $E_0$ decreases phytoplankton growth.

### 4.4 Discussion

In this section, we first compare our case study results of the impact of temporal variability on time-averaged GPP in dynamic equilibrium (application 1) and unbound $B(t)$ (application 2) to the literature. Next, we discuss the implications of our results for both modelers and experimental scientists. Finally, we discuss the limitations of our approach, focusing on non-local processes and temporal variability in phytoplankton characteristics affecting $B(t)$ dynamics.

#### 4.4.1 The impact of temporal variability on phytoplankton growth

We constructed an idealized model to study the impact of temporal variability in light climate and advective water flow on time-averaged GPP and $B(t)$ dynamics. Our approach allows for an extensive analysis by separating the individual impact of temporal variability in $E_0$, $k_d$, and $d$ and their interactions on time-averaged GPP and (exponential) growth of $B(t)$.
In our first application, we found that temporal fluctuations in light climate impact time-averaged GPP: temporal variability in \( k_d \) and \( d \) increase, whereas temporal variability in \( E_0 \) decrease time-averaged GPP and has a dominant impact (a decrease up to 50\%, depending on the season). The dominant impact of temporal variability in \( E_0 \) complies with the results of Cox et al. (2015). They developed a novel technique to estimate GPP using oxygen time series. A core assumption in this method states that GPP follows a truncated sinusoidal pattern over time with a 24 h period. Consequently, the method assumes that the main contribution of temporal variability in the forcings is due to temporal variability in \( E_0 \) (cf. the 24 h period).

In our second application, we studied the impact of temporal variability in light climate and advective water flow on \( B(t) \) dynamics in the exponential growth phase (i.e., phytoplankton bloom). Our results comply with the results of Desmit et al. (2005) who studied the impact of temporal variability in \( E_0, k_d, \) and \( d \) within a 40-day time frame for a specific case in the Scheldt estuary. Firstly, the exponential increase in \( B(t) \) is very sensitive to temporal variability in \( E_0, k_d, \) and \( d \) (Fig. 4.9). Secondly, temporal variability in \( k_d \) and \( d \) increases the exponential growth, whereas temporal variability in \( E_0 \) decreases the exponential growth (Fig. 4.9). Moreover, the magnitude of the exponential increase is determined by the ratio of time-averaged depth \( \langle d \rangle \) and corresponding temporal fluctuations \( \langle d'(t) \rangle \), which agrees with our definition of \( \delta(t) \) in Eq. (4.11). Last, a clear link exists between our implementation of advective transport of \( B(t) \) and the one presented in Desmit et al. (2005). To include advective transport of \( B(t) \), Desmit et al. (2005) postulated an additional term in Eq. (4.1) that is proportional to the derivative of \( d(t) \), which agrees with our result in Eq. (4.50) [from Eq. (4.28) follows that \( \partial_t d(t) \sim \sin(t) \)]. However, our approach generalizes the work of Desmit et al. (2005) who only studied a specific case in the Scheldt estuary. In contrast to Desmit et al. (2005), our approach is more generic and can be applied to other estuarine well-mixed, light-limited systems. Moreover, our framework allows us to easily compute the impact of temporal fluctuations in the individual parameters and their interactions on phytoplankton growth because we derived an explicit analytical solution for phytoplankton growth and applied a second-order moment approximation.

We thus conclude that our results of the impact of temporal variability on time-averaged GPP and exponential phytoplankton growth comply with results found in the literature and that they allow for a more generic analysis of well-mixed, light-limited estuaries.

### 4.4.2 Implications of our findings for modelers and experimental scientists

Our generic approach may be used to study present-day challenges by both modelers and experimental scientists. Modelers may use our approach to easily assess the error by neglecting temporal fluctuations in parameters affecting \( B(t) \) dynamics. As mentioned in the introduction, various models have been used to model \( B(t) \) dynamics. Although complex tide-explicit models exist (Arndt et al., 2011), various models often keep certain
parameters constant to reduce complexity and allow for fast analytical solution procedures (Dijkstra et al., 2017). Examples are tidally averaged and daily-averaged numerical models to compute estuarine PP (Brinkman, 1993; Soetaert et al., 1994). The light extinction coefficient \( k_d \), which is an important parameter in light-limited estuarine systems, is one of the main parameters which contributes to the complexity of PP dynamics because it is affected by suspended particulate matter (SPM) dynamics. Modeling SPM dynamics, even the corresponding large-scale estuarine turbidity maxima, is difficult because they result from a complex concurrence of ETM formation mechanisms [see Burchard et al. (2018) for a recent review]. However, our analysis framework provides a tool to assess the error made by neglecting such complex temporal variability in parameters affecting PP dynamics. As such, it may help to assess whether a given model with given accuracy and performance is sufficient to realistically simulate and study phytoplankton dynamics. This is particularly important when strong spatial and seasonal gradients in temporal variability in \( E_0, k_d \) (cf. SPM), and \( d \) exist. After all, this temporal variability can falsely be attributed (cf. calibrated) to spatial and seasonal variations in intrinsic features of phytoplankton dynamics such as \( P_{\text{max}}, \alpha, \) and \( m \). Finally, to use our framework for a model assessment, only basic knowledge on time-average parameter values and corresponding S1, M2, and M4 amplitudes is required.

Our results are also particularly useful for experimental scientists interested in data-based estimation of GPP. They may partly answer the long-standing question of why bottle incubations underestimate in situ GPP. Often, bottle incubations are used to estimate GPP in estuaries and coasts. This means that the parameters of the Platt equation \( \alpha \) and \( P_{\text{max}} \) in Eq. (4.2) are determined by incubating a sample at different light intensities (Vegter and De Visscher, 1984; Kromkamp and Peene, 1995). Consequently, the Platt equation needs to be numerically integrated over depth and time. This requires time series of \( E_0, k_d, \) and \( d \). Whereas time series of incident irradiance are often available, \( k_d \) is often estimated as an average on a short time interval during sampling, and linearly interpolated between consecutive sampling (Kromkamp and Peene, 2005). Our results show that in systems with large temporal variability in \( k_d \), this approach leads to a systematic underestimation of time-averaged PP. Indeed, temporal variability in \( k_d \) has a positive impact on GPP (Fig. 4.8). It has often been found that bottle incubations underestimate PP when compared to in situ methods. The reason why this occurs is still poorly understood (Westberry et al., 2012). In addition to so-called ‘bottle effects’ (Swaney et al., 1999) and physiological adaptation effects (Halsey et al., 2010), our results show that large variability in light attenuation can additionally result in an underestimation when this variability is not taken into account.

4.4.3 Non-local processes and temporal variability in phytoplankton characteristics affecting the phytoplankton dynamics

Although our approach is generically applicable to well-mixed, light-limited estuaries, an extrapolation of our findings to other estuarine cases should be taken with care.

As a first step, we approximated the non-local processes caused by advective and
diffusive transport of $B(t)$ by a term that is proportional to the derivative of the water depth. As mentioned above, this complies with the postulation presented in Desmit et al. (2005). By applying this idealization, we partly neglected the impact of temporal variability caused by complex interactions between $B(t)$ and the water flow. We assumed that $B(t)$ growth is mainly caused by local processes, which complies with the model study carried out by Dijkstra et al. (2019) who focused on the phytoplankton dynamics in the Delaware River Estuary. However, we are aware that this may not be valid in the Scheldt estuary. To correct for complex, non-local interactions between the water flow and $B(t)$, our model can still be used by coupling the model to a hydrodynamical model. By doing so, the impact of individual temporal variability in light climate on $B(t)$ growth can be investigated, including these complex non-local $B(t)$ and water flow interactions, as done by Dijkstra et al. (2019).

Additionally, we focused on temporal variability in parameters affecting the light climate $E_0$, $k_d$, and $d$ on an hourly-daily time scale and kept most other parameters fixed. For example, we assumed balanced growth conditions and the absence of significant temporal variations in the phytoplankton characteristics (e.g., $P_{\text{max}}$, $\alpha$) on this hourly-daily time scale. In further research, the same approach as presented in this study can be applied assuming temporal variability in the latter parameters. This may cause additional (correlation) terms in the second-order approximation of $\Lambda$ [Eq. (4.10)] and thus alter time-averaged GPP and $B(t)$ dynamics.

4.5 Conclusions

We constructed an idealized model to analyze the impact of temporal variability in solar irradiance at the water surface $E_0$, exponential light extinction coefficient $k_d$, and water depth $d$ on phytoplankton biomass $B(t)$ and corresponding time-averaged gross primary production $\langle \text{GPP} \rangle$ in light-limited, vertically well-mixed systems. Apart from providing a full theoretical analysis, our approach allows us to quantify the impact of the temporal variability without a numerical model, to separate the impact of different sources of the temporal variability and their covariance, and it does so for a general light-limited, well-mixed tidal system.

To present the applicability of our idealized model, we applied the model to two cases in the Scheldt estuary. Our results showed that temporal variability in $E_0$, $k_d$, and $d$ can have an important impact on $\langle \text{GPP} \rangle$ in dynamic equilibrium and corresponding $B(t)$ dynamics. In the first application, we showed that $\langle \text{GPP} \rangle$ in dynamic equilibrium is mainly impacted by temporal variability in $E_0$ and results in a 30 % decrease. In contrast, temporal variability in $k_d$ and $d$ increased $\langle \text{GPP} \rangle$. Furthermore, the results showed a seasonality; in winter, temporal variability in $k_d$ had a larger impact on $\langle \text{GPP} \rangle$ than in summer, due to the larger tidal amplitude in $k_d$ in winter. Finally, our idealized model correctly predicted the order of magnitude of the impact of temporal variability of the individual parameters.
In the second application, we showed that during a phytoplankton bloom, temporal variability significantly contributes to the exponential $B(t)$ growth and generates additional low-frequency fluctuations similar to the spring-neap cycle. Moreover, we showed that temporal variability may delay the onset of the phytoplankton bloom by two weeks and may decrease $B(t)$ by a factor of 14 after two weeks. Again, our approach allowed us to apply an extensive analysis in which we showed that these low-frequency fluctuations in $B(t)$ are due to covariance of temporal variability in $E_0$ and $k_d$, and $d$.

List of symbols

Latin

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a$</td>
<td>Parameter defining the truncated sinusoidal behavior of $E_0$</td>
</tr>
<tr>
<td>$\dot{B}$</td>
<td>Depth-averaged phytoplankton biomass concentration</td>
</tr>
<tr>
<td>$B$</td>
<td>Local depth-averaged phytoplankton biomass concentration</td>
</tr>
<tr>
<td>$B_0$</td>
<td>Initial depth-averaged phytoplankton biomass concentration (at $t = 0$)</td>
</tr>
<tr>
<td>$b$</td>
<td>Parameter defining the truncated sinusoidal behavior of $E_0$</td>
</tr>
<tr>
<td>C:Chl-a</td>
<td>Carbon to Chlorophyll-a ratio in a phytoplankton cell</td>
</tr>
<tr>
<td>$\tilde{\text{Cov}}$</td>
<td>Statistical covariance operator in which the parameter of interest is divided by its time average</td>
</tr>
<tr>
<td>$\hat{D}_2$</td>
<td>Amplitude corresponding to the M2 tidal component of $d'$</td>
</tr>
<tr>
<td>$d$</td>
<td>Water depth</td>
</tr>
<tr>
<td>$d'$</td>
<td>Temporal variability of $d$ relative to its time average</td>
</tr>
<tr>
<td>$\langle d \rangle$</td>
<td>Time average of $d$</td>
</tr>
<tr>
<td>$E_0$</td>
<td>Solar irradiance at the water surface</td>
</tr>
<tr>
<td>$E_0'$</td>
<td>Temporal variability of $E_0$ relative to its time average</td>
</tr>
<tr>
<td>$\langle E_0 \rangle$</td>
<td>Time average of $E_0$</td>
</tr>
<tr>
<td>$\hat{E}_1$</td>
<td>Amplitude corresponding to the $\omega_{S1}$ component of $E_0'$</td>
</tr>
<tr>
<td>$\hat{E}_2$</td>
<td>Amplitude corresponding to the $2\omega_{S1}$ component of $E_0'$</td>
</tr>
<tr>
<td>$\hat{E}_3$</td>
<td>Amplitude corresponding to the $3\omega_{S1}$ component of $E_0'$</td>
</tr>
<tr>
<td>$\hat{E}_n$</td>
<td>Amplitude corresponding to the $n\omega_{S1}$ component of $E_0'$</td>
</tr>
<tr>
<td>$f(x)$</td>
<td>Linear function that captures the spatial dependence of $\tilde{B}$</td>
</tr>
<tr>
<td>$\langle \text{GPP} \rangle$</td>
<td>Time-averaged gross primary production</td>
</tr>
<tr>
<td>$I$</td>
<td>Light intensity in the water column</td>
</tr>
<tr>
<td>$\hat{K}_2$</td>
<td>Amplitude corresponding to the M2 tidal component of $k_d'$</td>
</tr>
<tr>
<td>$\hat{K}_4$</td>
<td>Amplitude corresponding to the M4 tidal component of $k_d'$</td>
</tr>
<tr>
<td>$K_h$</td>
<td>Horizontal diffusivity coefficient</td>
</tr>
<tr>
<td>$k_d$</td>
<td>Exponential light extinction coefficient</td>
</tr>
<tr>
<td>$k_d'$</td>
<td>Temporal variability of $k_d$ relative to its time average</td>
</tr>
<tr>
<td>$\langle k_d \rangle$</td>
<td>Time average of $k_d$</td>
</tr>
<tr>
<td>$m$</td>
<td>Mortality rate of phytoplankton</td>
</tr>
<tr>
<td>$m^*$</td>
<td>Integrated mortality rate, including the effects of advective transport of phytoplankton</td>
</tr>
<tr>
<td>$\text{mfun}_0$</td>
<td>$\Lambda$ without temporal variability in its parameters</td>
</tr>
</tbody>
</table>
mfun_{\delta} \quad \text{First-order mfun function corresponding to } \delta
mfun_{\delta^2} \quad \text{Second-order mfun function corresponding to } \delta
mfun_{\delta\epsilon} \quad \text{Second-order mfun function corresponding to the correlation of } \delta \text{ and } \epsilon
mfun_{\epsilon} \quad \text{First-order mfun function corresponding to } \epsilon
mfun_{\epsilon^2} \quad \text{Second-order mfun function corresponding to } \epsilon
mfun_{\epsilon\kappa} \quad \text{Second-order mfun function corresponding to the correlation of } \epsilon \text{ and } \kappa
mfun_{\kappa} \quad \text{First-order mfun function corresponding to } \kappa
mfun_{\kappa^2} \quad \text{Second-order mfun function corresponding to } \kappa
mfun_{\kappa\delta} \quad \text{Second-order mfun function corresponding to the correlation of } \kappa \text{ and } \delta
N \quad \text{Total number of field observations}
n \quad \text{Natural number}
\mathcal{O} \quad \text{Order of}
\text{P}_{\text{max}} \quad \text{Maximum photosynthetic rate}
t \quad \text{Time}
u \quad \text{Local depth-averaged water flow velocity in the longitudinal dimension}
\tilde{u} \quad \text{Water flow velocity in the longitudinal dimension}
\text{Var} \quad \text{Statistical variance operator in which the parameter of interest is divided by its time average}
x \quad \text{Spatial coordinate in the longitudinal dimension}
x^* \quad \text{Location of interest in the longitudinal dimension}
z \quad \text{Spatial coordinate in the vertical dimension}
\tilde{z} \quad \text{Nondimensional parameter (i.e., } k_d z) \text{ defining } \tilde{\Lambda}

\textbf{Greek}

\alpha \quad \text{Photosynthetic growth efficiency}
\beta \quad \text{Time-averaged } \tilde{\beta}
\gamma \quad \text{Amplitude corresponding to the advective transport of phytoplankton}
\tilde{\beta} \quad \text{Nondimensional parameter (i.e., } \alpha E_0 / P_{\text{max}}) \text{ defining } \tilde{\Lambda}
\delta \quad \text{Temporal variability of } d \text{ divided by its time average (i.e., } d' / \langle d \rangle)
\partial_t \quad \text{Partial derivative to } t
\epsilon \quad \text{Temporal variability of } E_0 \text{ divided by its time average (i.e., } E_0' / \langle d \rangle)
\theta \quad \text{Phase corresponding to the M2 tidal component of } d'
\kappa \quad \text{Temporal variability of } k_d \text{ divided by its time average (i.e., } k_d' / \langle d \rangle)
\Lambda \quad \text{Depth-averaged Platt light limitation function}
\tilde{\Lambda} \quad \text{Platt light limitation function}
\lambda \quad \tilde{z} \text{ using time-averaged parameters (i.e., } \langle k_d \rangle / \langle d \rangle)
\rho_B \quad \text{Respiration and excretion coefficient of phytoplankton}
\Phi_{M2} \quad \text{Phase corresponding to the M2 tidal component of } k_d'
\Phi_{M4} \quad \text{Phase corresponding to the M4 tidal component of } k_d'
\omega_{M2} \quad \text{Angular frequency corresponding to the semidiurnal lunar M2 tide}
\omega_{M4} \quad \text{Angular frequency corresponding to the M4 tide}
\omega_{S1} \quad \text{Diurnal angular frequency}
Appendix 4.A: analytical solution of DAP

Substitution of \( q = \tilde{\beta} \cdot \exp(-k_d z) \) in Eq. 4.6 allows us to analytically solve the DAP. The result reads as

\[
\frac{1}{d} \int_0^d \tilde{\Lambda}(z) dz = 1 + \frac{1}{d} \left[ E_1(\tilde{\beta}) - E_1(\tilde{\beta} \exp(-\tilde{d})) \right],
\]

in which we used the definition of the exponential integral \( E_1(x) \) (Bender and Orszag, 2013):

\[
E_1(x) = \int_x^\infty \frac{\exp(-q)}{q} dq
\]

where

\[
\tilde{d} = k_d \cdot d.
\]

Consequently, the DAP is defined by two dimensionless numbers \( \tilde{d} \) and \( \tilde{\beta} \).

Appendix 4.B: application of a Taylor expansion to \( \Lambda \): definition of the mfun functions

Consider a function \( f \) that depend on a set of variables \( a = \{k_d, E_0, d\} \). Furthermore, assume that the variables are constructed as a mean value \( \langle k_d \rangle, \langle E_0 \rangle, \langle d \rangle \) plus a (small) variation \( k'_d, E'_0, \text{ and } d' \), respectively. Consequently, the Taylor expansion at the mean values \( \langle k_d \rangle, \langle E_0 \rangle, \langle d \rangle \) upto second-order terms yields

\[
\begin{align*}
&f \left( \langle k_d \rangle + k'_d(t), \langle E_0 \rangle + E'_0(t), \langle d \rangle + d'(t) \right) \approx f \left( \langle a \rangle \right) \\
&+ \frac{\partial f}{\partial k_d} \bigg|_{a=\langle a \rangle} k'_d(t) + \frac{\partial f}{\partial E_0} \bigg|_{a=\langle a \rangle} E'_0(t) + \frac{\partial f}{\partial d} \bigg|_{a=\langle a \rangle} d'(t) \\
&+ \frac{1}{2} \frac{\partial^2 f}{\partial k_d^2} \bigg|_{a=\langle a \rangle} [k'_d(t)]^2 + \frac{1}{2} \frac{\partial^2 f}{\partial E_0^2} \bigg|_{a=\langle a \rangle} [E'_0(t)]^2 \\
&+ \frac{1}{2} \frac{\partial^2 f}{\partial d^2} \bigg|_{a=\langle a \rangle} [d'(t)]^2 + \frac{\partial^2 f}{\partial k_d \partial E_0} \bigg|_{a=\langle a \rangle} k'_d(t) E'_0(t) \\
&+ \frac{\partial^2 f}{\partial d \partial E_0} \bigg|_{a=\langle a \rangle} d'(t) E'_0(t). \quad (4.32)
\end{align*}
\]

Applying this Taylor expansion to the DAP using the following property of the exponential integral \( E_1 \) (Bender and Orszag, 2013):

\[
\frac{dE_1(x)}{dx} = -\frac{\exp[-x]}{x},
\]

(4.33)
yields the definitions of the mfun functions:

\[ mfun_0 = \frac{E_1(\beta) - E_1(e^{-\lambda}\beta)}{\lambda} + 1, \quad (4.34) \]

\[ mfun_{\kappa/\delta} = \frac{1}{mfun_0} \cdot \left[ -e^\beta(e^{-\lambda}) - mfun_0 + 1 \right], \quad (4.35) \]

\[ mfun_\epsilon = \frac{1}{mfun_0} \cdot \left[ \frac{e^{\beta(-e^{-\lambda})} - e^{-\beta}}{\lambda} \right], \quad (4.36) \]

\[ mfun_{\kappa^2/\delta^2} = \frac{1}{mfun_0} \cdot \frac{\beta \lambda}{2} \left( e^{\beta(-e^{-\lambda})} - \lambda + mfun_0 - 1 \right), \quad (4.37) \]

\[ mfun_{\epsilon^2} = \frac{1}{mfun_0} \cdot \frac{1}{2} \left( -\beta e^\beta(-e^{-\lambda}) - e^\beta(-e^{-\lambda}) + e^{-\beta} \beta + e^{-\beta} \right), \quad (4.38) \]

\[ mfun_{e\kappa/\epsilon \delta} = \frac{1}{mfun_0} \cdot \left[ \beta e^\beta(-e^{-\lambda}) - \frac{e^{\beta(-e^{-\lambda})} - e^{-\beta}}{\lambda} \right], \quad (4.39) \]

\[ mfun_{\kappa \delta} = \frac{1}{mfun_0} \cdot \left[ \beta \lambda \left( e^{\beta(-e^{-\lambda})} - \lambda \right) + e^\beta(-e^{-\lambda}) + mfun_0 - 1 \right]. \quad (4.40) \]

In the R programming language, the package `expint` can be used to implement these mfun functions.

**Appendix 4.C: solution of phytoplankton biomass concentration**

We rewrite Eq. (4.1) for depth-averaged phytoplankton biomass concentration \( \tilde{B}(t) \):

\[
\frac{\partial \tilde{B}(t, x)}{\partial t} + \frac{1}{d} \int_0^d \left( \frac{\partial}{\partial x}(\tilde{u}\tilde{B}) + \frac{\partial}{\partial x} \left( K_h \frac{\partial}{\partial x} \tilde{B} \right) \right) \quad \text{change in } \tilde{B}(t) \text{ due to transport} \\
= \tilde{B}(x, t) \cdot [P_{\max} \cdot \Lambda \cdot (1 - \rho_B) - m], \quad (4.41)
\]

Scaling of the longitudinal terms in Eq. (4.41) shows that the longitudinal diffusion term is \( \sim 10^3 \) times smaller than the longitudinal advection term. To obtain the latter result, we assumed that typical scales for \( \tilde{u} \), \( x \), and \( K_h \) are 1 m s\(^{-1}\), 10\(^5\) m, and 10\(^2\) m\(^2\) s\(^{-1}\), respectively (de Swart et al., 2009). Consequently, the change in \( \tilde{B}(t) \) due to transport \( \approx \frac{\partial}{\partial x} \left( \tilde{u}(x, t)\tilde{B}(x, t) \right) \), \( (4.42) \)
where $\tilde{u}$ is the depth-averaged velocity in the longitudinal direction. This results in

$$
\frac{\partial \tilde{B}(t, x)}{\partial t} = \tilde{B}(x, t) \cdot [P_{\text{max}} \cdot \Lambda \cdot (1 - \rho_B) - m]
- \frac{\partial}{\partial x} \left[ \tilde{u}(x, t) \tilde{B}(x, t) \right].
$$

(4.43)

We need further simplification to solve this differential equation analytically. The change in phytoplankton biomass concentration due to advection is given by

$$
\frac{dB}{dt}_{\text{advection}} = - \frac{\partial (\tilde{u} \cdot \tilde{B})}{\partial x}
= - \left[ \tilde{u} \frac{\partial \tilde{B}}{\partial x} + \tilde{B} \frac{\partial \tilde{u}}{\partial x} \right].
$$

(4.44)

We assume that

$$
\tilde{B}(x, t) = B(t) \cdot f(x).
$$

(4.45)

Alternatively put, the phytoplankton biomass concentration $\tilde{B}(x,t)$ only scales to the magnitude of the biomass $B(t)$ in time. Here, $B(t)$ is the local carbon biomass concentration at the location $x = x^*$ where the Chl-a sensor is deployed. The assumption states that the spatial distribution $f(x)$ of $\tilde{B}(x,t)$ is time-independent. For example, if $f(x)$ is linear, we assume it stays linear over time. Because, in this study, we are interested in high-frequency dynamics (\~days$^{-1}$), we argue that this assumption is acceptable. Furthermore, we state that $f(x)$ can be approximated by a linear function (first-order Taylor expansion); we argue that the impact due to advection is a local phenomenon near the location of the Chl-a sensor $x^*$:

$$
\frac{\partial \tilde{B}(x,t)}{\partial x} \approx B(t) \cdot \frac{df(x)}{dx} \bigg|_{x=x^*}.
$$

(4.46)

Similarly, we assume that

$$
\frac{\partial \tilde{u}(x,t)}{\partial x} \approx u(t) \cdot \frac{dg(x)}{dx} \bigg|_{x=x^*},
$$

(4.47)

where $u(t)$ is the water velocity in the $x$ direction towards the mouth. Furthermore, we assume that the water velocity $u(t)$ mainly consists of a subtidal and an M2 tidal constituent:

$$
u(t) = v_0 + A_v \cdot \sin(\omega_{M2} \cdot t + \Phi_{M2} + \theta),
$$

(4.48)

in which $v_0 > 0$ is the residual velocity and $A_v$ is the amplitude of the M2 tidal constituent. To obtain Eq. (4.48), we assumed a phase shift between the horizontal tide (current velocity) and vertical tide (water level $d$) of approximately $-\pi/2$ (van Rijn, 2010). If this is not valid, the results can be generalized using an additional phase shift. This is out of the scope of this study. In Eq. (4.48), $A_v$ has the same sign as $D_2$. If we define $A_v$ as being positive, we have

$$
u(t) = v_0 + |A_v| \cdot \text{sign}(D_2) \cdot \sin(\omega_{M2} \cdot t + \Phi_{M2} + \theta),
$$

(4.49)
where \( \text{sign()} \) denotes the sign function. Combining Eqs. (4.44), (4.46), (4.47), and (4.49), the change in phytoplankton biomass concentration due to advection yields

\[
\left. \frac{dB}{dt} \right|_{\text{advection}} = -\left[ \tilde{v}_0^* + \tilde{\gamma} \cdot \sin(\omega M_2 \cdot t + \Phi M_2 + \theta) \right] \cdot B(t),
\]

(4.50)
in which

\[
\tilde{\gamma} = |A_v| \text{sign}(\hat{D}_2) \cdot \left[ g(x) \left. \frac{df(x)}{dx} \right|_{x=x^*} + f(x) \left. \frac{dg(x)}{dx} \right|_{x=x^*} \right] \text{ and }
\]

(4.51)

\[
\tilde{v}_0^* = v_0 \cdot \left[ g(x) \left. \frac{df(x)}{dx} \right|_{x=x^*} + f(x) \left. \frac{dg(x)}{dx} \right|_{x=x^*} \right].
\]

(4.52)

By filling in the latter results into the differential equation for \( B(x,t) \) in Eq. (4.43) at \( x = x^* \), we have

\[
\frac{\partial B(t)}{\partial t} = B(t) \cdot \left[ P_{\max} \cdot A \cdot (1 - \rho_B) - m \right] + B(t) \cdot \left[ -v_0^* - \gamma \cdot \sin(\omega M_2 \cdot t + \Phi M_2 + \theta) \right],
\]

(4.53)

with

\[
\gamma = |A_v| \text{sign}(\hat{D}_2) \cdot \left[ \left. \frac{df(x)}{dx} \right|_{x=x^*} + \left. \frac{dg(x)}{dx} \right|_{x=x^*} \right] \text{ and }
\]

(4.54)

\[
v_0^* = v_0 \cdot \left[ \left. \frac{df(x)}{dx} \right|_{x=x^*} + \left. \frac{dg(x)}{dx} \right|_{x=x^*} \right],
\]

(4.55)
in which we used \( f(x) \bigg|_{x=x^*} = 1 \) and \( g(x) \bigg|_{x=x^*} = 1 \). The term due to advection in Eq. (4.53) is linear to \( B(t) \). Consequently, this differential equation can be solved analytically. Because the \( v_0^* \)-term results in an \( \exp(v_0^* \cdot t) \) factor, we define the mortality \( m^* \) as

\[
m^* = m + v_0^*.
\]

(4.56)

In the following, we estimate the order of magnitude of \( \gamma \), \( v_0^* \), and \( m^* \) at our case study site. In the Scheldt estuary, the water is pushed several kilometers \( \sim O(1) \) up and down over the longitudinal direction every approximately 12.4 h. So,

\[
\int_0^{6.2h} A_v \sin(M2 \cdot t) dt \sim O(1),
\]

(4.57)

\[
\Rightarrow A_v \sim O(1) \text{ km h}^{-1}.
\]

Furthermore, near our case study site, the phytoplankton biomass concentration increases with a factor of 1-10 \( \sim O(1) \) over a longitudinal distance of approximately 10 km [Maris and Meire, 2016]. Consequently,

\[
\frac{df(x)}{dx} \sim \frac{O(1)}{10 \text{ km}} \sim O(10^{-1}) \text{ km}^{-1}.
\]

(4.58)
Therefore, assuming that the longitudinal gradient in velocity $u$ is negligible ($df/dx \gg dg/dx$), we have

$$
\gamma \approx A_v \cdot \frac{df(x)}{dx} \sim \mathcal{O}(10^{-1}) \ h^{-1}.
$$

Similarly, if we assume $v_0 \sim \mathcal{O}(10^{-1})$ km h$^{-1}$, we have

$$
v_0^* \sim \mathcal{O}(10^{-2}) \ h^{-1}.
$$

Consequently, if we assume $m \sim \mathcal{O}(10^{-3})$ h$^{-1}$ (Desmit et al., 2005), we have

$$
m^* \sim \mathcal{O}(10^{-2}) \ h^{-1}.
$$

**Appendix 4.D: application 2: calibration of the amplitudes and phases**

We calibrate the amplitudes and phases using the postulated expressions:

$$
E'_0(t) = \sum_{n=1}^{3} \hat{E}_n \cos(n \omega_{S1} t),
$$

$$
k'_d(t) = \hat{K}_2 \cos(\omega_{M2} t + \Phi_{M2}) + \hat{K}_4 \cos(\omega_{M4} t + \Phi_{M4}), \quad \text{and}
$$

$$
d'(t) = \hat{D}_2 \cos(\omega_{M2} t + \Phi_{M2} + \theta),
$$

and the data sets of $E_0$, $k_d$, and $d$ at Schellebelle. This calibration results in Figs. 4.D.1a-c, respectively. We utilized the `nls()` function of the R programming language to obtain these data fits. We used subsets to diminish the impact of (low-frequency) temporal variability we did not include in the idealized time dependence in Eqs. (4.62)-(4.64). For example, we excluded long-term periodic temporal fluctuations (e.g., spring-neap) and aperiodic temporal fluctuations (e.g., the impact of clouds).

**Appendix 4.E: time integration of $\Lambda(t)$**

$$
\int \kappa(t) dt = \frac{1}{\langle k_d \rangle} \left[ \frac{\hat{K}_2 \sin (\omega_{M2} t + \Phi_{M2})}{\omega_{M2}} + \frac{\hat{K}_4 \sin (\omega_{M4} t + \Phi_{M4})}{\omega_{M4}} \right]
$$

(4.65)
Figure 4.D.1: The calibration of the postulated time dependencies (solid line) to a subset of the observation (dots) in the Scheldt estuary at Schellebelle (Belgium) of (a) the light extinction coefficient $k_d(t)$, (b) solar irradiance at the water surface $E_0(t)$, and (c) water depth $d(t)$.

\[
\int e(t) dt = \frac{1}{\langle E_0 \rangle} \left[ \frac{\hat{E}_1 \sin (\omega_{S1} t)}{\omega_{S1}} + \frac{1}{2} \frac{\hat{E}_2 \sin (2 \omega_{S1} t)}{\omega_{S1}} + \frac{1}{3} \frac{\hat{E}_3 \sin (3 \omega_{S1} t)}{\omega_{S1}} \right] \tag{4.66}
\]

\[
\int \delta(t) dt = \frac{1}{\langle d \rangle} \left[ \frac{\hat{D}_2 \sin (\omega_{M2} t + \Phi_{M2} + \theta)}{\omega_{M2}} \right] \tag{4.67}
\]
\[
\int \epsilon^2(t) dt = \frac{1}{\langle E_0 \rangle^2} \left[ 1/2 \hat{E}_1^2 t + 1/2 \hat{E}_2^2 t + 1/2 \hat{E}_3^2 t + \frac{\hat{E}_2 \hat{E}_1 + \hat{E}_3 \hat{E}_2}{\omega_{S1}} \sin (\omega_{S1} t) \right.
\]
\[
+ 1/2 \frac{\hat{E}_2 \hat{E}_1 + \hat{E}_3 \hat{E}_1}{\omega_{S1}} \sin (2 \omega_{S1} t) \left. + 1/3 \frac{\hat{E}_2 \hat{E}_1}{\omega_{S1}} \sin (3 \omega_{S1} t) \right. \\
+ 1/5 \frac{\hat{E}_3 \hat{E}_2}{\omega_{S1}} \sin (5 \omega_{S1} t) \\
+ 1/12 \frac{\hat{E}_3^2}{\omega_{S1}} \sin (6 \omega_{S1} t) \\
+ \left. 1/4 \frac{\hat{E}_3 \hat{E}_1 + 1/2 \hat{E}_3^2}{\omega_{S1}} \sin (4 \omega_{S1} t) \right] \\
\] (4.68)

\[
\int \kappa^2(t) dt = \frac{1}{\langle k_d \rangle^2} \left[ 1/4 \frac{\hat{K}_2^2}{\omega_{M2}} \sin (2 \omega_{M2} t + 2 \Phi_{M2}) \right.
\]
\[
+ 1/2 \hat{K}_2^2 t \left. + \hat{K}_2 \hat{K}_4 \sin ((\omega_{M2} - \omega_{M4}) t + \Phi_{M2} - \Phi_{M4}) \right. \\
+ \hat{K}_2 \hat{K}_4 \sin ((\omega_{M2} + \omega_{M4}) t + \Phi_{M2} + \Phi_{M4}) \\
+ 1/4 \frac{\hat{K}_4^2}{\omega_{M4}} \sin (2 \omega_{M4} t + 2 \Phi_{M4}) \\
+ 1/2 \hat{K}_4^2 t \right] \\
\] (4.69)

\[
\int \delta^2(t) dt = \frac{1}{\langle d \rangle^2} \left[ \frac{\hat{D}_2^2}{\omega_{M2}} 1/2 \cos (\omega_{M2} t + \Phi_{M2} + \theta) \times \sin (\omega_{M2} t + \Phi_{M2} + \theta) \right.
\]
\[
+ \hat{D}_2^2 \frac{\omega_{M2} t + \Phi_{M2} + \theta}{2 \omega_{M2}} \left. \right] \\
\] (4.70)
\[
\int \delta(t) \epsilon(t) dt = \frac{\hat{D}_2}{\langle E_0 \rangle \langle d \rangle} \left[ \\
\frac{1}{2} \frac{\hat{E}_1 \sin((\omega_{M2} - \omega_{S1}) t + \Phi_{M2} + \theta)}{\omega_{M2} - \omega_{S1}} \\
+ \frac{1}{2} \frac{\hat{E}_1 \sin((\omega_{M2} + \omega_{S1}) t + \Phi_{M2} + \theta)}{\omega_{M2} + \omega_{S1}} \\
+ \frac{1}{2} \frac{\hat{E}_2 \sin((\omega_{M2} - 2\omega_{S1}) t + \Phi_{M2} + \theta)}{\omega_{M2} - 2\omega_{S1}} \\
+ \frac{1}{2} \frac{\hat{E}_2 \sin((\omega_{M2} + 2\omega_{S1}) t + \Phi_{M2} + \theta)}{\omega_{M2} + 2\omega_{S1}} \\
+ \frac{1}{2} \frac{\hat{E}_3 \sin((\omega_{M2} - 3\omega_{S1}) t + \Phi_{M2} + \theta)}{\omega_{M2} - 3\omega_{S1}} \\
+ \frac{1}{2} \frac{\hat{E}_3 \sin((\omega_{M2} + 3\omega_{S1}) t + \Phi_{M2} + \theta)}{\omega_{M2} + 3\omega_{S1}} \right] \quad (4.71)
\]
Appendix 4.F: the impact of temporal variability in light climate on the onset of a phytoplankton bloom

We estimate the impact of temporal fluctuations in $E_0$, $k_d$, and $d$ on the onset of a phytoplankton bloom using our idealized model. After two weeks, the modeled phytoplankton biomass when no fluctuations are present is approximately a factor of 14 larger than observed. Using Eqs. (4.22) and (4.24), we estimate the difference in $\beta$ which is required to
obtain the same difference in phytoplankton biomass after 14 days:

\[
\frac{B_{\text{no fluct.}}(t = 14\ \text{days}, \beta + \Delta \beta)}{B_{\text{no fluct.}}(t = 14\ \text{days}, \beta)} = 14, \\
\Rightarrow \text{mfun}_0(\lambda, \beta + \Delta \beta) - \text{mfun}_0(\lambda, \beta) = \\
\frac{1}{(1 - \rho_B)P_{\text{max}} 14 \ \text{days}}, \\
\Rightarrow \Delta \beta \approx 1.3. \quad (4.74)
\]

Using Eq. (4.12), we compute the corresponding difference in solar irradiance \( \Delta < E_0 > \approx 31 \ \text{J s}^{-1}\ \text{m}^{-2} \). We use time series of \( E_0 \) observed in 2016 at Stabroek (Fig. 4.F.1) to compute the number of days required to increase the (daily-averaged) solar irradiance \( E_0 \) (low freq. var in Fig. 4.F.1) by 31 J s\(^{-1}\) m\(^{-2}\) in April, which equals approximately two weeks. Consequently, temporal variability in light climate may delay the onset of a phytoplankton bloom by approximately two weeks.


Chapter 5

Applying the sediment transport - phytoplankton model: the disappearance of phytoplankton blooms in spring in the brackish region in the Scheldt estuary

Abstract

Estuaries often show regions in which Chlorophyll-a (Chl-a) accumulates. The location and magnitude corresponding to such accumulation result from a complex interplay between processes such as river flushing, salinity, nutrients, grazing on phytoplankton, and the light climate in the water column. An example is the multi-annual evolution of the estuary-scale Chl-a distribution in the Scheldt estuary (Belgium/Netherlands) in spring. From 2004-2007, we observed a limited spring bloom in the brackish region (km 60-90 from the mouth, salinity $\sim$ 1-10 ppt). This bloom intensified in 2008-2014 and disappeared after 2015. This multi-annual evolution of Chl-a has been linked to simultaneous multi-annual trends in the suspended particulate matter (SPM) distribution and the improvement of the water quality, which affects grazing on phytoplankton by zooplankton. However, this hypothesis has not been systematically investigated. In this contribution, we apply a modeling approach in which observations are the core. We first analyze multi-annual in situ observations covering the full estuary. These observations include the SPM concentration, zooplankton abundance, and other variables affecting the Chl-a concentration. They show a multi-annual estuary-scale evolution not only in the SPM distribution but also in zooplankton abundance, freshwater discharge, and maximum photosynthetic rate. Next, we apply a model approach supported by these observations to constrain the processes and corresponding parameter variability that may have caused the observed change in Chl-a. Our results suggest that a change in SPM alone cannot explain the Chl-a observations. Instead, a multi-annual change in mortality rate, which we can attribute to both grazing by zooplankton and phytoplankton community characteristics (i.e., mortality dependence on salinity), may explain the multi-annual estuary-scale evolution of Chl-a in spring. Different model parameter choices may thus lead to similar model results, which is known as equifinality. Our results highlight that insight into the zooplankton dynamics and phytoplankton community characteristics is essential to understand the phytoplankton (cf. Chl-a) dynamics in the Scheldt estuary and that additional data regarding mortality and grazing rates is required to further constrain the model parameters.

5.1 Introduction

Estuaries regularly exhibit zones with locally elevated Chlorophyll-a (Chl-a) concentrations, which result from a complex interaction between physical, transport-related processes and chemical-biotic factors that determine net local phytoplankton growth. Such processes are governed by water temperature variations (Eppley, 1972), river flushing (Filardo and Dunstan, 1985, Liu and de Swart, 2015), salinity variations (Lucas et al., 1998), grazing on phytoplankton (Alpine and Cloern, 1992, Lionard et al., 2005), nutrient dynamics (Tilman et al., 1982, Cira et al., 2016), and the light climate in the water column (Sverdrup, 1953, Desmit et al., 2005).

Human influences may cause gradual (i.e., multi-annual) changes in multiple of these
interacting processes. Examples are the multi-annual changes in suspended particulate matter (SPM) dynamics caused by channel deepening in the Ems estuary (Winterwerp and Wang, 2013; Dijkstra et al., 2019c) and the multi-annual evolution in nutrients in the Scheldt estuary resulting from an increase in wastewater treatment capacity (Brion et al., 2015). Determining the exact factors that may have caused the observed changes in phytoplankton dynamics is challenging due to the high complexity and because many of the biological interactions are poorly constrained by available data, especially when considering multi-annual time scales.

In view of multi-annual changes in phytoplankton dynamics and the various interacting processes, the Scheldt estuary is an interesting example. Between 2004-2018, the estuary showed the appearance and disappearance of a phytoplankton spring (Apr-May) bloom in the brackish region (km 60-90 from the mouth, salinity ∼ 1-10 ppt) (Maris and Meire, 2017). From 2004 until 2007, almost no spring bloom was observed in the brackish region. Such a spring bloom was consistently observed between 2008-2014 but disappeared after 2015. Covering the same period, Cox et al. (2019) reported a multi-annual estuary-scale change in SPM dynamics in the Scheldt estuary. From 2009 onwards, a change in the estuarine turbidity maximum dynamics and an overall increase in SPM concentration were observed. Simultaneously, the water quality in the Scheldt estuary improved drastically, mainly because of a significant increase in wastewater treatment capacity in Brussels around 2006 (Brion et al., 2015). This resulted in increasing oxygen concentrations and changes in the zooplankton community and abundance. The reported changes in SPM and zooplankton dynamics have been hypothesized to link to the multi-annual disappearance of phytoplankton blooms (Maris and Meire, 2017). However, this has not been systematically investigated, which is necessary given the complex interplay between factors affecting phytoplankton growth.

As discussed by Franks (2009), the choice of an appropriate modeling approach to acquire insight into the phytoplankton(-zooplankton) dynamics depends on the research questions and data availability. Arndt et al. (2011), Naithani et al. (2016), and Gypens et al. (2013) explicitly resolved the phytoplankton-zooplankton(-nutrient) dynamics over one year in the Scheldt estuary using a complex model that includes multiple phytoplankton and zooplankton groups. This resulted in valuable insight into the transient behavior of phytoplankton and zooplankton groups covering the full estuary in 1995, 2003, and 2006, respectively. However, using such models to study multi-annual changes is challenging. The main reason is that (long-term) experimental data is often unavailable, which has three important consequences. Firstly, some of the modeled planktonic groups cannot be observationally validated. Secondly, such models require many (∼ dozens) calibration parameters that are often poorly constrained (e.g., maximum grazing rate, mortality rate per species). These parameters are generally calibrated by fitting them to data and assumed to be fixed in time. Although assuming fixed parameters may be acceptable when focusing on one year, this assumption may be invalid when interested in multi-annual trend changes, suggesting that (some of these) parameters must have changed over time. Thirdly, different model input parameter choices may lead to similar model results, which is known as equifinality. Equifinality has been studied using sediment-transport (van Maren and Cronin, 2016) and planktonic ecosystem models (Friedrichs et al., 2006; 2007) applied to estuarine and marine systems. This phenomenon especially occurs when using
more complex models because the number of model parameters increases by as much as
the square of the number of state variables (Denman and Pea, 2002).

In this contribution, we aim to answer whether the appearance and disappearance of
the phytoplankton bloom in the brackish zone in the Scheldt estuary should be attributed
to changes in SPM or changes in the phytoplankton or zooplankton properties. We choose
our model such that it is mainly data-driven and most of its parameters directly follow
from observations. We aim to minimize the number of variables and calibration param-
eters that we cannot validate using data, requiring the combining of several biological
factors into lumped parameters related to sediment, phytoplankton properties, and zoo-
plankton grazing. Hence, the model is used to constrain which of these combined sets of
processes may explain the observed changes. Although it may still occur using our model
approach, we thereby limit the effect of equifinality.

This contribution is structured as follows. We first introduce the Scheldt estuary, the
methodology to obtain the observations, and the model approach in Section 5.2. In Section
5.3 we show the multi-annual observations of Chl-a and factors impacting phytoplankton
growth in the Scheldt estuary in spring. Next, we present the results of our model
experiments: we calibrate the model, apply a sensitivity analysis of factors that may
explain the disappearance of phytoplankton accumulation in the brackish region, and run
different model scenarios. We study whether this multi-annual trend in phytoplankton
accumulation may be constrained by an individual multi-annual change in grazing by
zooplankton or other processes contributing to the phytoplankton mortality rate. In
Section 5.4 we discuss the model results and approach. Finally, we conclude in Section
5.5.

5.2 Material and methods

5.2.1 Study area

The Scheldt estuary is a funnel-shaped estuary that flows through Belgium into the North
Sea near Vlissingen (Netherlands) over a distance of approximately 160 km (Fig. 5.1). Given its relatively small freshwater discharge compared to the tidal volumes, the Scheldt
estuary is tide-dominated (Baeyens et al., 1997; Meire et al., 2005). The total time-
averaged freshwater discharge \( Q \) in spring (Apr.-May) equaled 85, 81, and 72 m\(^3\) s\(^{-1}\)
zes29f-1066 \( \sim 1 \) km downstream from the Rupel tributary). The main tributaries of the
Scheldt estuary are the Rupel and the Dender. They are responsible for 64.2, 59.3, 63.3
% and 9.4, 9.2, 9.6 % of the total river discharge in 2004-2007, 2008-2014, and 2015-
2018, respectively (Waterinfo.be, 2019, deducted from gauge stations at the upstream
boundary zes57a-1066, at the Dender tributary den02a-1066, and downstream from the
Rupel tributary zes29f-1066 assuming conservation of mass). The Scheldt estuary is a
nutrient-rich estuary (Cox et al., 2009; Maris and Meire, 2017). The dissolved nitrogen,
phosphorous, and silicon concentrations in spring range from 0.1 mmol L\(^{-1}\), 0.001 mmol
L$^{-1}$, and 0.005 mmol L$^{-1}$ at the seaside boundary to 0.4 mmol L$^{-1}$, 0.007 mmol L$^{-1}$, and 0.13 mmol L$^{-1}$ at the upstream boundary, respectively. These concentrations are at least one order of magnitude larger than the half-saturation constants at which we expect nutrient depletion (Billen and Garnier, 1997; Lancelot et al., 2005; Arndt et al., 2011; Naithani et al., 2016).

5.2.2 In situ observations

Both the Belgian and Dutch part of the Scheldt estuary have been monitored intensively over the last two decades. In the Belgian region, various variables have been measured within the multi-annual OMES (Dutch: “Onderzoek Milieu Effecten Sigmaplan”) monitoring program (Maris and Meire, 2017), independently of the tidal phase and spring-neap tide biweekly or monthly at 16 fixed stations (Fig. 5.1). These variables include Chl-a, SPM, salinity, and phytoplankton characteristics, such as the maximum photosynthetic rate $P_{\text{max}}$ and growth efficiency $\alpha$. In the Dutch region, we only use observations of Chl-a and SPM conducted by Rijkswaterstaat at three stations in the main channel (Fig. 5.1). In the following, we briefly introduce the methodology used to obtain the observations presented in this contribution. For a detailed methodological description, we refer the reader to the OMES reports (Maris and Meire, 2017) and the website of Rijkswaterstaat (Rijkswaterstaat, 2020).
5.2.2.1 Chl-a and zooplankton abundance

Within the OMES monitoring program, sub-surface bucket samples were taken to estimate the Chl-a concentration and the mesozooplankton abundance between 2004-2018. The Chl-a concentration was estimated following the spectrophotometric method described in Rice et al. (2017) that corrects for turbidity, Chlorophyll-b, Chlorophyll-c, and Pheophytin pigments, using 50 ml water samples, a 1-cm pathway cuvette, and a Shimadzu UV-1700 spectrophotometer. The observations conducted by Rijkswaterstaat in the Dutch part of the Scheldt estuary were estimated using High-performance liquid chromatography (HPLC) after filtration and extraction.

To estimate the mesozooplankton abundance, 50-250 L sub-surface water samples were collected and filtered over a 50 µm mesh. Next, the mesozooplankton was fixed using formaldehyde and stained with erythrosine in the laboratory. Finally, the organisms were counted in a counting wheel under a binocular microscope using a subsample. A minimum of 500 individuals per subsample was counted (Le Coz et al., 2017). In the brackish region in spring, which is the main focus of this paper, the mesozooplankton community dominantly consists of calanoids (Appeltans, 2003; Mialet et al., 2011). Therefore, we divide the community into two groups: adult calanoids and adult non-calanoids (i.e., cladocerans, harpacticoids, and cyclopoids).

5.2.2.2 Turbidity and SPM concentration

Within the OMES campaign, turbidity depth profiles were measured in 2015-2018 using an Optical Backscatter point Sensor (OBS) of RBR type XR420 CTD+ at the 16 OMES stations. Simultaneously, two SPM samples were collected at approximately the water surface and half the water depth. These SPM samples were used to translate turbidity to SPM concentration (Horemans et al., 2020a). The number of spring campaigns between 2015-2018 equals 16.

To determine the SPM concentration, 1 L water samples were collected and filtered in the laboratory using a GF/C 50 mm filter. To remove salinity, the filters were rinsed with 3 × 50 ml demineralized water before gravimetrically determining the SPM concentrations (norm NBN-EN872). Also within the monitoring program of Rijkswaterstaat, SPM concentrations were gravimetrically determined after filtration on a glass microfiber filter.

5.2.2.3 Light extinction coefficients, temperature, and salinity

The light climate was measured by estimating the light extinction coefficient $k_d$. Two light sensors (LiCOR) measured the light intensity near the water surface $E_1$ and the light-intensity $E_2$ at a fixed distance $\Delta z = 40$ cm from the sub-surface sensor. Next, the light extinction coefficient was estimated as $k_d = \log(E_1/E_2)/\Delta z$, assuming exponential
decrease of light as a function of depth. To correct for small-scale temporal variability (cf. seconds) in the light climate, the time-averaged value of $k_d$ was estimated over a time interval of 3-5 minutes, using a sampling frequency of $1 \text{s}^{-1}$ \cite{Maris_and_Meire,2017}. An estimate at the water surface suffices because, given the high turbidity in the Scheldt estuary, the euphotic depth is relatively small ($\sim \text{dm}$) compared to the total water depth ($\sim \text{m}$). We thus expect phytoplankton growth only near the water surface, where we do not expect strong vertical stratification of SPM.

Temperature and specific conductivity were determined in situ using a WTW LF 318 instrument directly after taking the bucket samples. Specific conductivity was transformed to salinity using the Practical Salinity Scale 1978 \cite{Perkin_and_Lewis,1980}.

### 5.2.2.4 Photosynthetic parameters

To estimate the maximum photosynthetic rate $P_{\text{max}}$ and growth efficiency $\alpha$, the incubation method described in \cite{Kromkamp_and_Peene,1995} was applied using the incubator presented in \cite{Vegter_and_De_Visscher,1984} and assuming a photosynthesis-irradiance (P-I) curve introduced in \cite{Eilers_and_Peeters,1988}. Briefly explained, the Chl-a concentration was determined and water samples were placed at fixed distances from a constant light source. Each distance thus corresponds to a given solar irradiance $I$. Next, the water samples were incubated for approximately 2 hours, while gently being rotated to avoid settling. The photosynthesis was determined using a C-14 isotope method for each $I$, resulting in an estimate of the amount of carbon that is consumed per unit of time per unit of Chl-a. Finally, a P-I curve was constructed to estimate $P_{\text{max}}$ and $\alpha$.

### 5.2.3 Model set-up

Before presenting the technical details of our model approach, we introduce its core characteristic (see Fig. 5.2 for a schematic overview of the model approach). To constrain the processes and parameter variations as much as possible, we require that our model is mainly data-driven and most of its parameters directly follow from the observations. The remaining (lumped) parameters are calibration parameters and are subject to extensive sensitivity study. We do not explicitly resolve planktonic groups that we cannot validate using observations and only include processes and planktonic groups that are shown by the observations to have a dominant impact. Therefore, we only dynamically resolve freshwater and marine diatoms as they are dominantly abundant in the Scheldt estuary in spring \cite{Maris_and_Meire,2007,2009,2009,2013,2017}. With the exception of grazing by zooplankton, all processes contributing to phytoplankton mortality are lumped into parameters $m^{\text{fresh}}$ and $m^{\text{mar}}$ for the freshwater and marine diatoms, respectively, as they cannot be validated by data. Zooplankton is not resolved dynamically but observed zooplankton abundances are directly used in the model, hence eliminating the uncertainty of a dynamic zooplankton model. Observations allow us to distinguish between calanoids and non-calanoids, where calanoids are dominant in the brackish region in spring \cite{Appeltans,2003,Mialet_et_al,2011}. Grazing by
Figure 5.2: To constrain the processes and parameter variations as much as possible, we construct a model such that it is mainly data-driven and most parameters directly follow from the observations. The remaining (lumped) parameters are calibration parameters and are subject to extensive sensitivity analysis. We only dynamically resolve freshwater and marine diatoms as they are dominantly abundant in the Scheldt estuary in spring (Apr.-May). Zooplankton is not resolved dynamically but observed zooplankton abundances are directly used in the model. Observations allow us to distinguish between calanoids and non-calanoids. Grazing by zooplankton is not well constrained and thus included using calibration parameters $g_1$ and $g_2$. The hydrodynamics are explicitly resolved in the vertical and longitudinal direction. The sediment dynamics are both resolved and data-driven. We use the resulting SPM distributions in combination with observed light extinction coefficients to estimate the sediment-induced light extinction coefficient $k_c$. As we are interested in multi-annual changes in sediment characteristics, we also apply a sensitivity study of the Chl-a distribution to $k_c$. As nutrients are abundant, we do not focus on nutrient dynamics.
except possibly for a small part close to the mouth, which is not our area of interest (see Section 5.2.1). The model is thus driven by data-driven time-dependent parameters and four calibration parameters related to phytoplankton ($m_{\text{fresh}}^0, m_{\text{mar}}^0$), and zooplankton characteristics ($g_1, g_2$). To study the individual impact of potential multi-annual changes in SPM and phytoplankton and zooplankton characteristics on the multi-annual evolution of Chl-a accumulation, we consider four model scenarios:

1. We calibrate the parameters $m_{\text{fresh}}^0, m_{\text{mar}}^0, g_1,$ and $g_2$ for the three distinct periods and determine the minimal multi-annual change in these calibration parameters required to capture the accumulation of Chl-a in the brackish region in spring.

2. We test what multi-annual change in sediment characteristics (i.e., $k_c$) is required to capture the accumulation of Chl-a in 2008-2014 in the brackish region assuming no multi-annual change in $m_{\text{mar}}^0, g_1,$ and $g_2$ after 2007. Here, we do not focus on $m_{\text{fresh}}^0$ because we show that marine diatoms dominate the brackish region.

3. We assume a dominant impact of grazing by zooplankton on the mortality rate (i.e., $m_{\text{fresh}}^0 \approx 0$ and $m_{\text{mar}}^0 \approx 0$ s$^{-1}$) and test what multi-annual change in grazing parameters (i.e., $g_1$ and $g_2$) is required to capture the multi-annual evolution of Chl-a accumulation.

4. We neglect the impact of grazing by zooplankton on the mortality rate (i.e., $g_1 \approx 0$ and $g_2 \approx 0$ s$^{-1}$ L) and test what multi-annual change in the mortality rate parameters (i.e., $m_{\text{fresh}}^0$ and $m_{\text{mar}}^0$) is required to capture the multi-annual evolution of Chl-a accumulation.

5.2.3.1 iFlow model

We chose to implement this model in the iFlow model, which we briefly introduce in this section. The iFlow model is a process-based, width-averaged, idealized model (Dijkstra et al., 2017). The model solves for water motion and cohesive SPM trapping in tide-dominated estuaries by resolving the width-averaged shallow water and SPM mass balance equations in equilibrium condition. For this, we use an equidistant grid of 100 cells in the longitudinal and 50 in the vertical direction. The flocculation dynamics of cohesive SPM are resolved using a single-class dynamic flocculation model (Winterwerp, 2002; Horemans et al., 2020a). The model focuses on the estuary-scale hydro- and SPM dynamics only by approximating the estuary’s bathymetry and width by smooth profiles. The model resolves the tidal and subtidal dynamics of water motion and cohesive SPM concentration and provides approximate solutions of the complex and nonlinear set of equations for hydro- and SPM dynamics using a scaling and perturbation approach.

The hydrodynamics are forced at the upstream boundary and two main tributaries by a fixed water inflow and at the mouth by a tidal signal. Following Warner et al. (2005), the longitudinal salinity profile is implemented as a tide- and depth-independent profile (see the Appendix 5.A). This assumption is consistent with the Scheldt estuary being well-mixed (Baeyens et al., 1997). The SPM dynamics are forced by a constant inflow of
SPM that equals the product of the water discharge and subtidal SPM concentration at the upstream boundary, and by a fixed SPM concentration at the mouth. We assume that erosion of sediment scales to the magnitude of the bed shear stress. For details on the performance of this model for the Scheldt estuary, we refer to Horemans et al. (2020a). For this study, it suffices to mention that the M2 tidal and subtidal surface elevation correspond well to observations and that the magnitude of the SPM concentration and location of the estuarine turbidity maxima are reproduced.

We adapted iFlow’s phytoplankton module (Dijkstra et al., 2019a). The width-averaged differential equation for the phytoplankton concentration $$P_i$$ of phytoplankton group $$i$$ and corresponding boundary conditions read as (Dijkstra et al., 2019a)

$$
\partial_t P_i + u \partial_x P_i + (w - w_P) \partial_z P_i - \frac{1}{B} \partial_x (B K_h \partial_x P_i) - \partial_x (K_v \partial_z P_i)
\right)
\underbrace{\text{advection-diffusion}}_{(\mu - m)P_i},
\right)
\underbrace{\text{balance between local growth and mortality}}_{(\mu - m)P_i},
\right)

\tag{5.1}
$$

Here, $$t$$ represents time, $$x$$ and $$z$$ are the coordinates in the longitudinal and vertical direction, $$u$$ and $$w$$ are the water velocities in the longitudinal and vertical direction, $$w_P$$ is the constant settling velocity of phytoplankton cells, $$B$$ is the width of the estuary, $$K_h$$ and $$K_v$$ are the horizontal and vertical eddy diffusivities, the angle brackets denote averaging over a long time scale (i.e., larger than a tide or day ~ 1-2 weeks), $$-H$$ and $$\zeta$$ are the z-coordinates of the bed and water surface, $$P_{sea}$$ is the constant phytoplankton concentration at the seaside boundary, $$QP$$ is the constant influx of phytoplankton at the upstream boundary, and $$\mu$$ and $$m$$ are the growth and mortality rate of phytoplankton. We divide the model into two phytoplankton classes: freshwater diatoms $$P_{\text{fresh}}$$ and marine diatoms $$P_{\text{max}}$$. Following Naithani et al. (2016) most parameters of the two phytoplankton groups are equal, except the mortality rate $$m$$ and maximum growth rate $$\mu_{\text{max}}$$, which is ~ 1.6 times larger for marine diatoms. The mortality rate depends on salinity $$S$$ and the abundance of phytoplankton grazers $$Z$$:

$$
m = m^i_0 f_S(S) + f_Z(Z),
\right)
\tag{5.3}
$$

in which $$m^i_0$$ is a (calibrated) constant mortality rate parameter of phytoplankton group $$i$$ (i.e., the freshwater or marine diatoms) and $$f_S$$ and $$f_Z$$ are functions that determine the salinity and zooplankton dependence of the mortality rate $$m$$, respectively. Following Naithani et al. (2016) we assume the following (normalized) salinity stress:

$$
f_S(S) = \begin{cases} 
1.07^{1.07^{S_{\text{max}}}} & \text{freshwater diatoms,} \\
1.07^{5 \times 0.85^{S_{\text{upstream}}}} & \text{marine diatoms,} 
\end{cases}
\tag{5.4}
$$
in which $s_{sea}$ and $S_{upstream}$ are the salinity at the downstream and upstream boundary (in ppt), respectively. For the zooplankton dynamics, we follow a data-driven approach and parameterize the corresponding grazing by zooplankton. We include the two dominant zooplankton groups $Z_{calanoids}$ and $Z_{non-calanoids}$ (units ind. L$^{-1}$, where 'ind.' denotes 'individuals'), which directly follow from the observations. More specifically, we linearly interpolate the zooplankton abundance observations and extrapolate the zooplankton abundance in the downstream region where we do not have observations using the system-averaged abundance. In the literature, multiple zooplankton dependencies of the phytoplankton mortality rate have been studied (Steele and Henderson, 1992). We consider the following longitudinal variation in $m$ due to zooplankton abundance:

$$f_Z(Z_{calanoids}, Z_{non-calanoids}) = g_1 Z_{calanoids}(x) + g_2 Z_{non-calanoids}(x)$$  \hspace{1cm} (5.5)

in which $g_1$, and $g_2$ are grazing parameters that follow from calibration (units s$^{-1}$ L$^{-1}$).

Given that the Scheldt estuary is a turbid system, we use the Platt formulation for light limitation of the time-averaged growth rate $\mu$. This formulation is suitable for turbid systems as it does not consider an inverse relationship between $\mu$ and the photosynthetically active radiation (PAR) $E$ at large $E$ (cf. cell burning). Without nutrient limitation, $\mu$ then reads

$$\mu = \mu_{max}(T) \left[ 1 - \exp \left( \frac{\alpha}{P_{max}} E \right) \right],$$  \hspace{1cm} (5.6)

in which $T$ is the water temperature and the angle brackets again denote averaging over a long time scale (i.e., larger than a tide or day). The photosynthetically active radiation $E$ reads as

$$E(z, t; P^i, c) = E_{00}(t) \exp \left( k_{bg} z - k_c \int_0^z c(z, t) dz - k_P \int_0^z P^i(z, t) dz \right) \times$$

$$\begin{cases} \sin [\omega_E \ mod (t, 24h)] & \text{if } \ mod (t, 24h) < \pi/\omega_E, \\ 0 & \text{elsewhere} \end{cases}$$

\hspace{1cm} (5.7)

in which $E_{00}$ represents the maximum PAR during mid-day, $k_{bg}$, $k_c$, and $k_P$ are the background, sediment-induced, and self-shading exponential light extinction coefficients, respectively, $c$ is the SPM concentration, $\omega_E$ is the angular frequency for day length, and mod denotes the modulus operator. Following Eppley (1972), we postulate the following temperature dependence of the maximum growth rate $\mu_{max}(T)$:

$$\mu_{max}(T) = \mu_{00} \mu_{01} \left( \frac{T}{15^\circ C} \right),$$

\hspace{1cm} (5.8)

in which $\mu_{00}$ and $\mu_{01}$ are calibration parameters and $T$ is expressed in $^\circ C$.

As with the hydro- and SPM dynamics, the model solves the approximated phytoplankton dynamics in equilibrium conditions (Dijkstra et al., 2019a). By doing so, we do
not have to postulate initial conditions, which further simplifies our sensitivity analysis. We argue that this assumption of equilibrium conditions is acceptable because the accumulation of phytoplankton in the brackish region covers approximately two months, which is large compared to the time scale of a bloom (∼2-3 weeks). As shown by Regnier et al. (1997), the accuracy of their coupled reaction-transport in equilibrium conditions applied to the Scheldt estuary depends on the biological rates; higher rates (which are typical for the spring/summer months) result in higher model performance. We solve the marine and freshwater diatom dynamics separately and thereby neglect their coupling through shading by marine diatoms on freshwater diatoms and vice versa. This assumption is acceptable as we show later that freshwater and marine diatoms are spatially separated. In the region where we have similar concentrations of freshwater and marine (cf. coupling), self-shading is negligible due to the high SPM concentrations in this region. Last, for the implementation of time-averaged $\mu$, we use the approximated Platt light limitation function presented in Horemans et al. (2020b). By solving approximate solutions for the phytoplankton dynamics, our model approach comes with very low computation times (∼$s$) when compared to more realistic models (∼hours-days), allowing for an extensive sensitivity analysis.

5.2.3.2 Calibration and parameter values

To determine potential environmental changes that may have caused the disappearance of phytoplankton accumulation in the brackish region, we apply a sensitivity study and run four different model scenarios (see Section 5.2.3). Here, most model parameter values directly follow from observations. Using the $P_{max}$ and temperature observations, we derive the calibration parameters $\mu_{00}$ and $\mu_{01}$, which contain the temperature dependence of $\mu_{max}$ [Eq. (5.8), see the Appendix 5.B]. The influx of phytoplankton at the upstream boundary $QP$ follows from the Chl-a observations at the upstream boundary. To correct for the large temporal variability in discharges [Waterinfo.be, 2019] and correctly resolve the sediment dynamics in spring, we use the procedure of Horemans et al. (2020a) to determine representative values of the freshwater discharge and determine the erosion and flocculation characteristics. This results in a representative modeled SPM distribution in 2004-2007, 2008-2014, and 2015-2018. We refer the reader to the Appendix 5.C for the technical details. To ensure that we correctly model light extinction, we use these modeled SPM distributions in combination with the observations of the light extinction coefficient $k_d$ to estimate the sediment-induced light extinction coefficient $k_c$ (see the Appendix 5.D for the technical details). We estimate the grazing parameters $g_1$ and $g_2$ corresponding to the calanoids and non-calanoids, respectively, and mortality rate parameters $m_0^{fresh}$, $m_0^{mar}$ corresponding to freshwater and marine diatoms, respectively, by calibrating modeled Chl-a concentrations to the Chl-a observations. Here, we again use the calibration method described in Horemans et al. (2020a) in which the phytoplankton model results and observations (cf. Chl-a) are quantitatively compared. The model parameters that are the focus of this paper are summarized in Table 5.1. The full parameter list is provided in Table 5.A.1. The reported values follow from observations, calibration, Dijkstra et al. (2019a), Horemans et al. (2020a) and other published studies.
Table 5.1: Parameter values used in our model experiments based on observations (source), model calibration (calibrated), and the literature. If only one parameter value is presented, we used this value for all three periods.

<table>
<thead>
<tr>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mu_{\text{fresh.00}}$</td>
<td>Maximum growth rate at 0 °C of freshwater diatoms (source)</td>
<td>$0.96 \times 10^{-5}$</td>
<td>$1.04 \times 10^{-5}$</td>
<td>$0.86 \times 10^{-5}$</td>
<td>s$^{-1}$</td>
</tr>
<tr>
<td>$\mu_{\text{mar.00}}$</td>
<td>Maximum growth rate at 0 °C of marine diatoms (source)</td>
<td>$1.59 \times 10^{-5}$</td>
<td>$1.72 \times 10^{-5}$</td>
<td>$1.43 \times 10^{-5}$</td>
<td>s$^{-1}$</td>
</tr>
<tr>
<td>$\mu_{01}$</td>
<td>Calibration parameter in temperature dependence for $\mu_{\text{max}}$ (source)</td>
<td>1.10</td>
<td>1.07</td>
<td>1.05</td>
<td>/</td>
</tr>
<tr>
<td>$g_1$</td>
<td>Calanoids grazing parameter (calibrated)</td>
<td>$0.8 \times 10^{-7}$</td>
<td>$0.13 \times 10^{-7}$</td>
<td>$0.8 \times 10^{-7}$</td>
<td>s$^{-1}$ L</td>
</tr>
<tr>
<td>$g_2^{\text{fresh.}}$</td>
<td>Non-calanoids grazing parameter (calibrated)</td>
<td>$0.47 \times 10^{-7}$</td>
<td>$0.32 \times 10^{-7}$</td>
<td>$0.47 \times 10^{-7}$</td>
<td>s$^{-1}$ L</td>
</tr>
<tr>
<td>$m_{0}^{\text{fresh.}}$</td>
<td>Mortality rate parameter for freshwater diatoms (calibrated)</td>
<td>$1.89 \times 10^{-6}$</td>
<td>$3.30 \times 10^{-6}$</td>
<td>$8.30 \times 10^{-6}$</td>
<td>s$^{-1}$ L</td>
</tr>
<tr>
<td>$m_{0}^{\text{mar.}}$</td>
<td>Mortality rate parameter for marine diatoms (calibrated)</td>
<td>$3.21 \times 10^{-6}$</td>
<td>$1.06 \times 10^{-6}$</td>
<td>$3.35 \times 10^{-6}$</td>
<td>s$^{-1}$ L</td>
</tr>
<tr>
<td>$E_{\text{max}}$</td>
<td>Maximum PAR (source)</td>
<td>1007</td>
<td></td>
<td></td>
<td>μmol photons m$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>$\omega_E$</td>
<td>Angular frequency for day length (source)</td>
<td>0.215</td>
<td></td>
<td></td>
<td>d$^{-1}$</td>
</tr>
<tr>
<td>$T$</td>
<td>Water temperature (source)</td>
<td>14.3</td>
<td>14.7</td>
<td>14.7</td>
<td>°C</td>
</tr>
<tr>
<td>$P_{\text{sea}}$</td>
<td>Phytoplankton boundary concentration at the mouth (source)</td>
<td>15.9</td>
<td>17.1</td>
<td>15.8</td>
<td>μg L$^{-1}$</td>
</tr>
<tr>
<td>$Q \eta$</td>
<td>Influx of phytoplankton at the upstream boundary (source)</td>
<td>1.5</td>
<td>1.8</td>
<td>2.5</td>
<td>g s$^{-1}$</td>
</tr>
<tr>
<td>$w_P$</td>
<td>Settling velocity of phytoplankton cells (Sarthou et al., 2005)</td>
<td>1.15 \times 10^{-5}</td>
<td></td>
<td></td>
<td>m s$^{-1}$</td>
</tr>
<tr>
<td>$k_{\text{bg}}$</td>
<td>Background exponential light extinction coefficient (Pennock and Sharp, 1994)</td>
<td>0.095</td>
<td></td>
<td></td>
<td>m$^{-1}$</td>
</tr>
<tr>
<td>$k_P$</td>
<td>Self-shading exponential light extinction coefficient (Pennock and Sharp, 1994)</td>
<td>18</td>
<td></td>
<td></td>
<td>m$^2$ (mol N)$^{-1}$</td>
</tr>
<tr>
<td>$k_c$</td>
<td>Sediment-induced exponential light extinction coefficient (source)</td>
<td>81.4</td>
<td>77.9</td>
<td>72.0</td>
<td>m$^2$ kg$^{-1}$</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Growth efficiency (source)</td>
<td>0.0165</td>
<td>0.0168</td>
<td>0.0188</td>
<td>mg C (mg Chl-a)$^{-1}$ h$^{-1}$</td>
</tr>
</tbody>
</table>

Note: All units are in scientific notation unless otherwise specified.
5.3 Results

5.3.1 Evolution of Chl-a and corresponding environmental conditions of the in situ observations

5.3.1.1 Evolution of Chl-a and zooplankton

Figure 5.3 presents the evolution of the sub-surface Chl-a concentration in 2004-2018, showing a clear seasonality and corresponding phytoplankton blooms; at the upstream boundary (∼ km 160), the Chl-a concentration can reach values above 400 µg L$^{-1}$ in summer and, although local maxima are observed, decays in the downstream direction. We divide the time series into three distinct periods and focus on the time-averaged Chl-a concentration in spring (Apr.-May) (Fig. 5.3). In 2004-2007, we detect time-averaged Chl-a concentrations above 50 µg L$^{-1}$ in the upstream region, > km 80. In 2008-2014 and 2015-2018, this region is limited to > 100 km and > 110 km, respectively. In 2008-2014, we also observe concentrations > 50 µg L$^{-1}$ more downstream in the brackish region between km 60-90. The Chl-a concentrations are significantly larger in 2008-2014 in the brackish region compared to the concentrations in 2004-2007 (Welch t-test, p-value < 10$^{-3}$) and 2015-2018 (Welch t-test, p-value < 10$^{-12}$).

Figure 5.4 displays the time-averaged calanoids and non-calanoids abundance in spring for the three distinct periods considered. The shaded area depicts the standard error of the zooplankton abundance. The calanoids abundance (Fig. 5.4a) also shows distinct trends in the three periods considered. In 2004-2007, we observe a relatively low mean calanoids abundance between km 110-150, ranging up to approximately 5 ind. L$^{-1}$. Downstream from km 110, we observe an increase in calanoids, resulting in a local maximum of the mean values of calanoids of approximately 10 ind. L$^{-1}$, centered near km 90. In 2008-2014, the local maximum of the mean values in calanoids abundance shifts in the upstream direction and increases. The overall calanoid abundance increases, with a maximum of the mean values of approximately 17.5 ind. L$^{-1}$ at km 110. In 2015-2018, the local maximum of the mean values in calanoids abundance shifts further landwards to approximately km 140, with again a maximum of approximately 17.5 ind. L$^{-1}$. We thus observe a land-inward shift and estuary-scale increase of the local calanoids abundance over time. The calanoids concentrations are significantly larger between km 110-150 in 2015-2018 compared to 2004-2007 (Welch t-test, p-value < 10$^{-4}$). However, the system-averaged, volume-weighted calanoids abundances are 4.9 ind. L$^{-1}$, 6.43 ind. L$^{-1}$, and 4.3 ind. L$^{-1}$ in 2004-2007, 2008-2014, and 2015-2018, respectively. With ‘volume-weighted’, we mean that we weigh the abundance with the water volume corresponding to the location at which the abundance is estimated. So, we take into account the fact that the funnel-shaped estuary is narrow and shallow in the upstream region and we thus attribute more weight to observations in the downstream direction. At the upstream boundary, non-calanoids are dominantly present (Fig. 5.4b). On average, we observe an increase of the non-calanoids abundance in the landward direction on the estuary scale in all three periods. As illustrated by the large standard error, the differences of the non-calanoids
Figure 5.3: (a) Observed Chl-a concentration (µg L−1) in 2004-2018 and (b) time-averaged Chl-a concentration in spring. We observe a phytoplankton bloom in the brackish region (km 60-90) in spring in 2008-2014, which is absent in the other years considered. The Chl-a concentration also decreases faster in the downstream direction in more recent years (illustrated by the horizontal arrows).

5.3.1.2 Evolution of SPM and light extinction

Figure 5.5a shows the sub-surface time-averaged SPM concentration in the three periods considered in spring. In all three periods, the SPM concentrations range up to approximately 150 mg L−1. However, we observe significantly lower concentrations between approximately km 50-100 in 2004-2007 (Welch t-test, p-value < 10−5 and < 10−6 when compared to 2008-2014 and 2015-2018, respectively). The lower SPM concentrations are especially visible between km 70-80, where we have concentrations below 50 mg L−1 in 2004-2007 and up to 150 mg L−1 after 2007. Moreover, in 2015-2018, we observe the largest SPM concentrations between km 80-120 (Welch t-test, p-value < 10−3 and < 10−4 when compared to 2008-2014 and 2004-2007, respectively).

The time-averaged light extinction coefficient in spring shows a similar evolution to the SPM concentration (Fig. 5.5b), with significantly lower values of approximately 4 m−1 between km 50-100 in 2004-2007 compared to the values of approximately 7 m−1 after 2007 (Welch t-test, p-value < 10−8 and < 10−10 when compared to 2008-2014 and 2015-2018, respectively). We have the largest time-averaged values between km 80-120 in 2015-2018, which is consistent with the SPM observations (Welch t-test, p-value = 3.4 × 10−2 and < 10−5 when compared to 2008-2014 and 2004-2007, respectively).
Figure 5.4: Multi-annual time averages of (a) calanoids and (b) non-calanoids abundance in spring (Apr.-May). The shaded area depicts the standard error. We observe a dominant abundance of calanoids in the brackish region and a land-inward shift of calanoids in time. The non-calanoids are mainly situated at the upstream boundary.

Figure 5.5: Multi-annual time-averaged observations in spring (Apr.-May) in 2004-2018 of (a) the water surface SPM concentration and (b) the light extinction coefficient $k_d$. The error bars depict the standard error of the observations.

5.3.1.3 Evolution of discharge, salinity intrusion, and photosynthetic parameters

In spring (Apr.-May), the average discharge is 85, 81, and 72 m$^{-3}$ s$^{-1}$ in 2004-2007, 2008-2014, and 2015-2018, respectively. We thus observe a slight decrease in total freshwater discharge over time. We define the salinity intrusion as the distance from the mouth
at which the salinity equals 2 ppt. The corresponding time-averaged values in spring are 81 km, 79 km, and 83 km in 2004-2007, 2008-2014, and 2015-2018, respectively. In spring, the salinity intrusion does not show major changes during the study period 2004-2018. The time-and system-averaged maximum photosynthetic rate $P_{\text{max}}$ in spring is approximately equal in 2004-2007 and 2008-2014, but significantly lower in 2015-2018. The corresponding time-averaged values are 6.59, 6.44, and 4.31 mg C (mg Chl-a)$^{-1}$ h$^{-1}$, respectively. The corresponding time-and system-averaged growth efficiency $\alpha$ are 0.0165, 0.0168, and 0.0188 mg C (mg Chl-a)$^{-1}$ h$^{-1}$ [µmol photons m$^{-2}$ s$^{-1}$]$^{-1}$, respectively. For the monthly averaged data of the discharge, salinity intrusion, $P_{\text{max}}$, and $\alpha$ covering the full year, we refer the reader to the Appendix 5.E.

Multi-annual observations in the Scheldt estuary thus show an estuary-scale increasing or decreasing development in zooplankton abundance, SPM (cf. light extinction), $P_{\text{max}}$, and, to a minor extent, in the freshwater discharge in the Scheldt estuary in 2004-2018 in spring. Based on the observations alone, it is difficult to determine the change in parameters that is responsible for the evolution of Chl-a concentration. Therefore, we apply a complementary model approach in the following section.

5.3.2 Evolution of Chl-a studied using model experiments

To quantify the impact of the observed trends presented in the previous section and alterations of other factors affecting phytoplankton growth on the Chl-a concentration, we consider the four model scenarios presented in Section 5.2.3.

5.3.2.1 Scenario 1: minimum multi-annual change in calibration parameters required to capture the accumulation of Chl-a in the brackish region in spring

We calibrate the mortality rate parameters $m_{0}^{\text{fresh}}$ and $m_{0}^{\text{mar}}$ and grazing parameters $g_1$ and $g_2$ to the observed multi-annual time-averaged Chl-a concentrations for the three periods considered. The corresponding values are listed in Table 5.1. In 2004-2007 and 2015-2018, we capture the estuary-scale patterns of Chl-a by keeping the parameters $m_{0}^{\text{mar}}$, $g_1$, and $g_2$ more or less fixed and only changing $m_{0}^{\text{fresh}}$ (Figs. 5.6a and 5.6c). We require a significantly larger mortality rate parameter of the freshwater diatoms in 2015-2018 than in 2004-2007 ($m_{0}^{\text{fresh}} = 8.30 \times 10^{-6}$ versus $m_{0}^{\text{fresh}} = 1.89 \times 10^{-6}$ s$^{-1}$, respectively) to capture the faster decrease of the Chl-a concentrations in the downstream direction over the years 2004-2018 (depicted by the horizontal arrows in Fig. 5.3b). Here, it is important to note that the observed Chl-a values between km 50 and 100 are below the detection limit of 10 µg L$^{-1}$ and all modeled Chl-a concentrations lower than this limit are considered as equally good in the calibration. In 2008-2014, we only obtain the accumulation of Chl-a in the brackish region if we also assume a multi-annual evolution in parameters $m_{0}^{\text{mar}}$, $g_1$, and $g_2$ (see Table 5.1). The calibrated $m_{0}^{\text{mar}}$ and $g_1$ values are $\sim$ 3 and $\sim$ 7 times lower, respectively. If we were to assume no multi-annual evolution of parameters $m_{0}^{\text{mar}}$, $g_1$, and $g_2$ after 2007, we would not capture the accumulation of Chl-a.
Figure 5.6: Multi-annual time-averaged Chl-a observations (dots) and depth-averaged model result (dashed line) in spring (Apr.-May) in (a) 2004-2007, (b) 2008-2014, and (c) 2015-2018 (Scenario 1). (d) When we do not consider a multi-annual evolution of parameters $m_0^{\text{mar.}}$, $g_1$, and $g_2$ (Scenario 2), we do not capture the estuary-scale Chl-a distribution in 2008-2014 beyond km 60 (for which we have zooplankton data). A sensitivity analysis shows that by decreasing $k_c$ by a factor $\sim 3$ ($k_c = 25$ versus $k_c = 78 \text{ m}^2 \text{kg}^{-1}$), we also obtain accumulation of Chl-a in the brackish region. However, this difference is significantly larger than the variability of $k_c$ that follows from the observations.

in the brackish region (Fig. 5.6d). To summarize, to capture accumulation of Chl-a in the brackish region in 2008-2014, we require a (significant) multi-annual change in parameters $m_0^{\text{fresh.}}$, $m_0^{\text{mar.}}$, $g_1$, and $g_2$.  

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5.3.2.2 Scenario 2: multi-annual change in sediment characteristics required to capture the accumulation of Chl-a in the brackish region in spring

Keeping all parameters fixed to their values presented in Table 5.1 and assuming no multi-annual change after 2007 in parameters $m_{0\text{mar}}, g_1$, and $g_2$, a careful sensitivity analysis shows that variability in $\mu_0, Q, P_{\text{sea}},$ and $QP$ does not result in accumulation of Chl-a in the brackish region in 2008-2014 (for the details, see the Appendix 5.F). Only by decreasing $k_c$ by a factor $\sim 3$ ($k_c = 25$ versus $k_c = 78 \text{ m}^2 \text{ kg}^{-1}$), we obtain accumulation of Chl-a in the brackish region (Fig. 5.6d). This difference is significantly larger than the variability that follows from the observations, which is between $\sim 65$ and $80 \text{ m}^2 \text{ kg}^{-1}$ (for the details, see the Appendix 5.D). Therefore, a multi-annual change in sediment characteristics alone cannot explain the multi-annual evolution in the Chl-a distribution.

5.3.2.3 Scenario 3: the individual effect of grazing by zooplankton

In this section, we assume a dominant impact of grazing by zooplankton on the mortality rate (i.e., $m_{0\text{fresh}} \approx 0$ and $m_{0\text{mar}} \approx 0 \text{ s}^{-1}$). Calibration of the grazing parameters $g_1$ and $g_2$ to the Chl-a observations in 2004-2007 and 2015-2018, and the Chl-a observations in the brackish region only in 2008-2014 results in the modeled Chl-a concentration presented in Figs. 5.7a, 5.7c, and 5.7b, respectively. Our calibration results in larger grazing parameters because we neglected other processes contributing to the mortality rate (e.g., salinity stress). The grazing parameters are $g_1 = 2.5$ and $g_2 = 0.93 \times 10^{-7} \text{ s}^{-1} \text{ L}$, $g_1 = 0.51$ and $g_2 = 0.71 \times 10^{-7} \text{ s}^{-1} \text{ L}$, and $g_1 = 2.5$ and $g_2 = 0.93 \times 10^{-7} \text{ s}^{-1} \text{ L}$ in 2004-2007, 2008-2014, and 2015-2018, respectively. In 2004-2007 and 2015-2018, although we detect some local anomalies (for example, the underestimation near km 90 in 2004-2007), the model captures the Chl-a distribution on the estuary-scale using the same $g_1$ and $g_2$ values. In contrast, if we were to choose these calibrated grazing parameter values in 2008-2014, we would obtain a Chl-a distribution very similar to the case presented in Fig. 5.6d (all diatoms) and we would thus not capture the accumulation of Chl-a in the brackish region (not shown). Considering different values for $g_1$ and $g_2$ in 2008-2014, we can again model the estuary-scale Chl-a patterns. Here, the local minimum near km 100 is due to a local increase in mortality rate resulting from the high calanoid abundance in this region. If we were to consider only the calanoids (so a reduction of one calibration parameter), we would not obtain the clear local minimum in Chl-a near km 100 (not shown). Finally, choosing the $g_1$ and $g_2$ values corresponding to 2008-2014 in 2015-2018 results in a system-scale overestimation of Chl-a (not shown). To summarize, when only including the effect of grazing by zooplankton, we again require a (significant) multi-annual evolution of $g_1$ and $g_2$ to capture the accumulation of Chl-a in the brackish region in spring in 2008-2014.
Figure 5.7: Multi-annual time-averaged Chl-a observations (dots) and depth-averaged model result (dashed/solid line) in spring (Apr.-May) in (a) 2004-2007, (b) 2008-2014, and (c) 2015-2018 assuming a mortality rate exclusively caused by grazing (Scenario 3, denoted by ‘grazing only’) and neglecting the effect of grazing by zooplankton (Scenario 4, denoted by ‘freshwater diatoms’, ‘marine diatoms’, and ‘all diatoms’). Except for the Chl-a concentrations downstream from km 60 in 2008-2014 (for which we do not have zooplankton data), the model captures the estuary-scale Chl-a distribution. We can thus obtain the multi-annual evolution in Chl-a accumulation in the brackish region in spring (Apr.-May) by both a change in phytoplankton community characteristics and grazing by zooplankton. In Scenario 3 (cf. ‘grazing only’), we attribute the local minimum near km 100 in 2008-2014 to a local increase in mortality rate resulting from the high calanoid abundance in this region. In Scenario 4 (cf. ‘all diatoms’), this local minimum in Chl-a results from the spatial separation between marine and freshwater diatoms that is caused by salinity stress.
5.3.2.4 Scenario 4: neglecting the effect of grazing by zooplankton

In this section, we neglect the impact of grazing by zooplankton to the mortality rate (i.e., \( g_1 \approx 0 \) and \( g_2 \approx 0 \) s\(^{-1}\) L). We calibrate the mortality rate parameters \( m_0^{\text{fresh.}} \) and \( m_0^{\text{mar.}} \), while keeping all other parameters fixed to the calibrated values presented in Table 5.1 (Figs. 5.7a-5.7c). The calibration results in larger mortality rate parameters, which is due to the absence of grazing pressure. In 2004-2007, we capture the large-scale pattern of the Chl-a distribution using a \( \sim 70 \% \) larger mortality rate parameter for the marine diatoms \( (m_0^{\text{fresh.}} = 3.8 \times 10^{-6} \text{ versus } m_0^{\text{mar.}} = 6.4 \times 10^{-6} \text{ s}^{-1}) \). In 2008-2014, the model captures the Chl-a distribution beyond km 59 and the local minimum near \( \sim \) km 100. This local minimum results from a clear spatial separation between marine and freshwater diatoms that is caused by salinity stress. In Scenario 3, we attributed this minimum to a local increase in mortality rate resulting from the high calanoid abundance in this region. The model overestimates the Chl-a concentration in the marine region at \( \sim \) km 21 and 36. The accumulation of Chl-a in the brackish region mainly corresponds to marine diatoms. This accumulation requires a \( \sim 3 \) times lower mortality rate parameter for the marine diatoms \( (m_0^{\text{fresh.}} = 6.6 \times 10^{-6} \text{ versus } m_0^{\text{mar.}} = 2.1 \times 10^{-6} \text{ s}^{-1}) \). Upstream from the local minimum at \( \sim \) km 100, we mainly have freshwater diatoms. In 2015-2018, we again model the Chl-a distribution accurately on the estuary-scale and have a clear spatial separation between freshwater and marine diatoms. The marine diatoms have a significantly lower mortality rate parameter \( (m_0^{\text{fresh.}} = 6.7 \times 10^{-6} \text{ versus } m_0^{\text{mar.}} = 16.6 \times 10^{-6} \text{ s}^{-1}) \). In the following, we focus on the multi-annual evolution of the calibration parameters. The mortality rate parameter corresponding to the marine diatoms is equal in 2004-2007 and 2015-2018 \( (m_0^{\text{mar.}} = 6.5 \times 10^{-6} \text{ s}^{-1}) \), but significantly lower in 2008-2014 \( (m_0^{\text{mar.}} = 2.1 \times 10^{-6} \text{ s}^{-1}) \). As found before, the model also shows a multi-annual increase of \( m_0^{\text{fresh.}} \). For the freshwater diatoms, we have \( m_0^{\text{fresh.}} = 3.8 \times 10^{-6} \text{ s}^{-1}, m_0^{\text{fresh.}} = 6.6 \times 10^{-6} \text{ s}^{-1}, \) and \( m_0^{\text{fresh.}} = 16.6 \times 10^{-6} \text{ s}^{-1} \) in 2004-2007, 2008-2014, and 2015-2018, respectively. To summarize, when excluding the effect of grazing on the mortality rate, we again require a (significant) multi-annual evolution of \( m_0^{\text{fresh.}} \) and \( m_0^{\text{mar.}} \) to capture the accumulation of Chl-a in the brackish region in spring in 2008-2014.

5.4 Discussion

5.4.1 Suggested importance of grazing and phytoplankton community characteristics

We studied the appearance and disappearance of accumulation of Chl-a in the brackish region of the Scheldt estuary in spring in 2008-2014. To this end, we analyzed multi-annual observations of factors affecting phytoplankton growth and ran various model scenarios. The model approach allowed us to detect which combination of multi-annual parameter change may result in the multi-annual evolution of the Chl-a concentrations. Our results suggest that we require a multi-annual shift in phytoplankton mortality rate to capture the appearance and disappearance of Chl-a accumulation in the brackish region and that
other parameters (e.g., SPM) alone cannot explain this observed trend of Chl-a.

The multi-annual evolution in mortality rate may be attributed to either a change in phytoplankton community characteristics or grazing by zooplankton (Fig. 5.7) or a combination (Figs. 5.6a-c). We thus found different model input parameter choices leading to similar model results, which is known as equifinality.

Equifinality is important because in the literature various authors presented opposing explanations regarding the relative contribution of grazing by zooplankton and phytoplankton community characteristics (cf. salinity stress) to the mortality rate. It is not clear whether these opposing explanations are caused by real changes of the estuarine system or whether they may be attributed to equifinality. For example, on the one hand, a model study by Gypens et al. (2013) showed that salinity is a crucial driver for the spatial phytoplankton distribution in the Scheldt estuary in 2006, while grazing pressure has a negligible role. The authors thus stress the importance of salinity stress (cf. phytoplankton community characteristics) instead of grazing by zooplankton. On the other hand, a dominant impact of zooplankton grazing on the mortality rate was found by Calbet and Landry (2004) who studied 136 estuarine systems and showed that zooplankton grazes on average approximately 60% of the primary production. Moreover, by comparing Chl-a concentrations and biomass concentrations of the *Eurytemora affinis*-dominated zooplankton population in the St Lawrence estuary (Canada), Winkler et al. (2003) showed that *Eurytemora affinis* is likely to be a major cause of the downstream decrease in Chl-a concentration. This agrees with the multi-annual observations in the Scheldt estuary: in 1996, calanoid copepods, in casu *Eurytemora affinis*, dominated in the downstream brackish region and were quasi absent in the freshwater region. From 2009, they gradually developed more upstream to also become dominant there (Appeltans, 2003; Mialet et al., 2010, 2011; Chambord et al., 2016).

Although we may not further constrain the relative importance of grazing by zooplankton and phytoplankton community characteristics to the mortality rate, we can compare our model parameter values to the literature as a first verification of our model results. Our mortality rate (i.e., $m$) values $\sim 10^{-6} s^{-1}$ comply with the value of $\sim 1.1 \times 10^{-6} s^{-1}$ presented in Desmit et al. (2005) who studied a real-case in the Scheldt estuary near km 115. Additionally, incubation experiments carried out with adult *Eurytemora affinis* around km 80 in the Scheldt estuary during spring 2013 and 2014 show $q$ values between $1.54 \times 10^{-8}$ and $2.78 \times 10^{-6} s^{-1} L$ (Chambord et al., in prep.), overlapping with the modeled values in this study, but also showing large variability.

Therefore, to further constrain which multi-annual change in model parameters may have resulted in the multi-annual change in Chl-a accumulation, additional observations are required. Firstly, to determine the relative importance of grazing by zooplankton, observational studies on the mortality rate are needed. Measurements of rates (e.g., grazing rate) are better constraints than state variables (e.g., Chl-a) (Franks, 2009). For example, Friedrichs et al. (2007) showed that different model and parameter choices may fit the data equally well, although the grazing rates may differ by more than an order of magnitude. Secondly, a larger spatial and temporal resolution of the observations is recommended as different model and parameter choices may converge at the specific times
and locations for which data is present, while in between the observations the distinct models may show very distinct results (Friedrichs et al., 2007).

5.4.2 Model limitations

In this section, we present our main model limitations, compare our model approach to state-of-the-art models applied to the Scheldt estuary, and discuss its implication for other studies and future research.

Following Naithani et al. (2016), we assumed stenohaline marine and freshwater diatoms adapted to the extreme high or low salinities. However, in the literature, also euryhaline species were considered (Roubeix et al., 2008; Gypens et al., 2013). As pointed out by Gypens et al. (2013), the presence of euryhaline phytoplankton species may have a significant impact on the magnitude and distribution of both freshwater and marine phytoplankton. Additionally, in the summer of 2003, the phytoplankton community characteristics showed species with different salinity optima and rather restricted salinity tolerances (Muylaert et al., 2009). The existence of such a phytoplankton group adapted to more intermediate salinity may explain the overestimation of Chl-a by our model at ~km 21 and 36 (Fig. 5.7b).

Compared to models such as presented by Arndt et al. (2011), Gypens et al. (2013), and Naithani et al. (2016), we idealized the biochemical processes. Based on observations, we excluded nutrient limitation, only included two zooplankton groups using a data-driven approach, and only distinguished between freshwater and marine diatoms.

Although we made significant model assumptions, our model approach resulted in valuable insights. Before our work, the observed trend change in Chl-a in spring was poorly described and it was unclear whether this trend change is related to changes in physical characteristics (sediment, discharge, temperature) or changes in biological characteristics. In our contribution, we can constrain this to a change in biological characteristics related to phytoplankton mortality that seems to have some correlation with zooplankton grazing and phytoplankton community characteristics. As discussed by Oreskes et al. (1994), we used our model approach to illuminate which features of the systems mainly require further research and which empirical data is lacking.

To summarize, although a careful assessment of the model assumptions is required, our model is generally applicable to turbid nutrient-rich, tide-dominated estuaries. The approach is particularly useful to constrain parameter ranges, quantify model parameters in more advanced state-of-the-art models, and determine which empirical data is recommended for further research on this topic.
5.5 Conclusions

In this contribution, we studied the multi-annual estuary-scale evolution of the spring phytoplankton (cf. Chl-a) distribution in the Scheldt estuary. We focused on the appearance and disappearance of phytoplankton accumulation in the brackish region in spring in 2004-2018.

We first analyzed multi-annual in situ observations covering the SPM concentration, zooplankton abundance, and other variables affecting net phytoplankton growth, showing a multi-annual estuary-scale evolution of not only the SPM distribution and zooplankton abundance, but also of the freshwater discharge and maximum photosynthetic rate. Next, to detect the multi-annual evolution of these variables that can be linked to the evolution of phytoplankton, we employed a model approach in which the observations were the core. Our model allowed us to significantly constrain which evolution of variables may explain the evolution of phytoplankton; both a multi-annual change in mortality rate and corresponding grazing by zooplankton and phytoplankton community characteristics may have caused the multi-annual estuary-scale evolution of phytoplankton in spring. We were thus able to limit the number of model input parameter choices leading to similar model results, which is known as equifinality.

Although our model approach simplifies reality and shows (local) anomalies when comparing phytoplankton model results and observations, it allowed us to quantitatively determine the importance of various factors affecting phytoplankton growth on the estuary scale. This knowledge is important for moving forward using more complex numerically costly models. Our results highlight the importance of insight into the zooplankton dynamics and phytoplankton community characteristics to understand the phytoplankton dynamics in the Scheldt estuary. Further research and experimental validation are required to determine the mechanisms that may have caused these multi-annual estuary-scale changes in mortality rate, grazing, and phytoplankton community characteristics.

List of symbols

Latin

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>$B$</td>
<td>Width of the estuary</td>
</tr>
<tr>
<td>$c$</td>
<td>Suspended sediment concentration</td>
</tr>
<tr>
<td>$E$</td>
<td>Photosynthetically active radiation</td>
</tr>
<tr>
<td>$E_{00}$</td>
<td>Maximum photosynthetically active radiation</td>
</tr>
<tr>
<td>$E_1$</td>
<td>Sub-surface light intensity</td>
</tr>
<tr>
<td>$E_2$</td>
<td>Light intensity measured at a fixed distance $\Delta z$ from the sub-surface sensor</td>
</tr>
<tr>
<td>$f_S$</td>
<td>Function that determines the salinity stress of phytoplankton</td>
</tr>
<tr>
<td>$f_Z$</td>
<td>Function that determines the grazing pressure of phytoplankton</td>
</tr>
<tr>
<td>$g$</td>
<td>Grazing parameter</td>
</tr>
</tbody>
</table>
$g_1$  
Calanoids grazing parameter

$g_2$  
Non-calanoids grazing parameter

$H$  
Magnitude of the z-coordinate of the river bed

$K_h$  
Horizontal diffusivity coefficient

$K_v$  
Vertical eddy diffusivity coefficient

$k_{bg}$  
Background exponential light extinction coefficient

$k_c$  
Sediment-induced exponential light extinction coefficient

$k_d$  
Exponential light extinction coefficient

$k_P$  
Self-shading exponential light extinction coefficient

$m$  
Mortality rate of phytoplankton

$m_i^0$  
Mortality rate parameter of phytoplankton class $i$

$m_{\text{fresh.}}^0$  
Mortality rate parameter of the freshwater diatoms

$m_{\text{mar.}}^0$  
Mortality rate parameter of the marine diatoms

$P$  
Phytoplankton concentration

$P_i$  
Phytoplankton concentration of class $i$

$P_{\text{fresh.}}$  
Phytoplankton concentration of the freshwater diatoms

$P_{\text{mar.}}$  
Phytoplankton concentration of the marine diatoms

$P_{\text{max}}$  
Maximum photosynthetic rate

$P_{\text{sea}}$  
Phytoplankton boundary concentration at the mouth

$QP$  
Influx of phytoplankton at the upstream boundary

$S$  
Salinity

$s_{\text{upstream}}$  
Salinity at the upstream boundary

$s_{\text{sea}}$  
Salinity at the downstream boundary

$T$  
Water temperature

$t$  
Time

$u$  
Water flow velocity in the longitudinal dimension

$w$  
Water flow velocity in the vertical dimension

$w_P$  
Settling velocity of phytoplankton cells

$x$  
Spatial coordinate in the longitudinal dimension

$Z$  
Concentrations of phytoplankton grazers

$Z_{\text{calanoids}}$  
Calanoids concentration

$Z_{\text{non-calanoids}}$  
Non-calanoids concentration

$z$  
Spatial coordinate in the vertical dimension

Greek

\[\alpha\]  
Photosynthetic growth efficiency

$\Delta z$  
Distance between the sub-surface surface and second light sensor

$\partial_t$  
Partial derivative to $t$

$\partial_x$  
Partial derivative to $x$

$\partial_z$  
Partial derivative to $z$

$\zeta$  
$z$-coordinate of the water surface

$\mu$  
Growth rate of phytoplankton

$\mu_{00}$  
Maximum growth rate at $0 ^\circ C$

$\mu_{\text{fresh.}}^{00}$  
Maximum growth rate at $0 ^\circ C$ of the freshwater diatoms

$\mu_{\text{mar.}}^{00}$  
Maximum growth rate at $0 ^\circ C$ of the marine diatoms
Appendix 5.A: salinity profile

Following [Warner et al. (2005)] we fit the salinity data measured in spring in the Scheldt estuary to the following postulated salinity distribution:

\[
\frac{s_{\text{sea}}}{2} \left( 1 - \tanh \frac{x - x_{c}^{\text{Sal.}}}{x_{L}^{\text{Sal.}}} \right),
\]

(5.9)

where \( s_{\text{sea}} \) is the salinity boundary condition at the mouth and \( x_{c}^{\text{Sal.}} \) and \( x_{L}^{\text{Sal.}} \) are further undefined calibration parameters. Figure 5.A.1 shows the salinity data and corresponding data fit in spring for the three periods considered. The corresponding parameter values are listed in Table 5.A.1.

Figure 5.A.1: Measured salinity in the Scheldt estuary in spring and the corresponding data fit using Eq. 5.9.

Appendix 5.B: temperature dependence of \( \mu_{\text{max}} \)

Following [Eppley (1972)] we postulate the following temperature dependence of the maximum growth rate \( \mu_{\text{max}}(T) \):

\[
\mu_{\text{max}}(T) = \mu_{00} T^{\mu_{01}},
\]

(5.10)

in which \( T \) is the water temperature and \( \mu_{00} \) and \( \mu_{01} \) are calibration parameters. Figure 5.B.1a shows the multi-annual averaged temperature dependence in 2004-2018. Figure 5.B.1b shows the \( P_{\text{max}} \)-\( T \) plot for the 2015-2018 reference case and the corresponding
Figure 5.B.1: (a) The multi-annual time-averaged temperature observations between 2004-2018. (b) $P_{\text{max}}-T$ plot for the 2015-2018 reference case and the corresponding data fit using Eq. (5.10), resulting in $\mu_{00} = 1.00 \times 10^{-5}$ s$^{-1}$ and $\mu_{01} = 1.05$.

data fit following Eq. (5.10), resulting in $\mu_{00} = 1.00 \times 10^{-5}$ s$^{-1}$ and $\mu_{01} = 1.05$. Note that to translate $P_{\text{max}}$ to $\mu_{\text{max}}$, we have to divide by the C:Chla ratio. Similarly, we have $\mu_{00} = 1.21 \times 10^{-5}$ s$^{-1}$ and $\mu_{01} = 1.07$ and $\mu_{00} = 1.12 \times 10^{-5}$ s$^{-1}$ and $\mu_{01} = 1.10$ in 2008-2014 and 2004-2007, respectively.

We divide the phytoplankton abundance into freshwater and marine diatoms. Assuming that in the Belgian region of the Scheldt estuary (for which we have observations), phytoplankton consists of $\sim 75\%-25\%$ freshwater and marine diatoms, respectively, and assuming that $\mu_{00}$ of marine diatoms is $\sim 1.66$ larger [Naithani et al., 2016], we can compute the freshwater $\mu_{00}^{\text{fresh}}$ and marine $\mu_{00}^{\text{mar}}$. maximum growth rate at $0^\circ$ C:

$$\mu_{00}^{\text{fresh}} = \frac{\mu_{00}}{0.75 + 0.25 \times 1.66}$$  \hspace{0.5cm} (5.11)

$$\mu_{00}^{\text{mar}} = \frac{\mu_{00}}{1.56 + 0.25}$$  \hspace{0.5cm} (5.12)

Appendix 5.C: SPM distribution

Following [Horemans et al. (2020a)], we calibrate the erosion and flocculation characteristics by calibrating the residual SPM model output to the corresponding multi-annual residual SPM observations in 2015-2018 (Fig. 5.C.1a). The corresponding model parameters are the erosion parameter $M$ and $\lambda$, which determine the strength of the flocs. The optimal values are $\lambda = 65.9 \times 10^{-6}$ s$^{-1/2}$ m$^2$ and $M = 4 \times 10^{-3}$ s m$^{-1}$.

As reasoned by [Brouwer et al. (2018)] [Dijkstra et al. (2019b)] and [Horemans et al.]
between km 70-80 we have an increase in SPM due to dumping and dredging activities. However, this is not included in the model because the dredging and dumping activities act on a much smaller temporal scale (~ hours) and can thus be considered as a background SPM concentration, whereas our model computes the multi-annual SPM distribution. To guarantee that we capture the magnitude of the observed SPM concentrations at the dredging and dumping location, we add a background SPM concentration originated by dredging and dumping activities to our model results. More specifically, we replace the model concentration with the (smoothed) SPM observations between km 70 and 80 and beyond km 125. The resulting modeled SPM distribution is presented in Fig. 5.C.1).

To estimate the sensitivity of phytoplankton growth to the SPM concentration, we estimate the SPM distribution between 2004-2007 and 2008-2014, for which we only have water surface SPM observations. To this end, we first divide the water surface SPM concentration observed in 2004-2007 and 2008-2014 by the observations between 2015-2018. The resulting (interpolated) ratios are presented in Figs. 5.C.1a and 5.C.1b, respectively. Next, we use these ratios to scale the profiles of the modeled SPM concentrations in 2015-2018, thus keeping the shape of the vertical profiles the same. This results in the estimated modeled SPM distribution in 2004-2007 and 2008-2014 presented in Figs. 5.C.1c and 5.C.1d, respectively.

Appendix 5.D: sediment-induced light extinction coefficient

Within the OMES campaign, the exponential light extinction coefficient $k_d$ is computed by measuring the solar irradiance at the water surface $E_0$ and the light intensity at a depth $d$ of approximately 1 m. Neglecting background and self-shading light extinction (which is acceptable given the high turbidity in the Scheldt estuary), we can compute the sediment-induced light extinction $k_c$ as

$$k_d d = k_c \int_0^d c(z', t) dz', \quad (5.13)$$

in which $z'$ is the water depth and $c$ is the modeled SPM concentration (Fig. 5.C.1). Consequently, the time-averaged sediment-induced light extinction coefficient $\bar{k}_c$ can be
Figure 5.C.1: (a) The multi-annual time-averaged observations and (b) the modeled SPM concentration in spring 2015-2018. (c)-(d) Modeled SPM concentrations, which follow from scaling the vertical profiles presented in (a). The scaling is quantified by computing the ratio of the observed surface SPM concentrations of the various periods.
Figure 5.C.1: The (interpolated) water surface SPM observations divided by the observations in 2015-2018 between (a) 2004-2007 and (b) 2008-2014.

approximated by

\[
\bar{k}_c = \frac{k_c d}{\int_0^d c(z', t) \, dz'},
\]

\[
\approx \frac{k_d d}{\int_0^d c(z', t) \, dz'},
\]

(5.14)

which results in \( k_c \approx 72 \, \text{m}^2 \, \text{kg}^{-1} \) in 2015-2018 for \( d = 1 \, \text{m} \). To quantify the impact of variability in the integration depth \( d \) on the estimated value of \( k_c \), Fig. 5.D.1 shows the estimated \( k_c \) value as a function of \( d \). If we assume a vertically constant sediment concentration, Eq. (5.13) simplifies to the linear relationship \( k_d = k_c c(t) \), which results in a slightly lower \( k_c \) value of approximately 68 m\(^2\) kg\(^{-1}\). The corresponding data fit for 2015-2018 is presented in Fig. 5.D.1b. Similarly, \( k_c \) approximates 78 m\(^2\) kg\(^{-1}\) and 81 m\(^2\) kg\(^{-1}\) in 2008-2014 and 2004-2007, respectively.

Appendix 5.E: evolution in discharge, salinity intrusion, and photosynthetic parameters

Figure 5.E.1a shows the multi-annual, monthly-averaged discharge in the three periods considered. In spring (Apr.-May), the average discharge is 85, 81, and 72 m\(^{-3}\) s\(^{-1}\) in 2004-2007, 2008-2014, and 2015-2018, respectively. We thus observe a slight decrease in total freshwater discharge over time.
Figure 5.D.1: (a) Sediment-induced light extinction coefficient assuming a vertically non-constant SPM profile. Because the light extinction coefficient $k_c$ is measured at approximately 1 m depth, we use the corresponding $k_c$ value of 72 m$^2$ kg$^{-1}$ in our model (depicted in blue). (b) Estimation of $k_c$ in 2015-2018 assuming a vertically constant suspended sediment profile. By doing so, $k_c$ equals the slope of the linear relationship between $k_d$ and SPM (depicted by the blue line).

Figure 5.E.1b presents the monthly-averaged salinity intrusion, which is defined as the distance from the mouth at which the salinity equals 2 ppt. We typically observe the opposite trend compared to the freshwater discharge: a larger discharge comes with a smaller salt intrusion. However, this is not always valid. For example, in March, we have a similar time-averaged discharge in the three periods considered, while the salinity intrusion is significantly different. What is most important within the scope of this paper is the fact that in spring (Apr.-May) the salinity intrusion does not show major changes during the study period 2004-2018.

Figure 5.E.2a displays the monthly- and system-averaged maximum photosynthetic rate $P_{\text{max}}$ in the three periods considered. In spring (Apr.-May), $P_{\text{max}}$ is approximately equal in 2004-2007 and 2008-2014, but significantly lower in 2015-2018. The corresponding time-averaged values are 6.59, 6.44, and 4.31 mg C (mg Chl-a)$^{-1}$ h$^{-1}$, respectively.

Figure 5.E.2b shows the monthly- and system-averaged photosynthetic efficiency $\alpha$ in 2004-2018. What is most important within the scope of this paper is that $\alpha$ is not significantly different in all three periods 2004-2007, 2008-2014, and 2015-2018 in spring. The corresponding time-averaged values are 0.0165, 0.0168, and 0.0188 mg C (mg Chl-a)$^{-1}$ h$^{-1}$ [PAR]$^{-1}$, respectively.
Figure 5.E.1: Multi-annual monthly-averaged observations in 2004-2018 of (a) the total freshwater discharge (at the upstream boundary and tributaries) and (b) the salinity intrusion defined as the distance at which the salinity equals 2 ppt. The error bars depict the standard error of the observations.

Figure 5.E.2: Multi-annual monthly-averaged observations in 2004-2018 of the photosynthetic parameters (a) $P_{\text{max}}$ and (b) $\alpha$. The error bars depict the standard error of the observations. With ‘[PAR]’, we denote ‘the units of PAR’, i.e. $\mu$mol photons m$^{-2}$ s$^{-1}$.
Appendix 5.F: sensitivity study of model parameters

Figures 5.F.1a-d show the impact of variations in the sediment-induced light extinction coefficient $k_c$ and the maximum growth rate $\mu_{00}$ at 0 °C on the Chl-a model result for both the freshwater and marine diatoms in 2008-2014, keeping other parameters at their default values. The parameters $m_{0i}^{\text{max}}$, $g_1$, and $g_2$ are assumed to be fixed and set to the values corresponding to 2004-2007 and 2015-2018. We have to decrease $k_c$ by a factor of approximately 3 to obtain accumulation of Chl-a in the brackish region. This difference is significantly larger than the variability that follows from the observations, which is between approximately 65 and 80 m$^2$ kg$^{-1}$ (see Section 5.D). Similarly, at large $\mu_{00}$, we do not detect accumulation of Chl-a in the brackish region. Changes in the freshwater discharge $Q$, the influx of Chl-a at the upstream boundary $QP$, and the Chl-a concentration at the seaside boundary $P_{\text{sea}}$ in 2008-2014 only lead to changes in the upstream or downstream Chl-a concentration, but do not result in the accumulation of Chl-a in the brackish region (Figs. 5.F.2a-d). Therefore, changes in these parameters alone cannot explain the disappearance of accumulation of Chl-a in the brackish region.
Figure 5.F.1: Sensitivity study of the depth-averaged Chl-a concentration in 2008-2014 to the (a)-(b) sediment-induced light extinction coefficient $k_c$ and (c)-(d) the maximum growth rate $\mu_0$ at $0 \, ^\circ\mathrm{C}$ for both freshwater and marine diatoms. The default parameter values are depicted by the dashed horizontal lines.
Figure 5.F.2: Sensitivity study of the depth-averaged Chl-a concentration in 2008-2014 to the (a)-(b) the total freshwater discharge $Q$, (c) the Chl-a concentration at the seaside boundary $P_{\text{sea}}$, and (d) the phytoplankton influx at the upstream boundary $QP$. The default parameter values are depicted by the dashed horizontal lines.
<table>
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<td>$A^0$</td>
<td>Mz water level amplitude at x=0</td>
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<td>$A^1$</td>
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<td>$\phi^1$</td>
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<td>-1.3</td>
<td>-1.3</td>
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<td>Total river discharge to resolve the phytoplankton dynamics</td>
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<td>81</td>
<td>72</td>
<td>m$^3$s$^{-1}$</td>
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<td>$Q_{USS}$</td>
<td>Relative discharge contribution of the Upper Sea Scheldt boundary</td>
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<td>31.5%</td>
<td>27.1%</td>
<td>/</td>
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<tr>
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<td>Relative discharge contribution of the Rupel tributary</td>
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<td>59.3%</td>
<td>63.3%</td>
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<td>$Q_{Dender}$</td>
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<td>9.2%</td>
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<td>$Q_{ssea}$</td>
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<td>$c_{sea}$</td>
<td>Depth-averaged subtidal concentration at x=0</td>
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<td>Ratio of minimal aggregation and floc break-up parameter in spring</td>
<td>$65.9 \times 10^{-6}$</td>
<td>/</td>
<td>/</td>
<td>s$^{-1/2}$ m$^{2}$</td>
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<td>$\rho_{N}$</td>
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<td>mmol N L$^{-1}$</td>
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<td>$Phos_{sea}$</td>
<td>Phosphorus boundary concentration at the mouth</td>
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<td>Influx of nitrogen at the upstream boundary</td>
<td>3</td>
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<td>Influx of phosphorus at the upstream boundary</td>
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<td>/</td>
<td>mol P s$^{-1}$</td>
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<td>Influx of nitrogen at Rupel tributary</td>
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<td>/</td>
<td>mol N s$^{-1}$</td>
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<td>$Phos_{Rup}$</td>
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<td>$C:NP$</td>
<td>Ratio between carbon, nitrogen, and phosphorus (in mol)</td>
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<td>/</td>
<td>/</td>
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<td>$\mu_{max}$</td>
<td>Maximum growth rate at 0 °C of to freshwater diatoms (source)</td>
<td>$0.96 \times 10^{-5}$</td>
<td>$1.04 \times 10^{-5}$</td>
<td>$0.86 \times 10^{-5}$</td>
<td>s$^{-1}$</td>
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<tr>
<td>$\mu_{max}$</td>
<td>Maximum growth rate at 0 °C of to marine diatoms (source)</td>
<td>$1.59 \times 10^{-5}$</td>
<td>$1.72 \times 10^{-5}$</td>
<td>$1.43 \times 10^{-5}$</td>
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<td>Calibration parameter in temperature dependence for $\mu_{max}$</td>
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<td>1.07</td>
<td>1.05</td>
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<td>$0.8 \times 10^{-7}$</td>
<td>$0.13 \times 10^{-6}$</td>
<td>$0.8 \times 10^{-7}$</td>
<td>s$^{-1}$ L</td>
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<td>$g_2$</td>
<td>Non-calanoids grazing parameter (calibrated)</td>
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<td>$0.32 \times 10^{-6}$</td>
<td>$0.47 \times 10^{-7}$</td>
<td>s$^{-1}$ L</td>
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<tr>
<td>$m_{cal}$</td>
<td>Mortality rate parameter for freshwater diatoms calibrated</td>
<td>$1.89 \times 10^{-6}$</td>
<td>$3.30 \times 10^{-6}$</td>
<td>$2.90 \times 10^{-6}$</td>
<td>s$^{-1}$</td>
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<td>Mortality rate parameter for marine diatoms (calibrated)</td>
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<td>s$^{-1}$</td>
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<td>/</td>
<td>/</td>
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<td>14.7</td>
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<td>Marine phytoplankton boundary concentration at the mouth</td>
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<td>17.1</td>
<td>15.8</td>
<td>µg L$^{-1}$</td>
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<td>Influx of freshwater phytoplankton at the upstream boundary (source)</td>
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<td>1.8</td>
<td>2.5</td>
<td>g s$^{-1}$</td>
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<td>$w_{P}$</td>
<td>Setting velocity of phytoplankton cells (Sarthou et al., 2005)</td>
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<td>/</td>
<td>m s$^{-1}$</td>
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<tr>
<td>$k_{bg}$</td>
<td>Background exponential light extinction coefficient (Pennock and Sharp, 1994)</td>
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<td>Sediment-induced exponential light extinction coefficient (Pennock and Sharp, 1994)</td>
<td>81.4</td>
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<td>72.0</td>
<td>m$^{-2}$ kg$^{-1}$</td>
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<td>$\alpha$</td>
<td>Growth efficiency (source)</td>
<td>0.0165</td>
<td>0.0168</td>
<td>0.0188</td>
<td>[µmol photons m$^{-2}$ s$^{-1}$]$^{-1}$</td>
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Bibliography


In this chapter, we first conclude this thesis by answering the four main research questions we presented in Chapter 1. Next, we suggest further research opportunities that would naturally follow from our results.

### 6.1 Conclusions for our research questions

In this thesis, we studied the interconnection of SPM and phytoplankton on the large temporal- and spatial scale. We focused on a turbid, tide-dominated, and nutrient-rich estuary, being the Scheldt estuary. We combined the analysis of long-term observations covering the entire domain of the Scheldt estuary and a modeling approach. In Fig. 6.1, we repeat the outline of this thesis and present brief answers to our four main research questions corresponding to each chapter.

- **Chapter 5**: The observed disappearance of Chl-a accumulation in the brackish region in spring is due to a change in mortality rate.
- **Chapter 4**: Temporal variability in light climate may have a significant impact on phytoplankton growth.
- **Chapter 2**: Flocculation may have a large-scale impact on the SPM distribution.
- **Chapter 3**: Seasonal variations in biota are not necessary to explain seasonal variations in sediment dynamics.

Figure 6.1: Illustration of the interconnection between SPM and phytoplankton studied in this thesis, the corresponding chapters, and a brief answer to the four main research questions. SPM limits phytoplankton growth by deteriorating the light climate in the water column. Phytoplankton may affect SPM dynamics by altering the floc size and structure of flocs and excreting sticky substances that stimulate flocculation of SPM. In Chapter 2, we studied the impact of flocculation on SPM. In Chapter 3, we focused on the importance of biotically-induced flocculation on SPM. In Chapter 4, we analyzed the impact of temporal variability in light climate (cf. SPM) on phytoplankton. We combined all results and studied a specific case in the Scheldt estuary in Chapter 5.

- What is the impact of flocculation on the large-scale SPM dynamics?

In Chapter 2, we determined whether the small-scale flocculation process may impact the
estuary-scale formation of estuarine turbidity maxima (ETM). To this end, we extended the idealized, width-averaged hydro-sediment transport model known as iFlow by a flocculation model and applied the model to the Scheldt estuary in winter conditions. We showed that spatial and temporal variations in settling velocity caused by flocculation are necessary to capture the observed magnitude of the SPM concentrations. More specifically, a mechanism caused by the temporal interaction between the longitudinal water flow velocity and settling velocity prevents flushing out of the ETM at large river discharges. Additionally, it promotes land-inward sediment transport and may cause the formation of additional ETM. It may also promote downstream net sediment transport, which depends on the phase between the longitudinal water flow velocity and the M2 tidal signal in the settling velocity and can thus differ for various estuaries. To conclude, the small-scale flocculation process may have an estuary-scale impact on the SPM distribution.

- Can we detect a seasonal biotic impact on the SPM dynamics through flocculation and erosion?

In Chapter 3 we applied the modeling framework from Chapter 2 to study the importance of biotically-induced seasonality in flocculation and erosion on seasonality in ETM formation. Here, we again focused on the Scheldt estuary. By calibrating the biotically-affected model parameters to long-term, in situ turbidity observations, we showed that we do not require seasonality in biotically-affected flocculation and erosion parameters to reproduce the seasonality in ETM formation. The mechanism resulting in the seasonality in ETM formation is the seasonality in freshwater discharge; the higher discharges in winter flush out the upstream ETM observed in summer. Additionally, they increase the shear rate in the upstream region, which promotes break-up and thereby decreases the land-inward sediment transport caused by flocculation. Our conclusion was supported by long-term, in situ observations of turbidity, floc size, Chl-a, and sticky substances excreted by phytoplankton; the abiotic observations show seasonality, while seasonality in the sticky substances is absent on the estuary and seasonal scale. To conclude, we cannot detect an important seasonal biotic impact through flocculation and erosion on the seasonal estuary-scale SPM dynamics.

- Can we construct a model to easily assess the relative impact of temporal fluctuations in SPM on phytoplankton growth?

In Chapter 4 before using our results from Chapters 2 and 3 to construct a phytoplankton-SPM model, we developed a tool to locally study the impact of temporal variability in light climate on primary production without solving the complex SPM dynamics explicitly. The tool allowed us to determine the relative impact of temporal variability in the SPM dynamics without a priori assumptions regarding these dynamics. We aimed to know whether we may neglect temporal variations in these parameters and, if not, how we can correct for the introduced error by excluding this temporal variability. We focused on three factors affecting phytoplankton growth through light availability: solar irradiance at the water surface, exponential light extinction in the water column, and water depth.
(cf. ratio between euphotic/mixing depth). We again applied the tool to the Scheldt estuary. More specifically, we studied a phytoplankton bloom in spring and gross primary production in equilibrium conditions in summer and winter. We showed that our conceptual model correctly predicts the magnitude and sign of the introduced error. Moreover, we revealed that variability in solar irradiance has the largest impact on time-averaged primary production (PP) in dynamic equilibrium, resulting in a 30% decrease compared to time-invariant PP. Caused by this decrease, temporal fluctuations in solar irradiance may also result in a significant decrease of unbounded exponential phytoplankton growth and may delay the onset of a phytoplankton bloom by \( \sim 2 \) weeks. Our case study showed that temporal variability in light extinction coefficient and water depth has a less significant influence on phytoplankton growth. The reason is that temporal variability is relatively small compared to its time average for these two parameters. We showed that the effect of temporal fluctuations in light extinction is more extensive in winter than in summer because of the larger temporal variations in winter. The relative importance of temporal variability in the parameters is thus season- and system-dependent. To conclude, we constructed a conceptual model that allowed us to easily assess the relative impact of temporal fluctuations in SPM on phytoplankton growth. The tool can be easily implemented in models that explicitly resolve the SPM-phytoplankton dynamics, as done in Chapter [5].

- Can we link the disappearance of the spring-bloom in the brackish region in the Scheldt estuary between 2004-2018 to multi-annual changes of processes impacting phytoplankton growth and their corresponding model parameters?

In Chapter [5] we combined our previous results to construct a phytoplankton-sediment transport model. To show its applicability, we applied the model to investigate the multi-annual evolution of the estuary-scale phytoplankton distribution in spring in the brackish region in the Scheldt estuary. More specifically, we determined what change in factors affecting phytoplankton growth may have dominantly contributed to the appearance and disappearance of a spring phytoplankton bloom in the Scheldt estuary between 2004-2018. To this end, we combined field observations and a modeling approach in which we applied an extensive sensitivity analysis and focused on the effects of individual model parameters. Our results suggest that the observed change in SPM alone cannot explain the multi-annual evolution in phytoplankton blooms in spring. Instead, a multi-annual change in phytoplankton mortality rate, grazing by zooplankton, and phytoplankton community characteristics may result in this multi-annual estuary-scale evolution of phytoplankton abundance. On the one hand, the combined effect of a land-inward shift of the zooplankton distribution and a change in grazing characteristics may explain the occurrence of a spring bloom of phytoplankton in the brackish region in 2008-2014. On the other hand, we can attribute this temporary spring bloom to multi-annual alterations in the phytoplankton community characteristics and the resulting different mortality caused by, for example, salinity stress. We thus found different model input parameter choices leading to similar model results, which is known as equifinality. Additional observations of grazing and mortality rates for various phytoplankton taxa in the Scheldt estuary are required to constrain further which multi-annual change in model parameters may have resulted in the multi-annual change in phytoplankton accumulation.
6.2 General conclusion

We combined a model approach and analysis of long-term observations in the Scheldt estuary to study the coupled SPM-phytoplankton dynamics on the large spatial and temporal scale. First, we extended a hydro-sediment transport model by a flocculation model and showed that flocculation may have a drastic impact on the estuary-scale SPM distribution. Next, we applied the model to study the impact of biotically-induced flocculation on the estuary-scale seasonality in the SPM distribution. Here, we showed that biotically-induced seasonality in flocculation and erosion only has a minor impact on the seasonality in SPM on the estuary scale; the observed seasonality in SPM can be explained through seasonality in freshwater discharge. Next, before constructing a model that explicitly resolves the phytoplankton-SPM dynamics, we constructed a conceptual model to assess the relative impact of temporal variability in light climate (cf. SPM) on phytoplankton growth. We found that temporal fluctuations in light climate may reduce time-averaged PP and unbounded exponential phytoplankton growth and delay the onset of a spring bloom by $\sim 2$ weeks. Finally, we implemented this conceptual model and our previous results to construct a phytoplankton-sediment transport model. We applied the model to determine what processes may explain the multi-annual appearance and disappearance of phytoplankton accumulation in the brackish region in spring in the Scheldt estuary in 2004-2018. We showed that a change in mortality rate may explain this multi-annual trend. Additionally, we showed that the change in mortality rate may be caused by changes in grazing by zooplankton or phytoplankton community characteristics. Although we used our framework to study the Scheldt estuary, it may also be applied to other turbid, tide-dominated, and nutrient-rich estuaries.

6.3 Implication for the Scheldt estuary

In this section, we list some examples of potential implications of our results for the Scheldt estuary.

6.3.1 Management

In Chapter 4, we showed that temporal variability in water depth might increase phytoplankton growth. Intertidal areas are regions in which the relative contribution of temporal variability in water depth is expected to be significant as they are shallow and tidally-rewetted. Therefore, intertidal areas may positively affect phytoplankton growth based on our results. This may be important for managing the estuary; expanding the Scheldt estuary by reclaiming and reflooding land may result in an overall increase of phytoplankton growth, which is a crucial indicator for the ecological state of the estuary set by the European Water Framework Directive.
6.3.2 Holistic approach

We illustrated the potential importance of bio-physical-chemical interactions when studying SPM dynamics. More specifically, we showed that flocculation, which may be impacted by biochemical activity, could significantly affect ETM formation in Chapter 2. It may result in additional ETM and prevent flushing of the ETM at large freshwater discharges. These conclusions may be relevant for the Scheldt estuary because Cox et al. (2019) reported an overall increase in SPM concentrations over the last two decades. Although we showed that biotically-induced flocculation and erosion have a limited impact on the seasonality in ETM formation, this may not be true when focusing on multi-annual changes in the SPM distribution. Differently put, it may be that the reported overall increase in SPM in the Scheldt estuary over the last two decades is caused by multi-annual changes in the flocculation characteristics. This shows the importance of a holistic and interdisciplinary approach when studying multi-annual shifts in the SPM distribution.

6.3.3 Monitoring

In this thesis, we demonstrated that combining field observations (cf. monitoring) and modeling may result in valuable insights into the functioning of the SPM and phytoplankton dynamics. Monitoring programs may thus deliver important modeling insights. Conversely, model results may also impact and optimize monitoring programs. For example, we constrained the multi-annual appearance and disappearance of phytoplankton blooms in spring in the brackish region in the Scheldt estuary to a limited number of calibration parameters in Chapter 5. These modeling insights may adjust monitoring programs by focusing on these constrained parameters in the future. The coherence between monitoring and modeling insights is crucial when studying a dynamic and complex system as the Scheldt estuary, which is continuously changing.

6.4 Opportunities for further research

In this section, we suggest ideas for further research that naturally follow from our results and would extend the research scope of this thesis.

6.4.1 Application to other estuarine systems

Given the complexity of estuarine SPM dynamics, their dominant processes may be highly system-dependent. Therefore, various authors applied an intercomparison study of different estuaries to understand their complex dynamics. In this thesis, we focused on the Scheldt estuary. However, the iFlow model without flocculation has also been applied to the Ems estuary (Dijkstra et al., 2019b), Delaware river estuary (Dijkstra et al., 2019a),
and the Garonne tidal river (Jalón-Rojas et al., 2021). To study the individual impact of floculation, our model can thus be employed to these estuaries and other estuaries with similar characteristics, such as the Seine, Gironde, and Loire estuary. By doing so, we may reach different conclusions than found for the Scheldt estuary in Chapter 3; we may discover systems in which the aggregated effect of biotically-induced floculation and erosion may have a significant impact on the seasonality of the estuary-scale SPM distribution or in which floculation promotes net SPM export, which we calculated to be possible in Chapter 2.

6.4.2 Alternative model applications

Numerical models are a form of complex scientific hypotheses as they come with a list of assumptions (Oreskes et al., 1994). As explained by Franks (2009), the choice of an appropriate modeling approach depends on the research questions and data availability. Alternative model applications with different model assumptions are helpful to validate the model or hypothesis or answer additional research questions. For example, in an earlier iFlow model version, the settling velocity was considered constant. As we were interested in the effects of floculation on the SPM dynamics and measurements of floc characteristics that have recently been included in the OMES monitoring program, we considered floculation in the model. This resulted in a spatially and temporally varying settling velocity and valuable insight into the relative impact of floculation on ETM formation on the estuary scale (Chapters 2 and 3). In future research, alternative model extensions that come with different assumptions can be applied to, for example, capture local processes such as the effects of intertidal areas on the SPM dynamics.

6.4.3 The link between TEP, stickiness, and Chl-a

The link between TEP, stickiness, and Chl-a is ambiguous; various estuaries and seasons show different correlations. The reason is that, firstly, the TEP concentrations are often estimated using the colorimetric method described in Claquin et al. (2008). A disadvantage of this method is that TEP is not well-defined from a chemical-physical perspective because it has various sub-fractions and components. Consequently, translating TEP to stickiness (i.e., a physical and chemical characteristic) may be very challenging.

Secondly, as opposed to some results found in the literature, a clear link between Chl-a and TEP was absent in the Scheldt estuary. We may attribute this discrepancy to the complex interplay between various bio-physical-chemical processes that result in TEP production. Various authors linked TEP concentrations to environmental conditions such as SPM concentration (Malpezzi et al., 2013), salinity (Alldredge et al., 1993, Bar-Zeey et al., 2015), turbulence (Beauvais et al., 2006), and nutrient depletion (Corzo et al., 2000, Passow, 2002). Additionally, the TEP concentration also depends on phytoplankton age and mortality (Liu and Buskey, 2000, Ramaiah et al., 2001, Morelle et al., 2017), and the community composition and phytoplankton and bacteria species interactions (Alldredge...
Therefore, additional laboratory experiments may be set up in future research to study the impact of various phytoplankton taxa and environmental conditions on TEP production and its corresponding stickiness.

6.4.4 Refinement of the boundary conditions

We showed that the boundary conditions may significantly impact the phytoplankton and SPM concentrations. For example, an increased phytoplankton concentration at the up- and downstream boundary resulted in higher phytoplankton concentrations further away from its boundaries (Chapter 5). We also concluded that changes in phytoplankton community characteristics, which depend on the boundary conditions because marine and freshwater diatoms are imported from the down- and upstream boundaries, may have resulted in the multi-annual appearance and disappearance of the phytoplankton bloom in spring in the brackish region. This illustrates the importance of boundary conditions, which are in the current model version data-driven and not explicitly resolved. In future research, it may thus be interesting to refine the boundary conditions by, for example, extending our framework with a model that resolves the phytoplankton dynamics in the coastal region. This may result in valuable insights into the coastal-estuarine coupling.

6.4.5 Explicitly resolve the zooplankton dynamics

Various zooplankton-phytoplankton models have been applied in the literature, which showed that specific parameter choices may result in bifurcations and limit cycles (Steele and Henderson, 1992). Therefore, the zooplankton-phytoplankton coupling may also result in the appearance and disappearance of phytoplankton abundance, which we studied in Chapter 5. Because we aimed to eliminate the uncertainty of a dynamic zooplankton model and the temporal resolution of our field observations was relatively low (biweekly-monthly sampling), we used the observations of zooplankton concentrations directly in the model. We implemented the effect of grazing of phytoplankton by zooplankton parametrically, assuming equilibrium conditions. In future research, zooplankton-phytoplankton dynamics may be resolved explicitly. This may provide valuable insights into the dynamics on a smaller temporal scale (~ days-weeks), parameter stability, bifurcations, and limit cycles caused by the zooplankton-phytoplankton coupling.

6.4.6 Experimental validation of the calibrated model parameters

In Chapter 5 we concluded that the multi-annual appearance and disappearance of phytoplankton growth in the brackish region in spring in the Scheldt estuary may be caused by a change in both grazing by zooplankton and phytoplankton community characteristics. Also, in Chapter 3 we showed that both changes in primary particle density and floc
aggregation and break-up characteristics may result in a similar land-inward shift of the ETM. We thus found different model input parameter choices leading to similar model results, which is known as equifinality. To further constrain these parameters, additional experimental data is required. Therefore, given the potential importance of these parameters, our results can inspire and motivate experimental scientists to validate our model parameter values.
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Model and data availability

The iFlow code is open source and licensed under LGPL (GNU Lesser General Public License), which means that it may be used freely for non-commercial and commercial purposes. The version 2.9 code with the flocculation extension and an input file example are available through https://doi.org/10.5281/zenodo.4560637.

The data presented in this thesis in the Belgian and Dutch region of the Scheldt estuary is third-party data and was accessed through http://www.omes-monitoring.be/en/data and https://waterinfo.rws.nl/, respectively.


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List of publications

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Reports


Software

• Expansion of the width-averaged, hydrodynamics and sediment transport model called iFlow to include flocculation processes (2020): https://github.com/YoeriDijkstra/iFlow

• Model to analyze the impact of temporal variability in light availability on primary production (2020): https://ars.els-cdn.com/content/image/1-s2.0-S0304380020303574-mmcl.zip