

Wind and Phytoplankton Dynamics Drive Seasonal and Short-Term Variability of Suspended Matter in a Tidal Basin

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Abstract

Suspended particulate matter (SPM) is a key component of coastal ecosystems, modulating light availability, nutrient transport, and food web dynamics. Its variability is driven by a combination of physical and biological processes that interact across temporal and spatial scales. Using the Sylt-Rømø Bight as a natural laboratory and focusing on the period 2000-2019, in this study, we integrate statistical analysis of observational data from the Sylt Roads monitoring program and local meteorological stations, neural network modelling and Lagrangian transport simulations. This multi-method approach enables us to disentangle and quantify the relative roles of tidal and wind forcing, as well as biological processes in shaping SPM concentrations across various time scales, based on near-surface measurements at two monitoring stations.

The findings show that wind intensity dominates short-term SPM variability, particularly at the shallow station, where SPM responds rapidly to local wind-induced resuspension. At the deep station, the wind effects appear with a delay of ~5 days, aligning with tidally induced transport timescales (~133 hours) from shallower resuspension zones, as revealed by Lagrangian simulations. Seasonal patterns are further modulated by both reduced wind intensities and the onset of biological processes, such as phytoplankton blooms, which promote flocculation and subsequent settling in spring and summer. Neural network experiments highlight the shifting seasonal balance between physical and biological controls. The median concentration of SPM decreased by up to 80% from winter to summer. Approximately 40% of this seasonal difference can be attributed to weaker wind conditions, while the remaining ~40% is likely driven by biologically mediated sinking processes.

36 1 Introduction

37 Suspended particulate matter (SPM) is a key component of coastal systems, influencing a wide range of
38 physical and ecological processes. It consists of a mixture of small solid particles of both organic and
39 inorganic origin suspended in the water column with concentrations, size (Eisma, 1986) and
40 composition varying spatially and temporally (Schartau et al., 2019). The spatiotemporal variation in
41 SPM concentrations is driven by an interplay of hydrodynamic, meteorological, and biological factors,
42 which in turn, regulate nutrient availability, light penetration, and organic matter distribution. This
43 directly impacts ecosystem productivity, including the timing of the phytoplankton bloom in spring
44 (Cadée, 1986) and spatial gradients in primary productivity (Cloern, 1987; Colijn, 1982), and trophic
45 interactions (Dolch and Reise, 2010; Graf and Rosenberg, 1997).
46 In tidally energetic environments like the Wadden Sea, the suspended matter gradient is kept upright by
47 the density-driven coastward transport of bottom water and tidal straining (Becherer et al., 2016;
48 Burchard et al., 2008; Flöser et al., 2011). Sediment accumulation is further influenced by
49 hydrodynamic retention mechanisms such as enhanced settling due to the landward dissipation of
50 current velocity, the scour-lag (Dyer, 1995; Friedrichs and Aubrey, 1988) and settling lag effects
51 (Postma, 1967), and the tidal asymmetry formed by the presence of non-linear processes within the tidal
52 system (Dronkers, 1986; Fofonova et al., 2019; Friedrichs and Aubrey, 1988; Hagen et al., 2022). The
53 role of tidal currents in the variability of SPM concentrations is found to be significant through transport
54 processes (Bartholomä et al., 2009; Christiansen et al., 2006) as well as sediment erosion even under
55 calm and moderate weather conditions (Bartholomä et al., 2009; Lettmann et al., 2009). While tides
56 regulate advection and maintain a baseline shear stress, the co-occurrence of strong winds and wave-
57 induced turbulence during severe weather further enhances bottom stress and promotes additional
58 erosion.
59 Wind stress and wave action are critical in short-term SPM dynamics. Wind-induced resuspension,
60 particularly in shallow coastal areas, causes episodic increases in turbidity (Aarup, 2002; Fettweis et al.,
61 2012). Stronger and more persistent wind forcing during winter maintains higher SPM concentrations,
62 keeping fine sediments in suspension (de Jonge and van Beusekom, 1995; van Beusekom et al., 1999),
63 while calmer conditions in summer enable enhanced settling (Bale et al., 1985; Verney et al., 2009).
64 Beyond physical resuspension and transport mechanisms, biochemical processes also influence SPM
65 concentrations by modulating aggregation, stabilization, and vertical flux of particulate matter (de Jonge
66 and van Beusekom, 1995; van Beusekom and de Jonge, 2002; Verney et al., 2009). For example,
67 flocculation, the aggregation and breakup of particles, is a key mechanism by which biological activity
68 modulates SPM concentrations (Wotton, 2004; Eisma, 1986). Phytoplankton blooms in spring and
69 summer can promote flocculation, leading to enhanced particle settling and reduced SPM
70 concentrations in the water column (de Jonge & van Beusekom, 1995; Schartau et al., 2019). The
71 process is particularly enhanced by the occurrence of extracellular polymeric substances (EPS),
72 including marine gels such as transparent exopolymeric particles (TEPs), but can also occur due to
73 cohesive properties of fine-grained minerals like clays (Passow, 2002; Verney et al., 2009). The size
74 and cohesiveness of these biologically mediated flocs also govern their settling velocities and
75 resuspension thresholds, affecting how quickly particles are redistributed in the water column. Warmer
76 temperatures also accelerate the decomposition of organic matter (OM), which is nearly always present

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in natural flocs in various forms (detritus, adsorbed OM molecules, and living OM), and may range from a minor to a dominant fraction of total floc mass (Eisma, 1986; Engel and Schartau, 1999). This directly links SPM dynamics to food-web functioning (Wotton, 2004; Engel & Schartau, 1999). In addition, temperature also affects top-down controls such as zooplankton grazing, which alters SPM composition [and concentration through multiple pathways. Grazing modifies the SPM field not only through phytoplankton consumption but also via the fragmentation of aggregates, production of fecal pellets and organic material that promote microbial and TEP-mediated aggregation. The effects of these processes can either promote particle settling or enhance recycling and suspension, depending on community composition, feeding modes, and environmental conditions \(e.g., Passow, 2002; Toullec et al., 2019; Turner, 2002\).](#)

In the shallow coastal systems, light can reach the seafloor, stimulating benthic algae growth, provided water clarity allows sufficient light penetration (Loebl et al., 2007). These benthic processes further influence SPM concentrations through both stabilization and removal mechanisms. While microphytobenthos, consisting of benthic diatoms and cyanobacteria, produce biofilms that stabilize sediments and reduce resuspension (Stal, 2010), filter-feeding organisms, such as mussels (*Mytilus edulis*) and oysters (*Magallana gigas*), alter SPM dynamics by removing fine particles from suspension, affecting both sediment deposition rates and nutrient cycling (Graf and Rosenberg, 1997). Moreover, excessive nutrient loads can enhance phytoplankton blooms, whose decay products and extracellular polymeric substances facilitate biomineral floc formation (Passow, 2002; van Beusekom et al., 1999), which promotes OM sedimentation and alters benthic-pelagic coupling by influencing the availability of organic matter to both suspension and deposit feeders. These processes collectively shape coastal sediment dynamics, contributing to the long-term evolution of tidal flat environments.

While hydrodynamic and wind-driven influences on SPM have been extensively studied, their interaction with biological processes and relative contributions remain incompletely understood, despite growing interest. By leveraging long-term ecological monitoring data from the Sylt Roads program, this study aims to quantify the contributions of tidally induced and wind-driven resuspension and transport, as well as biologically mediated processes, to the spatiotemporal variability of SPM in the Sylt-Rømø Bight.

This analysis integrates high-resolution in situ measurements from two [Long-Term Ecological Research \(LTER\)](#) stations from 2000 to 2019, meteorological data from the station List (Sylt, Germany), and outputs from hydrodynamic and neural network modelling to evaluate SPM dynamics across short-term (hourly to daily) and seasonal timescales. The following research questions guide the study: (1) What are the dominant mechanisms driving SPM variability across different temporal scales? (2) How do these mechanisms differ between the two monitoring stations within the Sylt-Rømø Bight? (3) How important are biological processes in shaping observed SPM concentrations?

Deleted: by consuming phytoplankton and restructuring organic aggregates.

115 **2 Data and Methods**

116 **2.1 Area Description**

117 The investigations were carried out in the Sylt-Rømø Bight, a shallow, tidal basin in the northern
118 Wadden Sea (southeastern North Sea; see Fig. 1). The basin is semi-enclosed due to two causeways at
119 its northern and southern ends, isolating it from neighboring basins. Its only connection to the North Sea
120 is through the deep tidal inlet Lister Deep, between the islands of Sylt and Rømø. The bay spans
121 approximately 410 km² and features a highly variable topography, including extensive intertidal flats
122 (>45%), shallow subtidal zones (~35%), and deep tidal channels (~10%). The Sylt-Rømø Bight's
123 bathymetry is characterised by a mean water depth of approximately 4 m, with a maximum depth of
124 about 37 m observed in the tidal inlet Lister Deep. Most subtidal and ~72% of intertidal sediments are
125 sand-dominated. The basin also features a small sheltered embayment Königshafen, with an average
126 depth of ~2m and [encompasses large areas which become](#) exposed at low tides.
127 The tidal range in the bight averages 2 m, based on observations at the List tide gauge (E.U. Copernicus
128 Marine Service Information, doi.org/10.48670/moi-00036). Tidal forcing accounts for over 80% of
129 depth-averaged velocity variability under regular wind conditions, in the absence of storms, and over
130 90% during spring tides (Fofonova et al., 2019). The bight receives minimal fluvial input, with small
131 rivers such as the Vidå and Brede Å (see Fig.1), contributing only 4–10 m³/s of freshwater (Purkiani et
132 al., 2015). Water exchange with the open North Sea occurs exclusively through Lister Deep, a 2.8 km
133 wide tidal inlet. At the mouth of Lister Deep, a prominent ebb-tidal delta extends seaward, acting as
134 both a sediment trap and a pathway for sediment redistribution within the bight (Dissanayake et al.,
135 2012). Localized anthropogenic disturbance may also influence sediment availability and distribution in
136 the region, although detailed records of maritime traffic, dredging operations, and benthic trawling
137 specific to the Sylt-Rømø Bight are limited. However, studies from the broader Wadden Sea show that
138 anthropogenic activities such as bottom trawling (e.g., Bruns et al., 2023; Depestele et al., 2016),
139 dredging (de Jonge and de Jong, 2002; de Jonge, 1983; van Maren et al., 2015), and mussel farming
140 (Jansen et al., 2023) are recognized drivers of sediment disturbance and resuspension.

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Sylt-Rømø Bight

Legend

- tidal flats
- bivalve beds
- seagrass
- salt marshes

- Water depth
- >1-2 m
 - >2-5 m
 - >5-10 m
 - >10-20 m
 - >20 m

- tide gauge stations
- LTER sampling stations

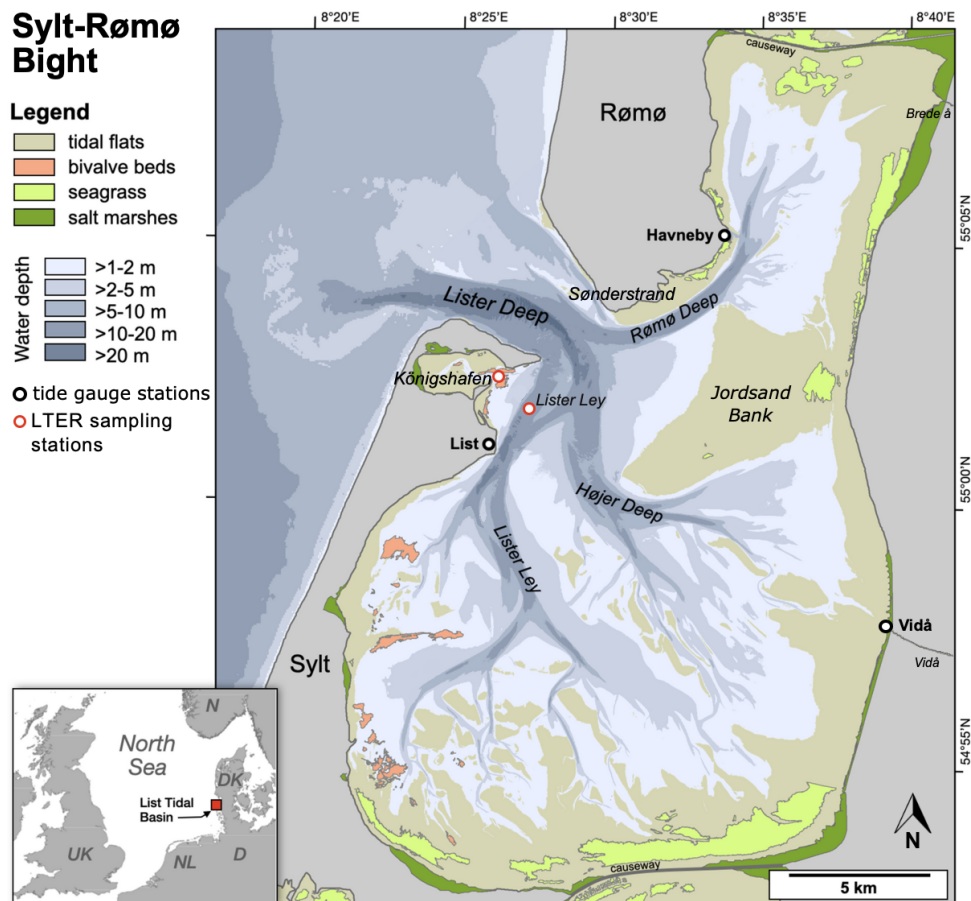


Figure 1: Map of the Sylt-Rømø Bight showing bathymetry, key habitats (tidal flats, bivalve beds, seagrass meadows, and salt marshes), LTER sampling stations (Deep Station – Lister Ley; Shallow Station – Königshafen), tide gauge locations (List, Havneby, Vidå), and rivers (Brede Å and Vidå). The basin is connected to the North Sea via the tidal inlet Lister Deep and laterally enclosed by causeways to the north and south.

2.2 FESOM-C Model

This study used the coastal hydrodynamic model FESOM-C (Androsov et al., 2019). FESOM-C is designed explicitly for high-resolution coastal applications and employs a finite-volume cell-vertex

151 discretization on unstructured meshes composed of triangles and quadrilaterals. This allows for flexible
152 spatial resolution down to several metres, suitable for simulating complex coastal dynamics (Fofonova
153 et al., 2019; Kuznetsov et al., 2020, 2024; Neder et al., 2022; Sprong et al., 2020; Sidorenko et al.,
154 2025).

155 2.2.1 Model Setup

156 The setup utilized an unstructured hybrid mesh of 208,345 nodes and 211,545 elements. Due to the
157 semi-enclosed state of the bight, the mesh contains a single open boundary at the seaward edge of the
158 domain, connecting the basin with the North Sea. The horizontal spatial resolution varies from up to 2
159 m in wetting-drying zones to 304 m in the deeper outer part (near the open boundary). The experiments
160 were carried out by running 2D barotropic simulations with the wetting/drying option enabled to
161 capture the periodic submergence and exposure of intertidal areas. The model timestep was set to ~ 0.25
162 seconds, with data output every ~ 20 minutes of simulation time. The bottom friction coefficient was
163 applied as 0.0025, a value identified as optimal in prior studies of the same study area when using
164 TPXO9 tidal solution (Fofonova et al., 2019; Konyssova et al., 2025).
165 The simulations are driven by tidal forcing alone, applied at the open boundary. For an accurate
166 simulation of the tidal dynamics, thirteen major tidal harmonic constituents (M2, S2, N2, K2, K1, O1,
167 P1, Q1, Mm, Mf, MN4, 2N, and S1) and two over-harmonics (M4, MS4) were prescribed by their
168 phases and amplitudes at the open boundary based on TPXO9 tidal atlas (Egbert and Erofeeva, 2002).
169 This selection of the tidal solution was justified by its robust performance and is one of the most
170 optimal for the North Sea (Fofonova et al., 2019). The current setup has been validated in a previous
171 work by Konyssova et al. (2025). The model's performance has been validated using tidal gauge (TG)
172 data from stations List, Vidå, and Havneby, (see Fig. 1; performance results are provided in the
173 Supplementary Materials, Table S1). Since the numerical setup remains unchanged, we refer to
174 Konyssova et al. (2025) for full validation details.

175 2.2.2 Lagrangian Module

176 To assess tidally driven spatial connectivity and transport timescales within the basin, we performed
177 Lagrangian simulations using FESOM-C Drift, a post-processing tool designed for particle tracking.
178 The model advects massless passive particles based on the velocity fields produced by the
179 hydrodynamic model. In this study, we use the term “passive tracers” to denote Lagrangian particles
180 without weight or settling properties. This is equivalent to virtual Lagrangian particles commonly
181 applied in particle tracking studies and should not be confused with dye experiments that represent
182 concentration changes in space and time.
183 The experiment involved releasing passive tracers from all grid elements within the domain that are
184 consistently inundated during every flood phase. We released about 90,000 tracers at three-hour
185 intervals over six weeks (169 iterations in total). Each tracer was tracked for up to three weeks and was
186 removed from the simulation once it reached either of the two Sylt Roads sampling stations (see Fig. 1
187 and Section 2.3.1). The iterative release process was designed to capture the full range of tidal
188 conditions and the complexity of hydrodynamic transport within the basin, ensuring that the results are

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statistically robust. Upon arrival at a station, tracers were immediately removed to prevent post-arrival movements from influencing the mapped source regions and transport pathways. The simulations were conducted independently for each station, focusing exclusively on the paths from the release locations to the respective sampling site. If a tracer did not reach the designated station within the simulation time frame, it was considered to originate from a region that falls outside the station's dominant transport pathways.

The first analysis approach assessed the source regions of passive tracers arriving at the sampling stations. This allows us to evaluate the connectivity between different subareas of the basin and the sampling sites. The probability of SPM originating from a given area was mapped based on the cumulative occurrence of tracer pathways across all iterations. Higher probability values indicate areas that more frequently serve as source regions or transport pathways for SPM reaching the sampling stations.

The second part of the analysis was conducted to estimate the mean transit time of the tracers reaching the sampling stations from shallow source zones, where resuspension typically occurs (defined here as areas <2 m deep, based on de Jonge & van Beusekom, 1995). The mean transit time over all implementations was calculated for all elements whose tracers reached the stations within the simulated three weeks. To quantify how long it typically takes for high tracer concentrations to reach the station, we computed a probability-weighted median transit time, where mean transit times were weighted by their probability values. This approach ensures that frequent transport pathways are given greater influence in the median transit time calculation, reducing bias from rare, low-probability trajectories.

2.3 Data

2.3.1 Biogeochemistry data

This study used data from the Sylt Roads long-term ecological monitoring program, focusing on a subset from 2000 to 2019 to ensure consistent methodology and regular sampling. From a broad range of hydrographic and biogeochemical parameters covered in the dataset, this study specifically analyzed suspended particulate matter (SPM; mg/L, filtered through 0.4 µm nucleopore filters, rinsed with distilled water, stored frozen, and dried at 60 °C) and chlorophyll-a (Chl-a; µg/L, filtered through GF/C filters (Whatman), stored at -20 °C, and extracted using 90% acetone and analyzed spectrophotometrically using the trichromatic method as described by Jeffrey and Humphrey, (1975). Both parameters were measured twice weekly at a sampling depth of 1 m below the surface at two primary stations: the deep station at the Lister Ley channel and the shallow station at the entrance of Königshafen embayment (see Fig. 1). The full dataset is publicly available on the data portal PANGAEA (<https://www.pangaea.de>) and the recent evaluation is detailed in Rick et al. (2023). To analyze seasonal variability, we defined the seasons based on observed cycles in Chl-a concentrations at the study site, rather than calendar or astronomical definitions. Specifically, we used:

- winter (November 20 – February 19, low biological activity, low Chl-a);
- spring (February 20 – May 31, phytoplankton bloom initiation and peak);
- summer (June 1 – September 19, post-bloom conditions, high light, reduced Chl-a);
- autumn (September 20 – November 20, transitional period).

229 **2.3.2 Meteorological data**

230 For the statistical analysis, we also downloaded the quality checked historical meteorological data for
231 station 3032, List auf Sylt, from Climate Data Center (CDC) of the Deutscher Wetterdienst (DWD).
232 The data includes hourly mean wind speed and wind direction (dataset ID: urn:x-
233 wmo:md:de:dwdcdc::obsgermany-climate-hourly-wind), and daily sunshine duration (dataset ID:
234 urn:wmo:md:de-dwd-cdc:obsgermany-climate-daily-kl).

235 **2.3.3 Sea Surface Height data**

236 Using the validated model setup (Fofonova et al., 2019; Konyssova et al., 2025), the sea surface height
237 (SSH) data were reconstructed for the deep and shallow stations. In particular, the amplitudes and
238 phases of the 15 harmonics mentioned above were obtained from the modeling output using Fast
239 Fourier Transform analysis. Subsequently, the SSH signal was reconstructed for the exact timestamps
240 of the LTER sampling using the *t_tide* package (Pawlowicz et al., 2002). This allowed us to estimate
241 the tidal elevation and phase (through the SSH and SSH gradients) at the time of sampling, which were
242 then included as input features in the statistical and neural network (NN) analyses. Their relationship is
243 illustrated and discussed in Supplementary Material, Figs. S3–S4.

244 **Table 1. The physical and hydrochemical parameters used in the study (short name, units, frequency, and source)**

Data	Unit	Frequency	Source
SPM	mg/L	twice weekly	Sylt Roads Marine Observatory
Chl-a	µg/L	twice weekly	Sylt Roads Marine Observatory
Wind speed and direction	m/s, degrees	hourly mean	Deutscher Wetterdienst
Light	hours	daily	Deutscher Wetterdienst
SSH	m	every 5 minutes	Model reconstruction using 13 harmonics and 2 over-harmonics from TPXO9, and verified with the tidal gauges

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245 **2.4 Neural Network**

246 To assess the relative contribution of biotic conditions to SPM concentrations, we employed a forward
247 NN to predict SPM based on environmental input parameters. The approach relies primarily on long-
248 term observational data, thereby enhancing the robustness of the results, complemented by model-
249 derived SSH and its temporal gradient data. Although numerical simulations are also powerful tools in
250 this context, any discrepancies they produce can largely be attributed to the choice of numerical
251 methods and the associated uncertainties, including unresolved processes, boundary conditions, and the
252 spatial and temporal resolution of both modelled and observed data (Skogen et al, 2021). While it does
253 not provide process-based understanding, the NN approach is particularly well-suited for this task, as it
254 offers a cost-efficient and data-driven means to capture complex, non-linear relationships between
255 influencing factors. We conducted several sensitivity experiments (not shown), varying both the

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network depth and the number of neurons per layer. For the current application, increasing the network depth further did not improve performance.

For the first part of the experiment, the primary regression task involved predicting SPM concentrations during the winter season, which is characterized by low biological activity, as indicated by minimal Chl-a concentrations. Focusing on winter allows for a clearer assessment of physical (abiotic) drivers, such as wind forcing, with reduced biological confounding. The input features, 19 in total, of the NN model include wind magnitude at the time of sampling and averages over a series of prior time intervals (6 to 240 hours, corresponding to the intervals analyzed in Subsection 3.2.2), dominant wind direction over 6 and 12 hours (even though the correlation analysis shows only a minor impact, the non-linear effects of wind direction may still be present), salinity, SSH, and the temporal gradient of SSH (computed using a forward scheme). The results of the winter model are presented in Section 3.3.1.

To extend the analysis across seasons, we applied the same model architecture to the full dataset, initially using the same input features as in the winter setup (results in Section 3.3.3), and subsequently incorporating two additional features: temperature and the weekly sum of sunshine hours before the measurement date (21 features in total, Section 3.3.4). The latter serve as a pragmatic proxy for both Chl-a concentration and benthic algae abundance. While we do not attempt to separate these mechanisms individually, this approach is exploratory and intended to approximate their combined influence on seasonal SPM variability.

The technical details of the NN architecture are provided in the Supplementary Material, together with the complete predictor list in Table S2.

3 Results

Figure 2 presents the time series of SPM concentrations at the deep and shallow stations in the Sylt-Rømø Bight from 2000 to 2019, based on data from the Sylt Roads monitoring program (Section 2.3.1; station locations shown in Fig. 1). Both stations display a pronounced seasonal cycle, with SPM concentrations typically peaking in winter and declining during summer. The deep station shows more frequent and sustained seasonal peaks in SPM concentrations throughout the time series, whereas the shallow station tends to exhibit higher concentrations during peak events. It is also important to note that the regular sampling at the shallow station was discontinued after 2013 following the replacement of the research vessel, resulting in reduced data coverage in subsequent years.



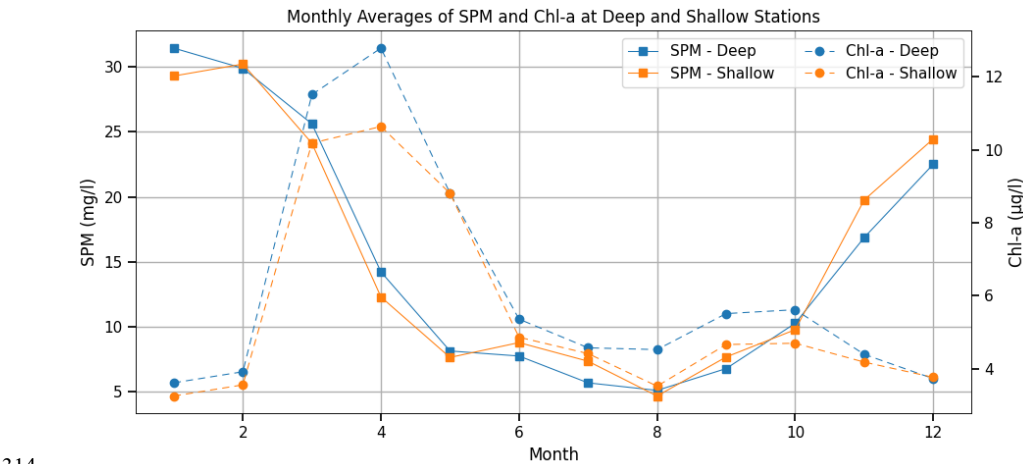
Figure 2: Time series of SPM for the considered years (2000–2019) for the deep (blue) and shallow stations (orange) gridded per year. Each subplot represents a calendar year with time on the x-axis and SPM concentration [mg/l] on the y-axis.

3.1 Seasonality of SPM concentrations

SPM concentrations show an apparent seasonality, which is further investigated in relation to biological activity and meteorological drivers in the subsequent analyses. Using Chl-a as a proxy for phytoplankton biomass, this subsection applies statistical analysis to examine how biological activity and wind forcing together shape seasonal patterns of SPM variability at both the deep and shallow stations.

299 **3.1.1 Role of biological processes in seasonal SPM variations**

300 Both stations exhibit distinct seasonal variations in SPM and Chl-a concentrations (Fig. 3). SPM
301 concentrations are highest in late autumn and winter, with median values of about ~28 mg/L ($33.9 \pm$
302 18.2 mg/L) at the deep and median of ~26 mg/L (38.6 ± 16.4 mg/L) at the shallow stations. In January,
303 the peak reaches over 60 mg/L at both stations, with individual events exceeding 70 mg/L. The decline
304 in concentrations is observed from February to May, reaching their lowest values in June to August
305 (around 2-3 mg/L), although occasional peaks above 30-40 mg/L still occur. From September onward,
306 SPM begins to increase. This pattern is similar at both stations, though the shallow station generally has
307 slightly higher SPM concentrations, suggesting potential differences in sediment availability or
308 resuspension dynamics.
309 Chlorophyll-a concentrations follow an inverse seasonal pattern. It remains low in December–January
310 with a median around 2 $\mu\text{g/L}$ (2.1 ± 0.7 $\mu\text{g/L}$) at both stations. It increases sharply in early spring, with
311 peaks in March–April exceeding 30 $\mu\text{g/L}$ at the deep station and 25 $\mu\text{g/L}$ at the shallow station.
312 Concentrations then drop in summer, followed by a secondary increase in August–October, when values
313 up to ~15-20 $\mu\text{g/L}$ are observed.



314 **Figure 3: Monthly averages of SPM and Chl-a at the deep (blue) and shallow (orange) stations for the considered years (2000–**
315 **2019). SPM concentrations (solid line with square marker, left axis) and Chl-a concentrations (dashed line with circle markers,**
316 **right axis) are displayed separately for clarity. The x-axis represents the months from January to December.**
317

318 The relationship between Chl-a and SPM varies across seasons (Fig. 4). During December–February,
319 biological activity is low and Chl-a constitutes a relatively constant fraction of total SPM, both subject
320 to the same resuspension-deposition processes. This co-settling behavior leads to strong positive
321 correlations observed at both stations. The highest values occur at the shallow station, with $R^2 = 0.84$ in
322 December and $R^2 = 0.75$ in January. At the deep station, correlations are relatively lower, reaching $R^2 =$

0.52 in December and $R^2 = 0.56$ in January. The strong correlation and the slope are in line with the resuspension of microphytobenthos as observed in winter by de Jonge and van Beusekom (1995). From March onward, the relationship becomes more complex. As Chl-a rises rapidly during the spring bloom and SPM concentrations decline, the correlation at both stations weakens with R^2 values dropping to near-zero in March. This divergence reflects differences in vertical distribution and settling behavior. While phytoplankton is a component of SPM and both are retained on the filter, their decoupling in spring and summer may reflect that not all phytoplankton is flocc-associated. Active phytoplankton may remain suspended near the surface, while denser biomineral flocs sink more rapidly, leading to weaker correlations in integrated surface samples. This non-linear and temporally variable influence of Chl-a also underlies the decision not to use it directly as a predictor in the neural network (Section 3.3), but instead to approximate biological activity through more general proxy variables such as temperature and light availability (see Table S2 in Supplementary Material). During June–August, as both the SPM and Chl-a concentrations reach low values, the correlation weakens further, with August showing almost no correlation at the shallow station. In November and December the correlation strengthens again as the role of biological activity reduces and physical drivers become dominant again.

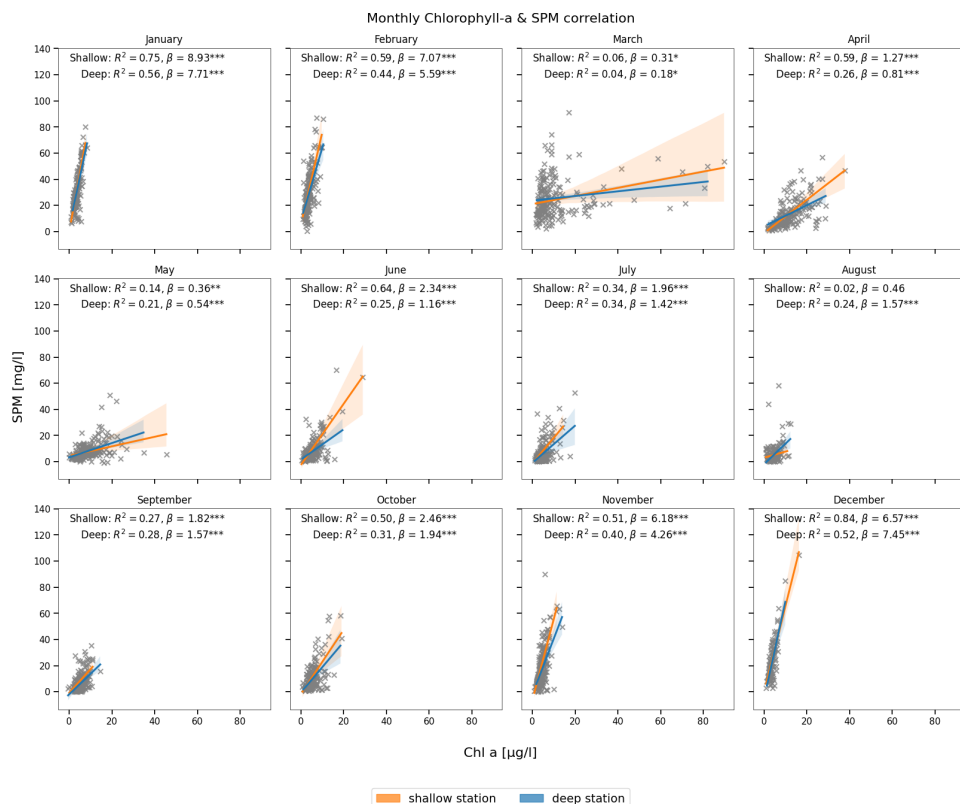
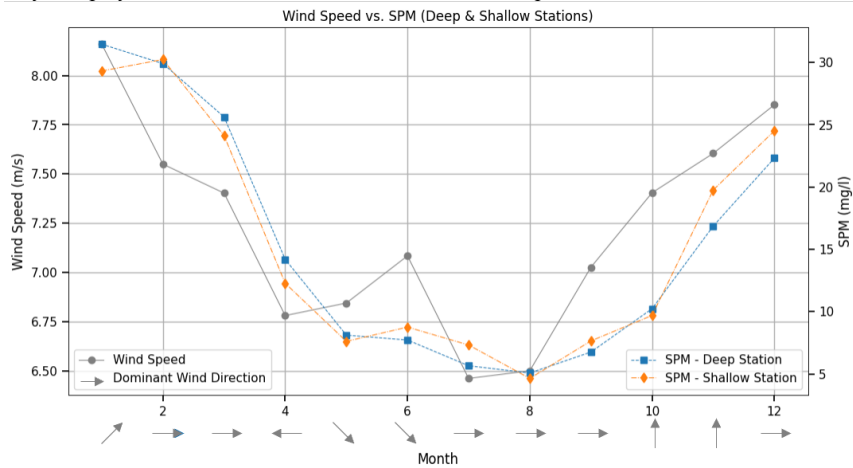


Figure 4: Monthly correlation between Chl-a (x-axis) and SPM (y-axis) concentrations at the deep (blue) and shallow (orange) stations. The coefficient of determination (R^2), slope (β), and statistical significance ($p < 0.001$, ** $p < 0.01$, * $p < 0.05$) are indicated in each panel.**

3.1.2 Role of wind forcing in seasonal SPM variations

The relationship between seasonal SPM concentrations and wind characteristics is illustrated in Fig. 5. Wind speeds are highest in January, averaging around 8.5 ± 2.8 m/s, with peaks reaching 14.2 ± 1.7 m/s, and gradually decline through spring months, reaching a minimum with mean speeds around 6.5 ± 3.1 m/s and minima of 2.0 ± 0.7 m/s in July–August. From August onward, wind speeds begin to rise again. This seasonal cycle aligns with the observed variability in SPM, with higher concentrations in winter and lower values in summer. Dominant wind directions also vary throughout the year, with

349 westerly and northwesterly winds prevailing in most months, easterly winds in April, and more
 350 southerly winds during October–November. Seasonal wind roses illustrating these patterns are shown in
 351 the Supplementary Material (Fig. S1). However, no clear linear relationship emerges between dominant
 352 wind direction and seasonal SPM concentrations. To account for potential non-linear interactions that
 353 may still play a role, wind direction was included as an input feature in the NN model.



354
 355 **Figure 5: Monthly averages of wind speed, SPM concentrations, and dominant wind direction (2000–2019).** The figure illustrates
 356 monthly mean wind speed (grey solid line, left axis) and SPM concentrations at the deep (blue squares) and shallow (orange
 357 diamonds) stations (right axis). Grey arrows along the x-axis represent the dominant wind direction for each month, with arrow
 358 orientation indicating the direction from which the wind originates, following standard meteorological convention.

359 While correlations are further explored in Section 3.2, the seasonal alignment of wind intensity and
 360 SPM concentrations suggests strong physical modulation.

361 3.2 Resuspension and Time Scales of Inner Basin Transport

362 Beyond seasonal variation, there are mechanisms responsible for the variability of SPM concentrations
 363 on shorter temporal scales, from hours to days. This subsection examines how wind-induced
 364 resuspension operates over different time frames and the role of tidal transport within the basin.

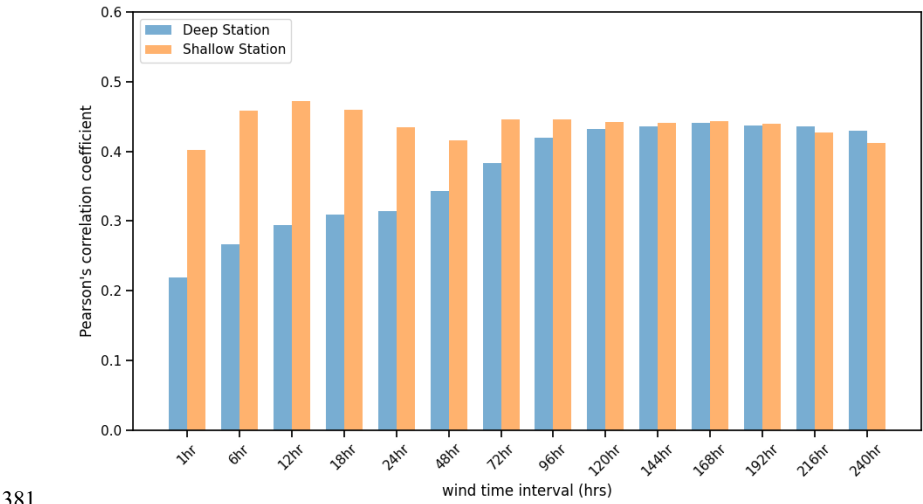
365 3.2.1 Role of wind forcing in short time SPM variations

366 To quantify the short-term response of SPM to wind forcing, we computed Pearson correlation
 367 coefficients between SPM concentrations and wind speed averaged over different time intervals,
 368 ranging from 1 hour to 240 hours. This approach, commonly referred to as “wind memory,” allows us
 369 to evaluate how the cumulative influence of past wind conditions affects SPM variability at different
 370 depths and over varying time scales. Figure 6 illustrates how the strength of this relationship evolves as

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373 wind speeds are averaged over progressively longer intervals, ranging from 1 hour to 240 hours. This
374 approach quantifies how SPM responds to the cumulative influence of past wind conditions over
375 varying time scales. The results show that correlation coefficients generally increase as wind memory
376 lengthens, reaching a peak around 12-18 hours at the shallow station and 120 hours at the deep station,
377 followed by a slight decline.
378 Monthly variations in the correlation patterns are provided in the Supplementary Material (Fig. S2),
379 showing that the correlation between wind speed and SPM is generally stronger in winter than in
380 summer, with more complex patterns during transitional months such as April and November.



381
382 **Figure 6:** Pearson correlation coefficients between SPM and wind speed averaged over different time intervals, ranging from 1
383 hour to 240 hours. The two stations (deep station as blue and shallow station as orange) are represented separately, showing how
384 wind memory influences SPM variability at different depths.

385 The correlation between instantaneous wind speed and SPM concentrations is strongest during winter
386 and early spring (Fig. 7), particularly at the shallow station, where the most substantial value occurs in
387 January ($R^2 = 0.44$) and April ($R^2 = 0.50$). In contrast, the deep station shows much weaker correlations,
388 with R^2 values often near zero, reflecting a lagged effect on SPM concentrations.
389 During June–August, the relationship weakens significantly, with the lowest correlations observed in
390 August for both stations. The shallow station maintains minor correlation ($R^2 \approx 0.15$) into late summer
391 and early autumn.

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Deleted: This suggests that wind-driven resuspension may play a relatively stronger role in regulating SPM concentrations in winter.

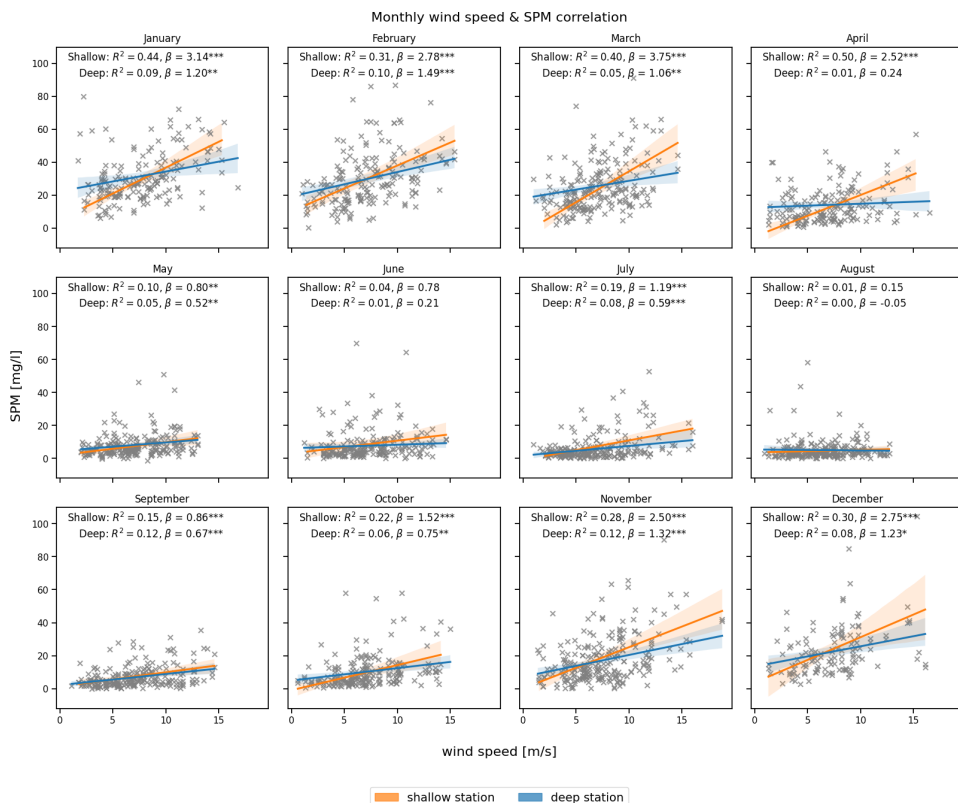


Figure 7: Monthly correlation between wind speed at the time of sampling (x-axis) and SPM concentration (y-axis) at the deep (blue) and shallow (orange) stations. The coefficient of determination (R^2), slope (β), and statistical significance (* $p < 0.001$, ** $p < 0.01$, * $p < 0.05$) are indicated in each panel.**

In contrast to the instantaneous impact of wind-speed, averaging wind speed over 120 hours (5 days) results in an improved correlation between wind and SPM at the deep station, as shown in Fig. 8, although the seasonal pattern remains similar. The stronger relationship is particularly evident in the winter and autumn months, especially at the deep station (e.g., $R^2 = 0.35$ in September, 0.24 in November and February). These findings indicate that the average wind-speed over several days have a stronger influence on SPM concentrations at the deep station than the wind at the time of sampling. The results highlight the role of wind forcing in modulating SPM concentration variability, particularly in shallow waters during winter and autumn. Naturally, the influence of wind is less pronounced in

deeper waters through direct resuspension mechanisms. However, it remains significant, with a time lag possibly caused by the transport of resuspended material from shallower zones to the deep station.

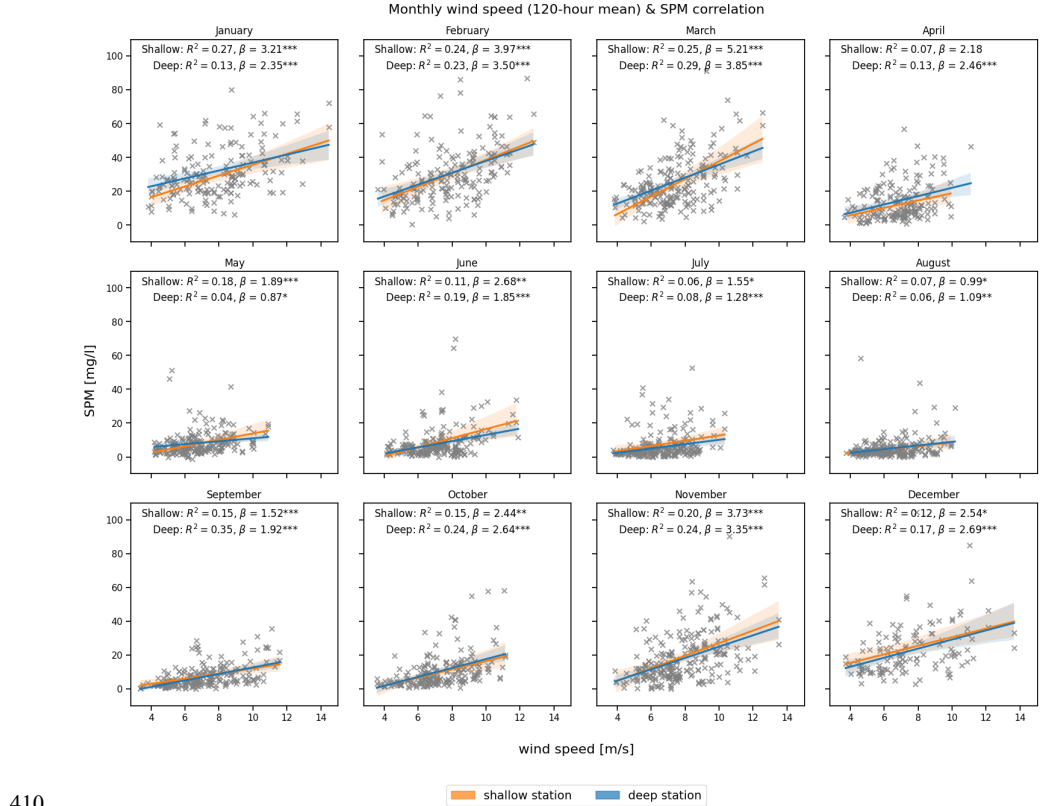


Figure 8: Monthly correlation between wind speed averaged over a 120-hour (5-day) period (x-axis) and SPM concentration (y-axis) at the deep (blue) and shallow (orange) stations. The coefficient of determination (R^2), slope (β), and statistical significance ($***p < 0.001$, $**p < 0.01$, $*p < 0.05$) are indicated in each panel.

3.2.2 Role of tidal forcing in short time SPM variations

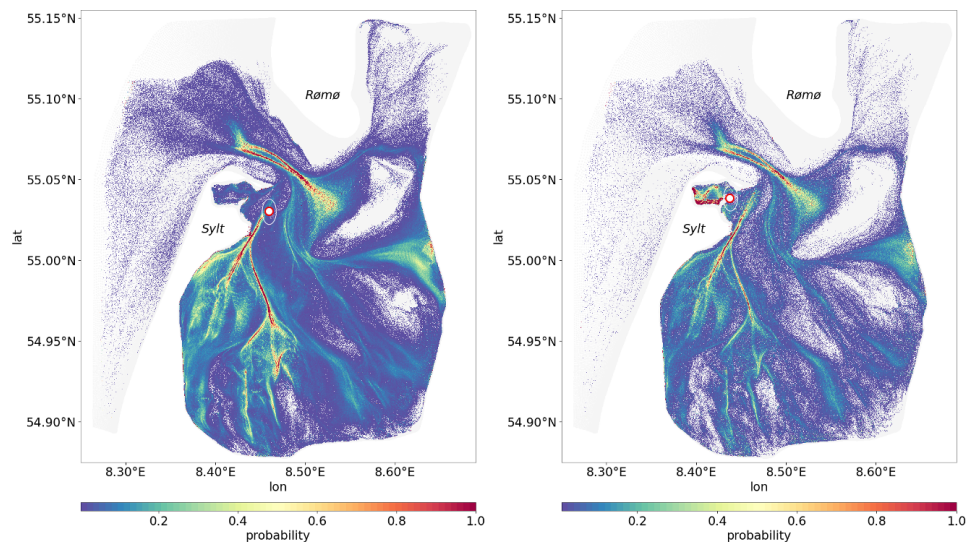


Figure 9: Probability distribution of passive tracer pathways to illustrate how frequently different areas act as source regions or transport corridors for passive tracers. Higher probability values (highlighted in yellow and red) indicate locations that more frequently contribute to SPM reaching the deep station (left panel) or shallow station (right panel).

Figure 9 shows the spatial probability distribution of passive tracer connectivity within the basin, derived from Lagrangian simulations forced solely by tides. This setup isolates the dominant physical driver of material transport in the Sylt-Rømø Bight, where tidal processes account for roughly 80% of velocity variability (Fofonova et al., 2019). While regular wind forcing (excluding storm events and uncommon multi-day episodes of strong winds from a single direction) enhances lateral dispersion, its net contribution to basin-wide transport over several days is relatively minor (Konyssova et al., 2025). Note, in this region, typical wind events last about 5–6 hours, defined as consecutive hours of wind from the same direction within an eight-sector classification (Rubinetti et al., 2023). Residual currents generated by non-linear tidal interactions establish consistent, directional transport pathways, which are effectively captured by the simulated tracer trajectories. The resulting network of tidally-induced transport pathways reveals apparent differences between the two stations. The shallow station is predominantly supplied by tracers originating in the northern parts of the bight, particularly Königshafen and its surroundings, reflecting localized and relatively rapid connections. In contrast, the deep station is influenced by broader and more distributed transport pathways, integrating material from a wider area of the basin.

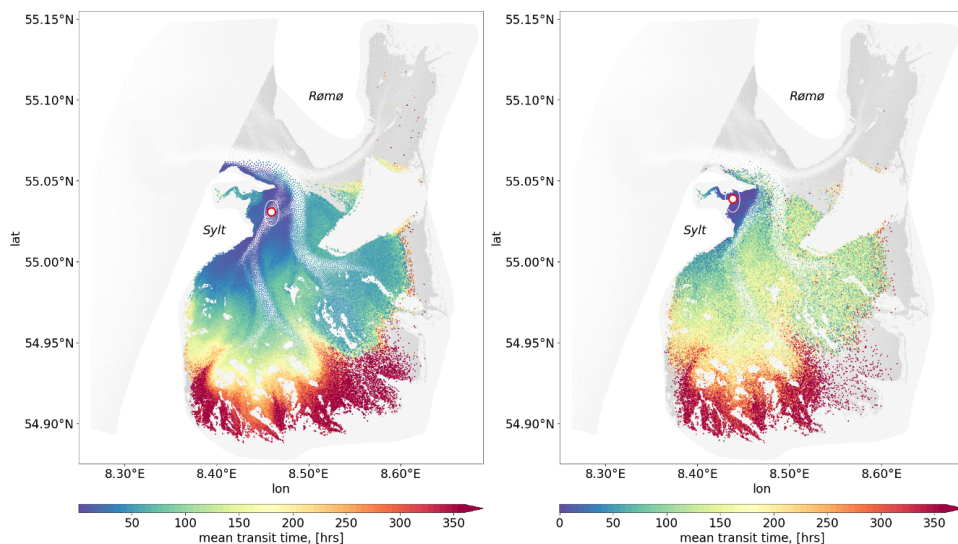


Figure 10: The mean transit time of passive tracers to reach the sampling station. The left panel corresponds to the deep station, while the right panel represents the shallow station. Grid elements that are consistently inundated with each flood phase, from which the passive tracers are released, are shown in grey. Elements that successfully reach one of the two sampling stations are colour-coded by their mean transit time (in hours), reflecting how long it takes for passive tracers to reach from a given location to each station.

For both stations, shorter transit times (depicted in blue and green, Fig. 10) are observed in areas closer to the stations, whereas regions further away, particularly in the inner tidal flats, exhibit longer transit times (yellow to red hues, Fig. 10). The transit time patterns reveal that SPM originating from intertidal areas and tidal channels follows particular pathways before reaching the deeper and shallower monitoring stations, with transport occurring on timescales of days to weeks. The difference between the two panels suggests that the deep and shallow stations receive material from primarily distinct but partially overlapping source regions.

Together, these figures illustrate the tidal connectivity of the two monitoring stations with their surrounding basin, highlighting likely source regions and typical transport pathways. To quantify the typical timescale for SPM transport, a probability-weighted transit time was computed, where individual transit times were weighted according to their interpolated probability values. This approach ensures that the estimated transit time reflects the most frequently occurring transport pathways rather than rare, low-probability trajectories. The resulting probability-weighted median transit time was 133.3 hours for the deep station, which aligns with the ~5-day wind-memory interval identified in Section 3.2.2, and 44.4 hours for the shallow station, highlighting the pronounced contrast in transport efficiency between the two locations [and likely reflecting differences in sediment source regions.](#) [Although not explicitly resolved in the simulations, this difference may also relate to grain-size](#)

composition, as the deeper channels typically contain coarser material, whereas the intertidal flats have finer, more cohesive sediments (Bartholomä et al., 2009). At the shallow station, which is situated within the Königshafen area, SPM concentrations respond to wind forcing almost immediately through local resuspension. The longer transit time therefore does not indicate a delayed response, but rather reflects the additional supply of tracers arriving from a wider surrounding area, which extends the median transit time beyond the short wind-driven response captured in the memory analysis. In contrast, the deep station integrates SPM from more complex, multi-step transport routes over longer timescales. These transit times provide a key reference for interpreting observed SPM fluctuations at each station and may help distinguish between short-term and cumulative drivers of SPM variability.

3.3 Neural Network

3.3.1 Winter SPM prediction (Baseline Model)

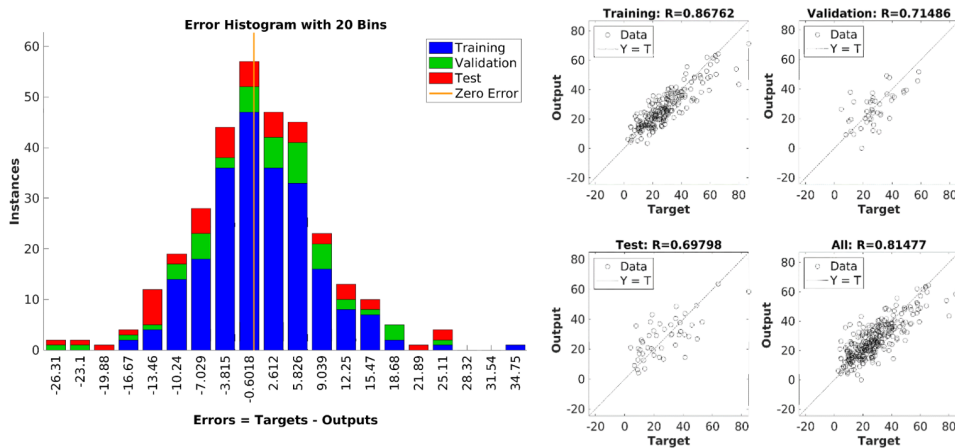


Figure 11: Performance of the neural network (NN) trained on the winter dataset for the deep station. Left: Error histogram showing the distribution of prediction errors (in mg/L) for training, validation, and test subsets. Right: Regression plots comparing predicted vs. observed SPM values, with correlation coefficients (R) for each data subset. Due to the overall qualitative similarity between the pictures for deep and shallow stations, only the deep station is presented here.

To evaluate how well physical drivers alone can explain SPM concentration variability, we first trained a baseline NN using only the winter dataset, when biological activity is minimal. This provides a reference to assess the contribution of wind and tidal forcing under conditions with limited biological influence.

The error for training, validation and testing has a normal distribution (Fig. 11, left). For the deep station, the root mean square error (RMSE), a measure of average prediction error magnitude, is 9.4 mg/L for the testing set. Note that the observed mean and median SPM concentrations in winter are 27.6

481 and 25.9 mg/L respectively. As a result of the application of NN to the winter dataset, they are 27.2 and
 482 25.5 mg/L respectively (Table 2). Regression analysis (Fig. 11, right) yields correlation coefficients
 483 between observed and simulated SPMs equal to ~ 0.87 , ~ 0.71 , ~ 0.7 and ~ 0.81 for training, validation,
 484 testing and all winter datasets, respectively.
 485 The results show that, by using tidal and wind forcing along with a proxy for baroclinic conditions
 486 (salinity), we can predict SPM concentrations during winter quite well without accounting for other
 487 factors.
 488 Notably, temperature, salinity, turbulent kinetic energy, and SPM concentrations themselves all
 489 influence flocculation processes, even without considering biological activity. Temperature was
 490 excluded from the feature set at this stage due to its strong seasonal cycle and its potential use as a
 491 proxy for biological conditions. Specifically, in our current approach, we cannot separate the influence
 492 of temperature on biological mechanisms from its physical effects (e.g. viscosity), which also impact
 493 flocculation.

494 **3.3.2 Applying of NN trained on winter data to other seasons**

495 Next, we applied the NN trained under winter conditions to other seasons. This approach is justified by
 496 the fact that the long-term dataset captures a representative range of wind conditions in both winter and
 497 summer, including calm periods and strong wind events. Furthermore, salinity remains relatively stable
 498 across seasons, supporting the applicability of the model. Using the trained winter model, we attempt to
 499 predict SPM concentrations for spring, summer, and autumn. The goal of this step is to estimate the
 500 influence of biological processes on SPM concentrations.
 501 For both stations, the results of the NN application show significantly higher SPM concentrations
 502 compared to observations (Table 2).

503 **Table 2. Mean, median values and correlation (R) of SPM concentrations, [mg/L], from observations VS predictions by NN**
 504 **trained on winter dataset for the deep and shallow stations**

seasons		Deep Station		Shallow Station	
		observed	predicted	observed	predicted
winter	mean, [mg/L]	27.6	27.2	28.5	24.7
	median, [mg/L]	25.9	25.5	24.4	26.5
		$R = 0.81$		$R = 0.81$	
spring	mean, [mg/L]	16.5	24.8	15.6	20.6
	median, [mg/L]	13.3	24.3	11.3	18.0
		$R = 0.29$		$R = 0.5$	
summer	mean, [mg/L]	6.15	17.7	6.8	18.5
	median, [mg/L]	4.22	16.9	3.9	16.1
		$R = 0.04$		$R = 0.35$	

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autumn	mean, [mg/L]	11.6	23.4	12.8	24.0
	median, [mg/L]	8.5	22.7	7.5	21.0
		$R = 0.44$		$R = 0.54$	

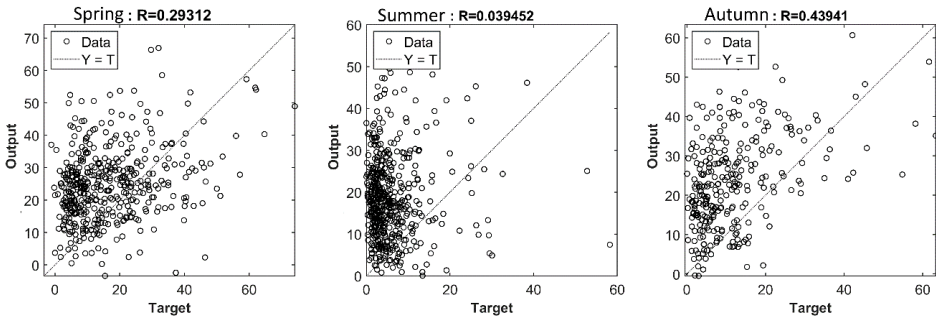


Figure 12: Regression analysis for NN trained on winter data and applied to spring, summer and autumn data vs observations. Results shown for the deep station; shallow station results are qualitatively similar and summarized in Table 2.

This discrepancy suggests that while calmer wind conditions alone can explain approximately 40% of the SPM reduction in summer compared to winter, they do not account for the full ~80% decrease observed in the data (see Table 2). The overestimation of SPM by the NN is clearly visible in the regression plots (Fig. 12), where most of the predicted values are positioned above the $Y = T$ line, particularly during summer and autumn. This does not necessarily imply that biological activity is weaker in spring, but rather that its impact is less well captured by the abiotic-only model, as biological processes contribute more strongly to variability than in autumn. The higher correlation between simulated and observed SPM in autumn likely reflects physical conditions that are more similar to winter, when the model was trained, including generally stronger winds (Section 3.2.2). In spring, enhanced biological activity and aggregation introduce variability that the model cannot account for, reducing correlation. During summer, [the regression coefficient dropping below 0.05 indicates that processes not represented in the abiotic model become dominant \(Fig. 12\). The subsequent analysis of all seasons presented in Section 3.3.3 points to biologically mediated mechanisms, such as flocculation and sediment stabilization, as the main contributors to this variability.](#)

3.3.3 NN trained at the data from all seasons

Previously we showed that there are factors, in particular biological processes, which significantly influence the SPM dynamics. To confirm this, we trained NN on all the dataset (not limited to winter) using the same input features representing only abiotic conditions, which we used in 3.3.1. As expected, the overall RMSE increases to approximately 11 mg/L, due to the omission of important features that influence SPM dynamics during the warm season (Supplementary Fig. S5). When the model is trained on data from all seasons, and applied specifically to winter conditions, it

Deleted: biological processes become the primary driver of SPM concentration variability, as evidenced by the regression coefficient dropping below 0.05 (Fig. 12).

underestimates SPM concentrations. For the deep station, the predicted mean and median are 19.17 mg/L and 17.2 mg/L, respectively, compared to observed values of 27.6 mg/L and 25.9 mg/L. At the shallow station, the model yields mean and median predictions of 20 mg/L and 17 mg/L, whereas the corresponding observed values are 28.45 mg/L and 24.35 mg/L. The regression coefficients also reflect this underperformance, with values of approximately $R \approx 0.66$ (deep) and 0.72 (shallow). The seasonal regression analysis further highlights these biases (Supplementary Fig. S7, upper row). A complete numerical breakdown of observed and predicted seasonal values is provided in Table S3.

3.3.4 Neural network trained on all-season data including biological proxy features

Including temperature and sunshine duration as proxies for biological activity improved overall model skill, reducing the test RMSE to ~10 mg/L. The regression analysis shows the correlation coefficient between observed and simulated SPM concentrations reached ~0.88, ~0.76, ~0.72 and ~0.84 for training, validation, testing and all datasets respectively (Supplementary Fig. S6). Seasonal regression plots also show marked improvements (Supplementary Fig. S7, lower row). For winter, R increased from 0.66 to 0.73 at the deep station and from 0.72 to 0.82 at the shallow station. The largest gains occurred in spring and summer, consistent with more substantial biological influence during these seasons. Table S3 provides the detailed seasonal comparison of observed and predicted means, medians, and correlations.

4 Discussion

This study set out to investigate the combined influence of wind forcing, tidal transport, and biological processes on SPM concentration variability in the Sylt-Rømø Bight. Using 20-year period monitoring data and applying statistical tools, neural network models, and Lagrangian simulations, we analyze the relative roles of these processes across seasonal and short-term timescales.

The variability observed in SPM concentrations reflects the inherent complexity of the system, where several physical and biological mechanisms interact across different spatial and temporal scales. Wind-driven resuspension, tidal advection, and biologically mediated effects each contribute to the observed dynamics, with their relative importance shifting seasonally. This complexity highlights that the variability in SPM concentrations cannot be attributed to a single driver but rather to the combined response of the system to multiple, overlapping processes.

In this chapter, we will further discuss topics that have not yet been fully addressed: the peculiarities of the Chl-a and SPM concentration relationship, biological processes that directly or indirectly influence SPM dynamics, further details on wind and tide control including SSH and SPM concentration relationship, details of NN related findings and study limitations.

4.1 Biological Interaction

The strong positive correlation between Chl-a and SPM concentrations in winter suggests that under low biological activity, such as limited phytoplankton growth, behaviour of both variables is driven mainly by physical processes such as wind-driven resuspension. This is consistent with previous

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observations in the Wadden Sea and German Bight (van Beusekom et al., 1999; van Beusekom and de Jonge, 2002). Together with SPM, the strong winds resuspend microphytobenthos attached to sediment particles (de Jonge and van Beusekom, 1995), consistent with observations by Hommersom et al. (2009), who found strong correlations between SPM and Chl-a in the Wadden Sea driven by resuspension. Such patterns support the interpretation that, under winter conditions with low biological production, a large fraction of Chl-a is contained within resuspended flocs, leading to the high correlations observed in our study at both stations. In contrast, the correlation between Chl-a and SPM concentrations begin to decline in spring, despite persistent wind forcing, and reaches a minimum in summer. This seasonal decoupling suggests that biological aggregation processes become increasingly dominant with the beginning of and following phytoplankton bloom, promoting the formation and subsequent rapid settling of flocs (Schartau et al., 2019; Maerz et al., 2016; Lunau et al., 2006). For example, the production of TEPs, secreted by phytoplankton and bacteria under certain physiological or nutrient conditions, strongly enhance particle stickiness and promote the aggregation of both organic and mineral particles (Passow, 2002). While our statistical analysis (however, not NN related efforts) uses Chl-a as a proxy for biological activity, it is worth noting that TEP production responds to additional drivers such as nutrient status and species composition, which can modulate flocculation potential even at similar Chl-a levels. In addition to the biological feedback, the increase in water temperature during summer also accelerates particle sinking rates by reducing water viscosity, further promoting sediment deposition (Maerz and Wirtz, 2009). This supports the view that the Chl-a–SPM relationship reflects an interplay of multiple biological and physical controls rather than a single, direct pathway.

While both spring and autumn are characterized by peaks in Chl-a concentration, their influence on SPM dynamics appears to differ. In spring, calmer conditions favor particle aggregation and settling, often leading to reduced SPM levels. In autumn, however, the secondary bloom coincides with a seasonal increase in wind speed (Fig. 5), resulting in elevated SPM concentrations despite high biological activity. This suggests that enhanced organic matter availability in autumn may promote flocculation, but stronger resuspension prevents effective settling. These seasonal contrasts highlight that biological influence on SPM is strongly modulated by concurrent physical forcing.

In parallel, benthic microbial processes such as biofilm formation by microphytobenthos enhance sediment stability and reduce its susceptibility to resuspension, thereby modulating the amount of SPM remaining in the water column (Andersen, 2000; Stal, 2010). Beyond microbial stabilization, benthic fauna also influences SPM dynamics through benthic–pelagic coupling. Filter-feeding benthos, such as bivalves, consume suspended particulate organic carbon (POC), effectively removing material from the water column and altering vertical fluxes of organic matter. In the Sylt-Rømø Bight, food web modelling by Baird et al. (2004) estimated that benthic consumers remove approximately $56.7 \text{ mgC m}^{-2} \text{ d}^{-1}$ of suspended POC – potentially a substantial fraction of total SPM, depending on the proportion of organic material within it. Benthic consumption likely constitutes a significant sink for fine, organic-rich particles, particularly in areas with dense benthic communities, as filter feeders remove SPM primarily through ingestion and the production of faecal pellets, which promote deposition of organic-rich material to seabed. Although the proportion of POC within SPM is known to depend on total SPM concentration (Schartau et al., 2019), robust SPM-to-POC conversion factors remain uncertain due to the unknown share of freshly formed organic matter. Although the effects of native species on benthic

pelagic coupling have been relatively well studied in the region (Baird et al., 2007), the role of introduced species remains less clear. For example, the introduction and spread of alien species such as the American razor clam (*Ensis leei*) may further amplify benthic filtration effects. In the Dutch Wadden Sea, they have shown to significantly alter trophic carbon flows, increasing carbon consumption by secondary producers and redirecting energy flows away from higher trophic levels (Jung et al., 2020). Although the abundance of *E. leei* in the Sylt-Rømø Bight has not yet been quantified, its potential influence on observed SPM concentrations provides a basis for further investigation as part of long-term ecological changes in the region.

4.2 Wind & Tide Control

The results suggest that the intensity of winds, rather than their dominant directions, exerts a stronger influence on the seasonal cycle of SPM. The seasonal patterns of wind speed, with maximum values in winter and minima in summer, aligns closely with the observed SPM concentrations, suggesting a physically mediated control during high-energy periods. During summer, the weaker relationship between wind speed and SPM concentrations points to the increasing importance of biological contribution. These may include enhanced flocculation and settling of particles, increased filtration by benthic organisms, reduced sediment availability due to biological stabilization, and overall calmer wind conditions. Although tidal forcing itself does not vary seasonally, its influence on SPM concentrations also likely changes throughout the year because the sediment properties and biological activity that determine sediment availability and erodibility vary over time.

Notably, correlations between wind speed and SPM are stronger at the shallow station compared to the deep station, as the shallow environments are more susceptible to wind-induced turbulence, including wave action. It should be noted that, since the study addresses observations of SPM concentrations, wind-induced resuspension is considered to include the effects of wave action. In addition, the area around the shallow station is characterized by relatively small depths. Therefore, at the shallow station, the short wind-memory intervals (12–18 hours) yield the strongest correlations. Lagrangian simulations support this interpretation by illustrating that the shallow station, located within the intertidal zone itself, receives SPM directly from resuspension at the site. While these simulations do not represent full sediment dynamics, they effectively illustrate dominant source areas and transport timescales. This relatively short response time aligns with earlier observations in tidal flat systems, where wind effects on SPM are immediate in intertidal zones but take longer to become apparent in deeper areas (de Jonge and van Beusekom, 1995). In contrast, the situation at the deep station is more complex. In addition to the larger distance between the observation point and the sea bottom, sediment characteristics in the channels are predominantly sandy, which typically requires stronger wind energy to be resuspended as opposed to the mud and finer material found in intertidal areas (Bartholomä et al., 2009). The delayed response seen in the wind memory analysis (~120 hours) and the tidally modulated transport pathways from our Lagrangian simulations (median transit time ~133 hours) likely reflects the combined influence of episodic wind-driven resuspension in shallow areas and subsequent advective transport to the deeper station (Friedrichs and Pery, 2001; de Jonge and van Beusekom, 1995). Similar multi-day lags in SPM response have been observed in other estuarine systems where tidally dominated transport pathways act

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on resuspended sediments originating from more energetic, nearshore zones (see also Winterwerp, 2001). Together, these results highlight a clear contrast between the two stations: rapid, localized sediment response in the shallow embayment versus slower, more integrated transport processes at the relatively deep channel. The Lagrangian tracer simulations provide additional spatial context, revealing weak connectivity between the north-eastern and southern sections of the Sylt-Rømø Bight under regular tidal conditions. Passive tracers released in the north-eastern region rarely reach either of the sampling stations. This indicates that the stations, while valuable for long-term monitoring, may not fully capture the spatial variability of SPM dynamics across the entire basin, especially on shorter time scales. Beyond local wind and tidal resuspension and transport within the Sylt-Rømø Bight, tidal phases also influence SPM concentrations. Naturally, higher SPM concentrations are associated with lower water levels under otherwise equal conditions, as the measurements are always taken at the surface (Fig. S3). Due to the large role of non-linear processes in the area there is also an asymmetry between flood and ebb in terms of duration as well as mean and maximum velocities.

4.3 Neural Network findings

The NN experiments provide a line of evidence supporting the seasonal shift in dominant SPM drivers independent of the statistical analysis. When trained only on winter data, where biological activity is minimal, the NN successfully captured winter SPM concentrations using solely physical drivers such as wind conditions within different time intervals, salinity, and tidal elevation (through SSH and SSH gradient). At both stations, this winter model performed well ($R \approx 0.81$), confirming that physical processes alone can account for most of the observed variability during this season. This reinforces the view that winter dynamics are dominated by abiotic drivers. However, when the same winter-trained model was applied to other seasons, it consistently overestimated SPM concentrations, especially in summer when biological activity is high and wind speed is generally weaker. For example, at the deep station, the observed mean SPM concentration decreased by $\sim 78\%$, whereas the model reproduced only a $\sim 36\%$ reduction, meaning that less than half of the observed seasonal decline could be explained by abiotic factors. Similarly, at the shallow station, observed SPM declined by 80% , while the model captured only a $\sim 42\%$ reduction, accounting for just over half of the change. These mismatches underscore the increasing importance of biological drivers during the summer months. Interestingly, although both stations showed similar magnitudes of overestimation, the shallow station retained higher predictive skill ($R \approx 0.35$) than the deep station ($R \approx 0.04$) in summer. This difference likely reflects the stronger and more immediate influence of wind forcing at the shallow site, where even in summer, intermittent wind events can rapidly mobilize sediments. The addition of biologically relevant features such as temperature and sunshine duration into the full-year NN model, in Section 3.3.4, significantly improved its performance, especially at the shallow station. While we do not explicitly resolve individual biological pathways, these proxies likely capture their cumulative effects as seen from the improved model fit indicating their aggregate influence is both detectable and substantial. These processes may include phytoplankton-driven aggregation and flocculation, microbial stabilization, and benthic-pelagic interactions such as benthic filtration. Our

692 interpretation remains exploratory rather than conclusive, as targeted biological data would be required
693 to quantify these mechanisms in detail.

694 4.4 Study Limitations

695 Although, in this study, we discuss the main driving mechanisms of SPM concentrations in the Sylt-
696 Rømø Bight, we also acknowledge that the system is far more complex and has more factors
697 influencing the fluctuations of SPM concentrations than we investigate here. Moreover, the SPM itself
698 is a parameter that is composed of various components and sizes in nature, and such a level of detail is
699 not readily available in the dataset. Incorporating measurements of sediment composition (organic
700 versus inorganic fractions) and their sizes would add more certainty into the roles of considered
701 mechanisms modifying SPM concentrations.

702 Our Lagrangian transport simulations use massless passive tracers, which do not account
703 for flocculation, deposition, or resuspension processes. Incorporating such processes would require a
704 different modeling framework and additional data that are not currently available, including detailed
705 habitat maps and reliable estimates of SPM fluxes from the open boundary. Nonetheless, despite these
706 simplifications, the model effectively captures key patterns of spatial connectivity pathways and
707 timescales driven by tidal dynamics.

708 Another limitation is that changes in the SPM input from the open boundary cannot be quantified. Net
709 coastward transport from the North Sea likely contributes to seasonal variability, for example, through
710 enhanced import or retention in spring and summer (Burchard et al., 2008; Postma, 1981), but we lack
711 data to resolve this numerically.

712 In addition, more precise SPM predictions would benefit from neural network architectures with
713 “memory”, such as Long Short-Term Memory (LSTM) models, which can account for the delayed
714 effects of biotic and abiotic conditions over several months. A North Sea-wide approach would be
715 required to capture these long-term dynamics and boundary-driven influences. Within such a
716 framework, it would also be appropriate to include additional predictors such as nutrient concentrations
717 and benthic processes to represent better the complex interactions driving SPM variability.

718 While this study does not claim to offer a comprehensive representation of all the processes in play, we
719 believe that our work presents new insights to understand better the baseline mechanisms of SPM
720 concentration variability within the basin and across multiple timescales.

721 5 Conclusions

722 This study investigated the primary drivers of SPM variability in the Sylt-Rømø Bight, a semi enclosed
723 basin with well-mixed conditions and hydrodynamics dominantly shaped by tides. We combined long-
724 term monitoring data (SPM, Chl-a, wind conditions, and light availability) with SSH reconstructed from
725 a validated hydrodynamic model, and applied statistical analyses and neural network modelling. In
726 addition, Lagrangian transport simulations were used to assess tidal connectivity and transport
727 timescales. Together, these approaches revealed new insights on SPM concentration variability across
728 multiple timescales and relative influence of the main driving mechanisms.

Our findings confirm that wind speed is the primary driver of short-term SPM variability, particularly at the shallow station, where SPM concentrations respond almost immediately to wind forcing. This rapid response, especially during winter and autumn, highlights the importance of wind-driven resuspension in shallow waters. In contrast, the deep station exhibits a more delayed response to wind forcing, with peak correlations occurring at longer wind memory intervals (~5 days). This lag reflects the fact that, at greater depths, direct resuspension due to immediate wind forcing plays a reduced role, while the transport of material from neighboring shallower areas becomes increasingly important. Tidal dynamics primarily regulate the advection processes within the basin, redistributing fine, easily resuspendable material from shallow to the deeper areas. Lagrangian simulations illustrate that SPM at the shallow station originates locally, predominantly from within or around the Königshafen embayment. Meanwhile, at the deep station, SPM is likely supplied from the intertidal and shallow areas by the tidally driven redistribution over a longer timescale (~133 hours), consistent with the observed wind-memory lag (~120 hours). These results highlight the fundamental distinction between localized, wind-driven resuspension and slower, broader-scale, tide-driven transport, both of which shape SPM variability but at different spatial and temporal scales.

On seasonal timescales, however, biological processes exert an increasingly strong control. The onset of the spring phytoplankton bloom coincides with a decline in SPM concentrations, likely due to the enhanced production of TEPs and other extracellular polymers that promote biological aggregation and flocculation, leading to enhanced particle settling. This strengthening of biological effects reduces the efficiency of physical resuspension even under comparable wind conditions, which results in the pronounced decline in SPM concentrations toward summer. The inverse seasonal patterns of Chl-a and SPM support this interpretation, aligning with previous studies that describe the role of phytoplankton in promoting flocculation and sedimentation in coastal systems. NN experiments suggest that calmer wind conditions alone can explain approximately ~40% of the observed summer SPM reduction compared to winter concentrations. Still, they do not account for up to ~80% decrease seen in the data. This substantial reduction is likely influenced by a variety of biologically related mechanisms, ranging from the microbial activity, production of EPS to zooplankton grazing. Further studies are needed to quantify the relative contributions of these individual mechanisms.

Overall, this study provides a comprehensive and quantitative assessment of how wind, tides, and biological activity interact to control SPM variability in a shallow, tidally dominated coastal system.

Deleted: dominant

Deleted: Seasonal analyses further emphasize the shifting balance between physical and biological controls in shaping SPM dynamics. While wind and tides dominate winter SPM variability, the

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Conflict of Interest

The authors declare that they have no conflict of interest.

Author Contributions

GK performed the data analysis and wrote the initial draft of the manuscript. GK, VS, and AA carried out the numerical simulations. JvB, GK, and VS conceptualized the study. SH, SR, IK, and KHW contributed to the discussion of methods and interpretation of the results. All authors contributed to manuscript review and editing.

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776 **Data Availability**

777 The source code of the FESOM-C model is publicly available via Zenodo:
778 <https://doi.org/10.5281/zenodo.2085177> (Androsov et al., 2018). The Sylt Roads observational dataset
779 is accessible through the PANGAEA data portal: <https://doi.org/10.1594/PANGAEA.873549>,
780 <https://doi.org/10.1594/PANGAEA.873547>, <https://doi.org/10.1594/PANGAEA.918018>,
781 <https://doi.org/10.1594/PANGAEA.918032>, <https://doi.org/10.1594/PANGAEA.918027>,
782 <https://doi.org/10.1594/PANGAEA.918023>, <https://doi.org/10.1594/PANGAEA.918033>,
783 <https://doi.org/10.1594/PANGAEA.918028>, <https://doi.org/10.1594/PANGAEA.918024>,
784 <https://doi.org/10.1594/PANGAEA.918034>, <https://doi.org/10.1594/PANGAEA.918029>,
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786 <https://doi.org/10.1594/PANGAEA.918030>, <https://doi.org/10.1594/PANGAEA.918026>,
787 <https://doi.org/10.1594/PANGAEA.918036>, and <https://doi.org/10.1594/PANGAEA.918031>.
788 Meteorological data, including hourly wind characteristics (station 3032, List auf Sylt; dataset ID: *urn:x-*
789 *wmo:md:de.dwd.cdc::obsgermany-climate-hourly-wind*) and daily sunshine duration (*urn:wmo:md:de-*
790 *dwd-cdc:obsgermany-climate-daily-kl*), are available from the Climate Data Center (CDC) of the
791 Deutscher Wetterdienst (DWD): https://opendata.dwd.de/climate_environment/. The Lagrangian model
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