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How does the resuspension of the biofilm alter the functioning of the
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39 Abstract

40 Intertidal mudflats are ecosystems submitted to natural hydrodynamical forcings during each tide. When the offshore water flows at high tide, a proportion of the biofilm produced at low 41 tide can be resuspended in the water column and interact with the pelagic food web. As a 42 consequence, the resuspension creates a link between the benthos and the pelagos, modifying 43 their properties and the stability of the meta-ecosystem they form together. The aim of this 44 study is to describe the consequences of the microbial biofilm resuspension on the pelagic 45 food web, and to investigate the question of the stability of the benthos-pelagos coupling 46 resulting from the biofilm resuspension. Two food webs were considered, corresponding to 47 48 different hydrodynamical conditions in summer condition: one allowing the biofilm massive resuspension, and one without resuspension, but with particle sedimentation. The Monte-49 Carlo Markov Chain Linear Modelling was used to estimate the unknown flows of the food 50 51 web. The comparison of the Ecological Network Analysis indices for the two food webs allowed defining their respective differences of structure and functioning. The results showed 52 that the massive resuspension of the microbial biofilm stimulates pelagic primary production 53 and microbial food web via a higher bacterivory. The higher activity of the whole system 54 coupled with both a drop in the specialization of the trophic pathways and a low cycling 55 56 activity demonstrated that when massive resuspension occurs, the system is disturbed. In contrast, when sedimentation occurs, the food webs show functioning features pointing out to 57 a higher stability of the whole system. 58

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- 60 Keywords: intertidal mudflat, biofilm resuspension, stability, meta-ecosystem
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63 **1. Introduction**

The structure and the functioning of food webs affect the emergent properties and thus the 64 stability of the ecosystem. As a consequence, describing the emergent properties of the 65 ecosystem is a prerequisite for establishing their stability. The literature on the subject is 66 diverse and can lead to controversial interpretations and conclusions. However, some trends 67 can be observed such as equilibrium between two extremes that allows the ecosystem to act 68 like a buffer to external perturbations. For instance, the coexistence of weak and strong 69 interactions is assumed to bring stability to the ecosystem (McCann et al., 1998), or the 70 asymmetry in the ecosystem ensures a higher stability (Rooney et al., 2006), or an ecosystem 71 that presents a balance between Ascendency and redundancy is supposed to be more stable 72 (Ulanowicz, 2003). Moreover, Levin (1999) proposed that a stable ecosystem pattern is 73 composed of subsystems strongly intraconnected but weakly interconnected. At larger scale, 74 75 this theory is transposable to the concept of the meta-ecosystem defined by Loreau et al. (2003) as a set of ecosystems connected by spatial flows. A set of ecosystems strongly 76 77 intraconnected weakly interconnected thus form a stable meta-ecosystem. In this study, we propose to consider the benthos and the pelagos as systems connected by spatial flows at high 78 tide to form a stable meta-system. We have used this concept in order to describe and to 79 understand the effects of the benthos-pelagos coupling on the properties of the food webs and 80 its consequences on the stability of the Brouage mudflat meta-system. 81

As bare intertidal mudflat, Brouage mudflat, is characterized by the development of a microbial biofilm at the surface of the sediments at diurnal low tide. This biofilm is usually mainly composed of brown micro-algae (diatoms) which constitute the microphytobenthos (Cariou-Le Gall and Blanchard, 1995) and prokaryotes, all of them linked by a matrix of extracellular polymeric substances (EPS) (Decho, 2000). The production of the biofilm is integrated to the benthic food web via the deposit feeders, especially *Peringia ulvae* (Haubois

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et al., 2005; Pascal et al., 2008a; Pascal et al., 2009), via the facultative suspension feeders
and via the meiofauna (Pascal et al., 2009; Pascal et al., 2008b; Pascal et al., 2008c). The
meio- and macrofauna are not only involved in the regulation of the microphytobenthos
biomass via the grazing, but also via the bioturbation and the biostabilisation of the sediment
whose effects are coupled with physical factors (tides and swell) (Blanchard et al., 2001).

When the tidal flow arises, the microphytobenthos biomass decreases at the surface of the 93 sediments (Guarini et al., 2000) for two reasons: i) the downward 'migration' of motile 94 diatoms into the sediments (Consalvey et al., 2004; Guarini et al., 2000; Herlory et al., 2004; 95 Ni Longphuirt et al., 2009) and ii) the resuspension of a part of the diatom stock into the 96 97 water column. The resuspension of the microphytobenthic biofilm is controlled by a complex interaction between physical and biological forcing. The physical resuspension of the 98 microphytobenthos depends on the bed shear stress which is induced by the tidal current 99 100 and/or the wind-waves (Blanchard et al., 2002; De Jonge and Van Beuselom, 1992). The erodability of the sediment is strongly variable in space and in time (Tolhurst et al., 2006) and 101 102 depends on biological factor modifying sediment properties such as macrofauna activities and 103 microbial biofilm setting up (Herman et al., 2001; Orvain et al., 2004). The microphytobenthos resuspension also depends on biological factors such as the ageing of the 104 biofilm (Orvain et al., 2004) and the content of exopolysaccharids (EPS) (Orvain et al., 105 present issue). When the microphytobenthic biofilm is in its exponential growth phase, it 106 stabilizes the sediments and counteracts the bioturbation due to macrofauna which favors its 107 resuspension (Orvain et al., 2004). In contrast, when the biofilm reaches its senescent phase, 108 the roughness of the biofilm as well as bacterial biofilm degradation are enhanced and the mat 109 is more easily resuspended (Orvain et al., 2004). The microphytobenthic diatom that are 110 resuspended in the water column are integrated to the pelagic food web and can be ingested 111 by suspension feeders, in the case of the Brouage mudflat especially by Crassostrea gigas, a 112

cultivated species (Riera and Richard, 1996) and *Cerastoderma edule* (Sauriau and Kang,2000).

In this study new in situ observations and experimentations were taken into account, 115 especially on the resuspension of the biofilm (Orvain et al., present issue) and the 116 consequences on the pelagic food web are evaluated by using a model describing trophic 117 pathways. Erosion experiments allowed to determine the critical shear velocity necessary 118 regarding the resuspension of benthic micro-organisms and resuspension rates (Dupuy et al., 119 120 present issue). The Lagrangian and Eulerian field surveys followed the future of the resuspended particles in the water column, respectively following the water mass or at a fixed 121 point (Guizien et al., 2013). Moreover grazing experiments and viral lysis experiments 122 (Montanié et al., present issue) were performed in order to determine the effect of 123 resuspension of benthic organisms on the pelagic food web sensu largo (i.e. including virus). 124 125 These refinements were incorporated in the framework of a trophic-flow model to better unravel the impact of the biofilm resuspension on the food web functioning by deciphering 126 127 the contribution of each flow in the contribution in the functioning during high-tide phase. At high tide, the rise of the tide and the chemical/physical/biological processes associated with 128 the increase of the water level on the Brouage mudflat create spatial flows linking benthic and 129 pelagic parts. Consequently, it can be considered as a 'meta-ecosystem' defined by Loreau et 130 al., (2003) as a set of ecosystems which are linked by spatial biotic and/or abiotic flows across 131 the ecosystem boundaries. We especially focused on one question: How does the resuspension 132 of the microbial biofilm at high tide modifies the stability of ecosystem in respect to the meta-133 ecosystem pattern? We explored this question by comparing the network organization of two 134 distinct food web models representing 2 scenarii of a summer situation. In the first model 135 scenario, the hydrodynamic conditions were extreme and sufficient to induce the resuspension 136 of the microphytobenthos (physical mass erosion, when bed shear stress $BSS > 3 \text{ cm.s}^{-1}$ on 137

Brouage mudflat). In this case the velocity of current stays superior to the critical sinking 138 velocity, thus no sedimentation is possible. While in the second case the physical forcing was 139 too weak to induce any resuspension, only limited and extremely low erosion of chla 140 (biological erosion when $BSS < 3 \text{ cm.s}^{-1}$) induced by the bioturbation of the macrofauna can 141 be observed (Orvain et al., present issue). Moreover the settling of pelagic particles (organic 142 or not) could occur and the sinking velocity enhanced by the pelletisation (Orvain et al., 143 present issue). The missing flows of the food web (i.e. flows which were not measured *in situ*) 144 were estimated by the Monte Carlo Markov Chain Linear Inverse Modelling (MCMC-LIM) 145 (Van den Meersche et al., 2009). This mathematical method explores a solution space defined 146 by constraints issued from in situ measurement and values issued from the literature. All 147 solutions proposed by the MCMC-LIM were used to calculate several ecological network 148 analysis (ENA) indices, describing the emergent properties of the ecosystem. 149

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151 **2. Material and Methods**

152 2.1. The study area

The Brouage intertidal mudflat is located at the French Atlantic coast in the bay of Marennes-153 Oléron (figure 1). The bay covers 150 km² and the Brouage mudflat, at the eastern part of the 154 bay, represents 68 km² at low tide. The averaged bottom slope is relatively flat (1:1000) and 155 the tidal area is large (up to 4 km). The sediment consists of silt and clay particles (95% < 63156 μ m) (Pascal et al., 2009). The current speeds in the bay range from 0.2 to 0.6 m.s⁻¹ and the 157 bed shear stress from 1.5 to 4 Pa (Bassoullet et al., 2000; Le Hir et al., 2000). The zone of 158 interest is located in the middle of the Brouage mudflat and is characterized by a typical ridge 159 and runnel bedform (Gouleau et al., 2000). 160

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162 2.2. Inverse modelling

Two summer food web models were performed: the first one simulated a high-tide situation with massive suspension of micro-organisms in the water column (resuspension model), and the second one also at high tide, where the physical forcing was not sufficient to induce resuspension (sedimentation model). As a consequence, an insignificant quantity of particles is suspended via biological factors as macrofauna bioturbation and it counterparts by a strong sedimentation of organic matter.

The inverse modelling can be divided into 3 steps: (i) determine the species that compose the food web, and all possible flows between them. Twenty-one compartments were listed (Table 1) linked by 115 or 118 flows for the models with and without resuspension, respectively. (ii) determine the mass balance of each compartment and constrains flow values by *in situ* measurements. (iii)limite possible values of flows by biological constraints.

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2.2.1. Considered compartments and flows

176 *2.2.1.1. Resuspension and sedimentation*

A portable erodimeter (Guizien et al., 2012) was used to estimate the erosion shear stress of *in situ* cohesive sediments. An increased shear stress (by small steps, each timed to last about 10 minutes) was applied to the mud surface and the suspension of micro-organisms inhabiting in the sediment was monitored by changes in water column micro-organisms concentration. From these experiments, it was defined that resuspension of the microorganisms and diatoms took place when the shear bed velocity was higher or equal to 3 cm.s⁻¹. Sedimentation could not occur at higher current velocity.

Sedimentation was only considered in the model without resuspension, because the lower current velocity (< 3 cm.s⁻¹) allowed particles to settle down on the bottom during the slack water. The sedimentation rate was estimated from the formula D=Ws*C where D is expressed in mgC.m⁻².h⁻¹, C is the concentration of particles in the water column (mgC.m⁻³) and Ws the

sinking velocity of particles (m.h⁻¹) (Krone, 1962). Concentration of both pelagic bacteria and 188 189 chlorophyll a were measured in situ. The minimal limit for the concentration of the particulate carbon corresponded to the pelagic particulate carbon produced during high tide. For defining 190 its maximal limit we considered that the import of carbon into the water column was equal to 191 the amount of the benthic particulate carbon resuspended in the water column and not 192 consumed. Consequently, it was assumed that the particulate carbon present in the water 193 194 column at high tide, when resuspension occurred was likely to be found in a similar quantity 195 at high tide n+1.

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2.2.1.2. The microphytobenthos

Primary production of the microphytobenthos is linked to the biomass of diatoms present in 198 the biofilm and to light conditions (Macintyre et al., 1996). During immersion, the turbidity of 199 200 the overlying water, stopping the penetration of light (Alpine and Cloern, 1988) inhibits benthic primary production (Denis and Desreumaux, 2009; Migné et al., 2009). Moreover, 201 202 just before the flood return, the diatoms move down into the sediment (Herlory et al., 2004; 203 Round and Palmer, 1966). In our models, no microphytobenthic production was considered. The only input to the microphytobenthos compartment is thus an import of carbon which 204 comes from the production of the previous diurnal low tide by the microphytobenthos, 205 206 andwhich was not consumed during low tide. The import of carbon to the high tide corresponded to the export of carbon determined for microphytobenthos in a low tide model 207 and was equal to 31.9 mgC.m⁻² per high tide(Saint-Béat et al., 2013). Moreover the secretion 208 209 of EPS, related to the activity of photosynthesis and migration (Underwood and Paterson, 2003), was supposed negligible at high tide (Hanlon et al., 2006). This was confirmed by a 210 survey of EPS concentration in a tidal mesocosm during a tidal cycle showing a fall of EPS 211 concentration in the biofilm during high tide (Agogué et al., present issue) 212

9.7mgC of benthic diatoms were resuspended per high tide (Dupuy et al., present issue); they
constitute a flow from the benthos to the pelagos that enhances the phytoplanktonic biomass
after resuspension (Brito et al., 2012; Koh et al., 2006). In this way, the microphytobenthos
constitutes a food resource for the secondary producers in pelagic and benthic ecosystems
(Guarini et al., 1998; Riera and Richard, 1996; Yoshino et al., 2012).

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2.2.1.3. Benthic bacteria

Biofilms (i.e an assemblage of benthic diatoms and bacteria) were reconstituted in a tidal 220 mesocosm: the production and the biomass of the bacteria were measured during 5 days at 221 low and high tides (Agogué et al., present issue). . The benthic bacterial production was 222 estimated by tritiated thymidine incorporation (Garet and Moriarty, 1996), whichwas 223 converted to numbers of cells using the ratio of 1.96*10¹⁷ cells per mol of thymidine 224 225 determined for this study. A mean increase of 24.3% of the bacterial production in comparison to the bacterial production at low tide was observed in mesocosm experiments. 226 The bacterial biomass was obtained from the mean cell volume calculated with Furhman's 227 formula (1981) and converted in carbon units from the formula 133.754*V^{0.438}(Vin µm³) 228 (Romanova and Sazhin, 2010). The carbon contain in a bacterium was thus estimated as equal 229 to 79 fg C. cell⁻¹ for a mean biovolume of 0.28 µm³. During erosion experiments at the 230 threshold velocity considered in this study, 58.2 mgC per high tide issued from the benthic 231 bacteria were suspended (Dupuy et al., present issue). A part of the suspended bacteria were 232 damaged or dead during the erosion process (40%) (Mallet et al., present issue), and thus they 233 integrated the pelagic particulate organic carbon compartment. On the contrary the remaining 234 suspended benthic bacteria, still active, were considered to increase the pelagic bacteria 235 biomass (Mallet et al., present issue). 236

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2.2.1.4. Infauna (meiofauna and macrofauna)

The microphytobenthos is the preferential resource of the benthic fauna but in case of unavailability of microphytobenthos, bacterivory becomes significant (Pascal et al., 2009). At high tide, bacterivory was supposed to be higher than the low tide values: the values of low tide used in the previous model (Saint-Béat et al., 2013) were thus integrated to high tide models as minimal values. The obligate as well as facultative suspension feeders were assumed to feed on particles from bacteria to mesozooplankton (Self and Jumars, 1988; Taghon, 1982).

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2.2.1.5. Phytoplankton

The primary production of the phytoplankton was estimated for six other sites in Marennes-248 Oléron Bay based on *in situ* measurement of chlorophyll *a* water concentration, salinity, 249 250 temperature and depth profiles of light attenuation (Struski and Bacher, 2006). The phytoplanktonic primary production is influenced by the erosion of sediment, limited light 251 penetration and by the resuspension of benthic diatoms which likely participate to the 252 phytoplanktonic production (Macintyre et al., 1996). In order to prevent any bias in the 253 estimation of the planktonic primary production, we constrained it by minimal and maximal 254 primary production values measured in summer for the different stations of the bay. These 255 stations are characterized by different hydrological conditions and are thus characterized by 256 different resuspension and turbidity, and consequently by a different light penetration. 257 Constraining the phytoplanktonic production by a range of possible values allowed to adjust 258 the probability density function for gross primary production according to the needs of the 259 food web based on the situation considered (i.e. with or without resuspension). 260

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2.2.1.6. Pelagic bacteria

The summer abundance and the summer production of the pelagic bacteria were measured in 263 situ (Ory et al., 2011). Bacteria were counted by epifluorescence microscopy after being fixed 264 with 0.02µm filtered formaldehyde (2% final concentration) and staining for 30 min with 265 Sybr Green I (for more details see Noble and Fuhrman, 1998; Ory et al., 2011). The 266 measurement of AMPase (V_{max}), which is considered as an indicator of the bacterial 267 production, was used to estimate the bacterial production from the equation $\log BP =$ 268 $0.9271*\log \text{Vmax} + 5.3641$ (r²=0.67, n=10, p=0.003). Pelagic bacteria being assumed to 269 contain 16 fgC per cell (Labry et al., 2002), the bacterial production was thus expressed in 270 carbon. 271

Experiments on viral lysis were conducted to test the interactions between bacteria, 272 heterotrophic nanoflagellates (HNF) and viruses in the water column. In artificial incubations, 273 they were re-combined to mimic the field assemblage with respect to the natural viral to 274 275 bacterial ratio (VRB) and the bacteria to flagellates ratios. Fractionation allowed creating experimental recombined treatments and then differentiating between the direct and indirect 276 277 interactions of the presence/absence of the virus and HNF. Through in vitro experiments, bacterial losses induced by viruses and flagellates were estimated by comparing the reduction 278 of the bacteria cell production (MBP) between the different experimental assemblages. 279

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2.2.1.7. Benthic and pelagic viruses

Viral lysis was estimated from viral production (i.e. net increase of viral abundance divided by the time of the experiment) within 2 L bottle incubations in the presence or absence of benthic particulates in order to determine the effect of the biofilm resuspension on the viral lysis. Bacterial mortality due to viral lysis was calculated from the viral production divided by the burst-size viruses (i.e. number of viruses produced by a bacterium at burst-time) which was estimated as 33 in this study (Montanié, pers. comm.). The quantity of viruses produced per time unit was converted into carbon considering that one virus contains 0.2 fgC
(Magagnini et al., 2007; Suttle, 2005).

The viral lysis of benthic bacteria at high tide was considered to be similar to the one at low tide, thus we considered that 40 % of the bacterial production was lost by viral lysis (Saint-Béat et al., 2013). At high tide, the benthic viruses were suspended and integrated into the pelagic virus compartment. 1.29 mgC per high tide of virus per m² were resuspended in the water column at the critical shear bed velocity of 3cm.s⁻¹ (Dupuy et al., present issue) determined by the erosion experiment (see above).

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2.2.1.8. Ciliates and heterotrophic Nanoflagellates

The biomasses were expressed in carbon from a conversion factor of 19 μ gC. μ m-³ (Putt and Stoecker, 1989) by considering equivalent spherical diameter (ESD). The abundances and biomass of Ciliates (ESD < 50 μ m) and heterotrophic nanoflagellates (2 μ m < ESD < 10 μ m) were estimated *in situ*. The seawater was filtered onto 0.8 μ m Nucleopore black filter. The ciliates and nanoflagellates were enumerated by epifluorescence microscopy. Cells were first fixed by the glutaraldehyde and the paraformaldehyde and stained with the lugol 1% and DAPI respectively.

In the models, phytoplankton and pelagic bacteria are prey of the compartments ciliates and nanoflagellates, which are themselves the preys of ciliates. The nanoflagellates also potentially graze the viruses (Bettarel et al., 2005; Manage et al., 2002).

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2.2.1.9. Mesozooplankton

Mesozooplankton ($200\mu m < ESD < 400\mu m$) play a central role in the pelagic food web in the Marennes-Oléron Bay (Sautour and Castel, 1998) and show a variable diet (Vincent and Hartmann, 2001). The resources of mesozooplankton, as confirmed by bottle grazer experiments, in the area of Brouage mudflat are phytoplankton, either directly or indirectly
via ciliates and the heterotrophic nanoflagellates (Azémar et al., 2007). This diet can be
completed by the ingestion of detritic matter (David et al., 2006).

316 The abundance and the biomass of the mesozooplankton were estimated during study period. The mesozooplankton was sampled using a 200µm mesh WP2 net, preserved in buffered 317 formaldehyde with Na(Bo₃)₄ (final concentration 5%) and counted under binocular 318 microscope. The biomasses were expressed in carbon by multiplying the dry weight by 0.4 319 (Simard et al., 1985). These measurements were completed by bottle grazer experiments to 320 test the effects of the biofilm suspension on the grazing of mesozooplankton. Water from the 321 erodimeter (after erosion experiments) was mixed with filtered pelagic water (200 µm, in 322 order to exclude mesozooplankton) in different proportions (20, 40 or 70%). This mix was 323 transferred in 1.13 L Nalgene bottles and and incubated during 12 or 24 h in presence/absence 324 (temoin) of mesozooplankton predators collected in situ. The resuspended biofilm in the water 325 disturbs the trophic interactions of the mesozooplankton, especially through an inhibition of 326 327 the grazing of phytoplankton and heterotrophic nanoflagellates by the mesozooplankton (Hartmann, pers. comm.). Thus in the model with the resuspension no flow between 328 phytoplankton and nanoflagellates towards mesozooplankton was considered. 329

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2.2.1.10. Grazing Fishes

Main species able to graze on mudflat at high tide are mullets (*Liza ramada* and *Liza aurata*). Observed individuals arrive on the mudflat with an empty stomach, while they leave it with a full stomach (Carpentier et al., present issue). Thus the mullets were considered as a vector of carbon export. Since, the abundance of individuals going about the Brouage mudflat at high tide could not be measured *in situ*, grazing traces left by mullets on mudflat were considered as a proxy of their grazing pressure. Presence of traces was estimated from pictures of one square meter quadrats (expressed by surface of sediment removed by mullets by square meter). In addition, experiments on the feeding behavior of mullets were conducted in mesocoms to assess the volume of sediment ingested per individual at each tide (Como et al., present issue)). The coupling with field pictures finally allowed estimating the density of fishes per square meter.

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2.2.1.11. Imports and exports

For all benthic compartments, we considered that the production during the previous low tide 345 was not totally consumed, thus imports of carbon from the diurnal low tide were taken into 346 account. These import values corresponded to the mean export values of the low tide model 347 (Saint-Béat et al., 2013). Export was considered for both models regarding the 348 microphytobenthos and the macrofauna, while export was considered only in the model 349 350 without resuspension regarding the benthic particulate carbon and the dissolved particulate carbon. For the pelagic compartment, no import from the open sea was considered in the 351 model with resuspension and an import from the open sea of pelagic particulate carbon was 352 considered in the model without resuspension. We supposed that the production of the 353 pelagos was totally consumed during the high tide, thus no export from the bay to the open 354 sea of carbon was considered except for the pelagic particulate carbon, when the suspension 355 occurred. 356

- 357
- 358 2.2.2. Equations

The second step characterizes the mass balances of each compartment (listed in Table S1) and flows measured in the field. These two elements (*i.e.* mass balance and equations) were written within an equation: A * x = b where x was vector that contained possible flows, the matrix A expressed the mass balance and the field observation as a combination of

coefficients of the carbon flows and the vector b contained value of mass balances and values 363 of known flows (Vézina, 1989). The mass balances correspond to the report of inputs and 364 outputs for each compartment of the food web. By default, a compartment is considered to be 365 at the equilibrium (*i.e.* a constant biomass). Concerning the model with suspension we needed 366 to consider standing stock of the benthic particulate carbon and the benthic dissolved carbon 367 in deficit. Indeed, for these two compartments, there was a net change in mass equal to 368 resuspension term for particulate carbon and equal to the minimal value necessary to the 369 running of model for the dissolved organic carbon. In the case without resuspension, we 370 considered that the biomass loss of pelagic bacteria was equal to the value of the 371 372 sedimentation. The other sets of equations corresponded to the values of flows, which were 373 measured in situ.

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2.2.3. Inequalities

At the last step, some biological constraints were added to the mass balances and flow values. 376 These constraints were obtained from the literature and limit the possible solutions of flows to 377 realistic values. The information was added to the model with the inequality: $G * x \le h$, 378 where x remains the vector containing flows, G is a matrix that contains the coefficients of the 379 biological constraints and the vector h is composed of values of biological constraints 380 (Vézina, 1989). For the benthic compartments the set of inequalities of the low tide model 381 (Saint-Béat et al., 2013) was used. When the constraints corresponded to a value of flows, the 382 383 value of this constraint was updated according to the time of high tide (8h). The inequalities 384 for the pelagic compartments were grouped in the Table 2.

The sedimentation flows values were limited, considering two different sinking velocities, which is different according to the particle size (De La Rocha and Passow, 2007). The minimal sinking velocity corresponded to the sinking velocity of a single isolated particle. We considered a minimal sinking velocity of 0.05 m.d⁻¹ for free bacteria (Lapoussière et al., 2011), 0.25 m.d⁻¹ for chlorophyll *a* (Lapoussière et al., 2011) and 2.32 m.d⁻¹ for particulate carbon (Burns and Rosa, 1980). The maximal sinking velocity referred to the sinking velocity of 'marine snow', that are defined as organic aggregates with a diameter > 0.5 mm (Alldredge and Silver, 1988). The maximum sinking velocity chosen for this study was 16 m.d⁻¹ (Turner, 2002). We considered that pelagic dissolved carbon can fall with the aggregates formed by 'marine snow' and represents one third of the total carbon in aggregates (Alldredge, 2000).

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- 396

2.2.4. Calculation of solutions

The generated matrices (A, b, G and h) define a multi-dimensional space delimiting possible 397 solutions of the flows (x). The MCMC-LIM mirror (Van den Meersche et al., 2009) was used 398 to sample through that solution space in an attempt to map it completely. The MCMC-LIM, 399 400 based on the mirror technique defined by Van Den Meersche et al.(2009) and updated for LIM use by van Oevelen et al. (2010), calculates several solutions and allows a direct 401 402 characterization of the uncertainty. This modelling technique brings the advantage of 403 calculating a range of possible values for each flow (i.e. a probability density function). For each model (with resuspension and without resuspension), 500, 000 iterations with a jump of 404 0.5 were calculated. The length of jump and the number of iterations were determined to 405 406 cover the solution space as completely as possible. In this study the simulations were realized with a MATLAB[®] translation conceived by Alain Vézina and Lauriane Campo of the R-407 CRAN project package LIM-Solve created by Van den Meersche et al. (2009). 408

409

410

2.2.5. Network analysis

411 From the 500,000 solutions estimated by MCMC-LIM, seven ecological network analysis412 (ENA) indices were calculated. These indices allow assessing the structure and the

functioning of the two food webs. The magnitude of cycling within the system was described 413 by the Finn cycling Index (i.e. FCI). This index represents the fraction of flows involved in 414 the cycling (Finn, 1976). A cycle represents a series of transfers between components in an 415 416 ecosystem beginning and ending in the same compartment without going through the same compartment twice. The FCI is estimated by the ratio Tc/TST, where TST is the total system 417 throughput (i.e. sum of all flows) and Tc the amount devoted to cycling. Various global 418 indices describe the developmental and organizational state of the ecosystem (Ulanowicz, 419 1986). The TST measures the activity of the whole ecosystem. The TST can be considered as 420 the total power generated within the system (Baird et al., 1998). The AMI value is indicative 421 of the specialization of flows in the network (Ulanowicz, 2004). The probability of flows 422 between two compartments increases with the AMI value, and thus with the specialization of 423 flows. The Ascendency (A) which represents the state of organization within the ecosystem 424 (Ulanowicz, 1986), is described as the product of the TST and the average mutual information 425 (*i.e.* AMI). The development capacity (DC) is defined as the upper limit of Ascendency. The 426 427 relative Ascendency is the ratio A/DC and estimates the proportion of the network that is organized and thus efficient. The (DC - A) difference estimates the inefficient part of the 428 network, corresponding to the overheads (*i.e.* overheads on imports, exports and dissipation) 429 and redundancy, that measures the uncertainty associated to the presence of multiple or 430 parallel pathways among the compartments (Ulanowicz and Norden, 1990). The internal 431 Ascendency (A_i) and internal development capacity (DC_i) refer to internal exchanges alone 432 and exclude the exogenous flows. 433

These indices were estimated using MATLAB[©] routine written by Carole Lebreton and
Markus Schartau (GKSS Research Centre, Geesthacht, Germany) to calculate the index value
for every solution estimated by the LIM-MCMC.

437

438 2.2.6 Statistical test

Since the distribution of the data did not follow a normal distribution a non-parametric test 439 was used. The significance of the differences between the indices calculated for both 440 networks with and without resuspension was controlled by the Wilcoxon test ($\alpha = 0.01$). The 441 tested hypothesis was that the two data sets were issued from a continuous distribution with 442 equal medians. Statistical tests is possible because using the LIM-MCMC technique on 443 500,000 solutions, 500,000 values of each ENA index were also calculated, as consequence 444 we can use statistical tests which are not usually possible in such a context of food web 445 modelling using static methods and at this level of functional diversity. 446

447

448 **3. Results**

449 3.1. Flow values

450 Some differences in flow values between the two conditions (i.e. with or without suspension) were observed (Table S2). The pelagic primary production was higher with resuspension. On 451 the whole, consumption rates remained the same irrespective of the condition, except for the 452 bacterivory of heterotrophic nanoflagellates (doubled with resuspension), the bacterivory of 453 nematodes (consumption without resuspension was 7 times higher than the value during 454 resuspension), as well as herbivory of deposit-feeders (about twice higher without 455 resuspension than with resuspension) and consumption on nematodes by grazing fishes that 456 both doubled without suspension. The exudation of DOC by benthic bacteria increased 457 without suspension contrary to the exudation of DOC by pelagic bacteria that was five times 458 higher during resuspension. The mortality of phytoplankton (*i.e.* phyTOppc) was higher when 459 resuspension occurred. The egestion of nematodes without suspension was twice the egestion 460 during resuspension. The export of carbon from benthic compartment was higher without 461 suspension. 462

463

464 3.2. Compartment activities

Significant differences appeared according to the condition considered (Figure 2). The benthic activity was stimulated by the sedimentation of micro-organisms of the water column. In contrast, the resuspension of micro-organisms inhabiting in the sediment stimulated the pelagic activity.

Several pelagic compartments were affected by the resuspension of micro-organisms. The phytoplankton (phy), the pelagic non-living compartments (*i.e.* particulate compartment (ppc) and dissolved organic carbon (pdc)) as well as the pelagic bacteria (bcp) had a higher activity in the case of resuspension. On the contrary the microphytobenthos, the benthic bacteria, the benthic non-living compartments were more active when sedimentation occurred.

474

475 3.3. Diet and consumption

Herbivory tended to represent a more important part in the pelagic system (Table 3). In contrast, bacterivory was higher in the benthic compartment. The resuspension of the microorganisms to the water column had an effect on bacterivory, herbivory and the ratio between them. In the pelagic part, the resuspension favored the bacterivory. On the contrary, the herbivory was favored in the case without resuspension. The herbivory and bacterivory in benthos were both higher without resuspension.

The ratios showed that the herbivory was dominant in the pelagos. Nevertheless, when resuspension occurred, the decline in the ratio (almost divided by 2) was due to a fall of herbivory as well as a rise in bacterivory. A fall in the ratio was observed regarding the benthos due to a decline in the bacterivory lower than in herbivory.

486 None of the compartment was affected by the resuspension (Figure 3). Conversely, the diet of487 nematodes was drastically altered during the resuspension phase (B). Whatever the

resuspension occurred or not, the contribution of the microphytobenthos as a food item for nematodes did not change (about 15%). On the contrary, benthic bacteria and detritus contributed to nematodes diet almost equally in the case without resuspension (40 and 30%, respectively), while it mainly shifted to benthic particulate (about 80%) in resuspension situation.

The detritus contributed only slightly to the diet of deposit feeders. This group fed mainly on microphytobenthos and benthic bacteria. The contribution of each of these two groups changed with the resuspension. While the consumption on the benthic bacteria remained the same between resuspension and sedimentation phases, it corresponded to a higher contribution to the deposit-feeder diet during resuspension.

The facultative suspension feeders fed on the planktonic and benthic species. The contribution of the phytoplankton to the diet of this group was higher when the resuspension occurred. Consequently, the microphytobenthos was merely consumed. In contrast, in the case without resuspension, the microphytobenthos contributed to 40% of the consumption and the phytoplankton for 20%. The contribution of pelagic and particulate carbon tended to be lower with the resuspension.

The resuspension tended to have some consequences on the diet of the heterotrophic 504 nanoflagellates and mesozooplankton (Figure 4). The diet of the heterotrophic nanoflagellates 505 was more diversified during resuspension because of the contribution of virus (vrp). The 506 contribution of pelagic bacteria (bcp) was three times higher in the case of resuspension. 507 Consequently the contribution of the phytoplankton (phy) decreased. The most affected 508 compartment was the mesozooplankton. Without resuspension the mesozooplankton fed 509 equally on the phytoplankton, heterotrophic nanoflagellates, pelagic particulate carbon (ppc) 510 511 and ciliates. During the resuspension, the consumption on both heterotrophic nanoflagellates and phytoplankton was inhibited. 512

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513

514 3.4. ENA indices

Significant differences between values of the indices of the two situations were observed 515 516 (Figure 5). The TST was about 3 times higher when the resuspension occurred. The Ascendency followed the same trend. However, a lower organization of the system (i.e. lower 517 AMI value) was observed. The higher relative Ascendency, coupled with a lower AMI value 518 519 observed in the case of resuspension, suggested a lower diversity of flows. The overheads 520 based on the imports, exports, respiration and redundancy of the system were lower during resuspension. The internal normalized Ascendency tended to be similar for the two situations. 521 The proportion of flows involved in the cycling (i.e. FCI value) was lower during 522 resuspension event. 523

524

525 **4. Discussion**

526 4.1. Activity of the benthic and the pelagic compartments

527 The benthos was more active than the pelagos regarding simulation without resuspension; in 528 contrast the pelagic activity was higher during resuspension. This fact can be explained by a large import of particulate carbon to the pelagic particulate compartment. Without this import 529 530 of pelagic carbon, the pelagic activity was equal to the benthic activity during resuspension. The higher benthic activity was previously shown in the model of the Brouage mudflat food 531 web. In fact, the higher activity of the benthos was observed irrespective of the model 532 considered in previous studies by regarding annual budget (Leguerrier et al., 2004) or by 533 deciphering seasonal budgets (Degré et al., 2006). However, in our model, the ranking of the 534 compartments was modified. The main difference concerned the benthic bacteria and the 535 microphytobenthos. The benthic bacteria dominated the ecosystem; they were followed by the 536 benthic non-living detritus compartments. Surprisingly, the microphytobenthos was ranked 537

only 8th. The difference with previous models is the time-scale considered: a mean year 538 539 (Leguerrier et al., 2004) or a mean month (Degré et al., 2006) which is in dramatic contrast to the small scale mean immersion period integrated in our model. During immersion, because 540 541 of darkness due to sediment burying, the microphytobenthos production was null (Blanchard, 2006). As a consequence, the carbon input to this compartment exclusively originated from 542 the photosynthesis performed during the previous diurnal low tide, which might explain its 8th 543 rank in our model. The first pelagic compartments were ranked 4th and 5th and corresponded 544 545 to the phytoplankton and the dissolved organic carbon, respectively.

The resuspension of the microbial biofilm stimulated the activity of the phytoplankton, the 546 non-living compartments (detritus) and the pelagic bacteria. This stimulation was not only the 547 consequence of the input of new matter in the water column. The phytoplankton showed a 548 higher gross primary production when the resuspension of the microbial biofilm occurred. 549 550 Pelagic primary production was not imposed to the model; it was only constrained by minimal and maximal values of pelagic production found for different hydrological conditions in 551 552 Marennes-Oléron Bay. The higher production was somewhat surprising because the resuspension also generates a high turbidity and the reduction of the light penetration which 553 dramatically reduces phytoplankton photosynthesis (Billerbeck et al., 2007; Porter et al., 554 555 2010). When buried in the sediments, the microphytobenthos can remove nutrients from the overlying water and the sediment pore water (Macintyre et al., 1996). Such activity tends to 556 reduce the nutrient fluxes from the sediments to the water column which can limit the 557 phytoplankton production (Sarker et al., 2009; Sloth et al., 1996). When the 558 microphytobenthos resuspension occurs, the nutrient fluxes to the water column increase 559 which has a positive effect on the phytoplankton production even if light is limiting (Porter et 560 561 al., 2010). Paradoxically, the import of microphytobenthic diatoms to the water column contributes itself positively to the production of the phytoplankton (Macintyre et al., 1996). 562

563 By enriching water in particulate and dissolved carbon, which sustained a higher bacterial 564 activity as previously reported (Cotner et al., 2000; Poremba et al., 1999; Sloth et al., 1996), 565 the resuspension also stimulated heterotrophic production.

566 On the contrary, the settling of pelagic micro-organisms to the bottom of the water column had smaller but significant consequences on the benthic compartments. In our model, the 567 input of pelagic fresh matter to the benthos increased the stock of available carbon for higher 568 trophic levels but it did not affect the production of the different compartments. The higher 569 570 activity observed for the benthic compartments (mpb, bdc, bcb, bpc) (Figure 2) was linked to the deposit of pelagic diatoms, dissolved carbon, pelagic bacteria and particulate carbon 571 respectively at the surface of the sediments. The photosynthetic production of the pelagic 572 diatoms settled at the bottom of the water column during immersion is more probably limited 573 by the penetration of light than by the nutrient availability. It is supposed that, in contrast to 574 575 the light reaching the surface of the sediments (Billerbeck et al., 2007; Macintyre et al., 1996), nutrients are often not a limiting factor for the photosynthesis in the intertidal sediments 576 577 (Migné et al., 2004; Serôdio and Catarino, 2000). The high turbidity reported in the bay of Marennes-Oléron (Raillard and Mesenguen, 1994) likely stops the light penetration to the 578 sediment surface and strongly impairs the photosynthesis of pelagic diatoms. The effect of the 579 settling on the benthic bacterial production is more obvious. The bacterial production depends 580 on the substrate supply including organic carbon (Sander and Kalff, 1993). For instance in 581 Kiel Bight, the settling of detritus stimulates the benthic bacterial production (Meyer-Reil, 582 1983). Consequently, it is very likely that the benthic bacterial production is stimulated in the 583 Brouage mudflat during sedimentation. 584

585

586 4.2. Herbivory *versus* bacterivory

The massive resuspension event impacted the pelagic microbial food web. It stimulated the 587 bacterivory of the pelagos as reported before for heterotrophic nanoflagellates (Garstecki et 588 al., 2002). A higher quantity of carbon flowed through the virus (viral lysis was doubled) and 589 590 consequently more virus were consumed by heterotrophic nanoflagellates. The increase of pelagic bacterial abundance and production doubled the bacterivory rate of heterotrophic 591 nanoflagellates. In contrast, the resuspension tended to decrease the pelagic herbivory. In spite 592 of a higher gross primary production and a direct input of benthic diatoms biomass to the 593 594 water column, the phytoplankton was integrated to a lower proportion to the pelagic food web. Indeed, the grazing of phytoplankton by the mesozooplankton had been shown to be 595 inhibited during catastrophic erosion event (Hartmann, pers.comm.). Consequently, our 596 models showed that a lower part of the phytoplankton was consumed in the model with 597 resuspension, inducing a greater mortality of this comportment. This result of the models (*i.e.* 598 599 higher phyTOppc when resuspension occurs) is coherent with previous results of resuspension experiments (Porter et al., 2010). It thus appears that in conditions of resuspension, the 600 601 phytoplankton indirectly participated to the enrichment of the water column in dissolved 602 organic carbon and to the pelagic bacterial production.

The bacterivory dominated the benthic compartment during immersion irrespective of the 603 604 resuspension. At high tide, benthic diatoms moves down the sediment and cannot perform photosynthesis because of the absence of light (Cartaxana et al., 2011). Hence, the benthic 605 food web must be sustained by the input of carbon coming from the photosynthetic 606 production of the previous emersion and by the bacterial production. In our model, we 607 supposed that the meiofauna and the macrofauna showed a constant consumption along the 608 day irrespective of immersion/emersion periods. This hypothesis was based on the assumption 609 610 that the meiofauna and the deposit feeders had alternative resources since the microphytobenthos was not sufficient to sustain their respective metabolism. Detritus (van 611

Oevelen et al., 2006) bacteria were possible alternative resource for benthic fauna (Pascal et al., 2009; van Oevelen et al., 2006). Some isotopic analyses at the Brouage mudflat showed that the benthic detritus contributes to 11% in maximum to the deposit-feeders consumption (Richard, comm. pers.). We supposed that the alternative feeding resource was the benthic bacteria (see result section .

Previous studies on the bacterivory rates of the nematodes, the foraminifera and Peringia 617 ulvae in the Brouage mudflat indeed showed that bacteria can constitute a significant 618 619 alternative resource to the microphytobenthos under some conditions (Pascal et al., 2008a; Pascal et al., 2008b; Pascal et al., 2008c). When there was no resuspension, the herbivory and 620 621 the bacterivory were both stimulated, although the bacterivory was stimulated in a larger proportion due to the higher activity of the nematodes. The bacterivory thus remained 622 dominant in the benthic compartment irrespective of the physical forcing. The Sylt- Rømø 623 624 Bight in the North of Germany is composed of a mosaic of habitats including a mudflat. Its benthic food web is based on microphytobenthos and macrophytes. In this benthic ecosystem, 625 the herbivory dominates the bacterivory (Baird et al., 2004). Indeed, herbivory is more than 626 627 two times higher than bacterivory. Thus the ratio herbivory/bacterivory of the Sylt-Rømø Bight displays an opposite tendency than the ratio estimated for the Brouage mudflat. This 628 629 opposite tendency may be linked to the fact that this ratio was obtained from the food web for the whole bay on a long term. As a consequence, habitats with high and low primary 630 production were associated, thus the available carbon issued from the primary production was 631 632 more important. Moreover, the food web considered in Baird et al. (2004) represents a mean day as a consequence the difference between low tide and high tide was not visible and the 633 effect of the season was not considered contrary to present simulations, which focused on 634 summer conditions and high tide only In this study, that considered the summer period, little 635 carbon issued from the primary production at low tide was available. The consideration of the 636

food web for the whole year at the Brouage mudflat should abate the seasonal difference andchange the trend of herbivory/bacterivory ratio.

639

640 4.3. Functioning of the Brouage food web

For comparing the functional indices from our model to others, we took care of selecting 641 ecosystem models that coupled the pelagic and benthic compartments. Moreover the non-642 643 living compartments must be separated from bacteria, otherwise the ENA indices would be biased (Johnson et al., 2009). Values of relative Ascendency and internal relative Ascendency 644 were in general higher to those previously reported. Relative Ascendency ranged from 33.4 645 (Monaco and Ulanowicz, 1997) to 49.5 for the Chesapeake Bay (Baird et al., 1991) and the 646 internal Ascendency from 31.2 for the Delaware (Monaco and Ulanowicz, 1997) to 44.1 for 647 the Sundays Bay (Scharler and Baird, 2005). The cycling estimated without resuspension was 648 649 higher to the FCI (i.e. Finn Cycling Index) estimated in the Sylt- Rømø Bight (= 17.2%) and close to the value of Sundays Estuary (Scharler and Baird, 2005). The differences between 650 651 our study and previous ones are most probably based on the shorter time scale in our models. In the aforementioned studies, the ecosystems considered are estuaries which are subjected to 652 the tidal rhythm and thus which are controlled by strong temporal/physical forcing. It was 653 demonstrated how physical parameters can influence the ecological properties described by 654 the ENA indices (Niquil et al., 2012). The consequences of physical forcing like the 655 resuspension are smoothed when the considered networks use flows averaged over a mean 656 day, more representative of normal conditions without waves. Additionally, the biological 657 processes change according to the immersion and the emersion periods which impacts the 658 carbon budget (Migné et al., 2009). Consequently, when emersion and immersion are 659 considered separately (i.e. short time scale), it allows more precisely deciphering the 660 biological and physical processes that control the functioning of the food web. 661

When the massive resuspension of the microbial biofilm occurred, the enrichment of the 662 water column by the benthic particulate carbon (i.e. 3.584gC.m⁻².per high tide) mainly 663 supported the increased activity of the whole system and the decreased organization of the 664 network decreased. The higher value of Ascendency (2*10⁴ mgC.m⁻².per high tide against 665 $0.6*10^4$ mgC.m⁻².per high tide during sedimentation) was the consequence of a higher TST 666 (10000 mgC.m⁻².per high tide during resuspension and 3500 mgC.m⁻².per high tide during 667 sedimentation), in spite of a drop down in the specialization of the trophic way (measured by 668 AMI)This is a characteristic observation for a so-called "pulse eutrophication", an intermittent 669 increase of organic matter supply combined with physical factors (Patrício et al., 2004). 670 Moreover a high value of Ascendency derived from a very high TST, could disturb the 671 internal stability of the system (Ulanowicz, 2003). The lower overheads suggest a lower 672 resistance to the perturbation as proposed by Ulanowicz (2003). Thus the resuspension 673 674 decreased the resistance of the system to a perturbation. We propose that the massive resuspension event in the Brouage mudflat could be defined as a "pulse eutrophication" event 675 676 that regularly disturbs the meta-ecosystem. On the contrary the biological erosion coupled with a high sedimentation tended to reduce the perturbation in the meta-system. 677

During the massive resuspension, the lower proportion of cycling (i.e. lower value of FCI) 678 coupled with a high internal relative Ascendency (close to 50%) can be explained by the 679 limited integration to the planktonic food web of the non-living carbon suspended in the water 680 column. Indeed, in spite of a higher pelagic bacterial production and a higher detritivory, a 681 low quantity of carbon was recycled. As a consequence, almost all the carbon suspended in 682 the water column (i.e. 3.534 gC.m⁻².per high tide) was exported. Note that the export value 683 was not constrained in the model, thus this value reflected a real property of the network. The 684 larger the difference between the cycling and the internal relative Ascendency, the less 685 organized and more under pressure a system would be (Baird et al., 2007; Baird et al., 1991). 686

Thus the Brouage mudflat was less organized (confirmed by lower value of AMI) and 687 submitted to a higher stress during the resuspension. A similar relation (i.e. great difference) 688 between FCI and internal relative Ascendency was found for the mussel-bed in the Rømø-Sylt 689 Bight (Baird et al., 2007) and in an upwelling area (Baird et al., 1991). Baird et al (1991) 690 brought a significant nuance to the stress sense, by the distinction between ecosystems which 691 692 are under physically or chemically pressure. Indeed, the two constraints do not refer to the 693 same time scale. A chemical stress is in general recent and it has an exogenous origin to the considered ecosystem. In contrast, physical perturbations are older and the ecosystem can 694 have adapted to it. These differences explain how a low cycling value can be coupled with a 695 high internal relative Ascendency (Baird et al., 1991). During the massive resuspension phase, 696 and as expected, the Brouage mudflat obviously showed the characteristics of a system that is 697 physically perturbed. 698

699 When massive resuspension did not occur, the Brouage mudflat was characterized by a high specialization (i.e. high AMI) and by a relative Ascendency close to 60%. Such value 700 701 illustrates a state closed to the equilibrium between the efficient and the fraction of the 702 network that has not yet been organized (Bodini and Bondavalli, 2002); it is based on redundancy in the imports, the exports, the dissipation and on internal redundancy (Baird et 703 al., 2004). The equilibrium between both parts (relative Ascendency and overheads) is 704 705 supposed to bring sustainability to the ecosystem (Ulanowicz et al., 2009); the inefficient part being used as a reserve that brings the necessary flexibility for the ecosystem sustainability. 706 Moreover the high internal relative Ascendency is a strong sign that the system is relatively 707 708 mature (Baird et al., 1991). The lower difference between internal relative Ascendency and FCI supposed a higher organization and a less disturbed system (Baird et al., 2007). Hence, 709 710 without massive resuspension of the microbial biofilm, the Brouage system seems to be relatively mature and stable. 711

712

4.4. Conclusion: the stability of the Brouage meta-system

As defined by Loreau (2003), a meta-ecosystem corresponds to the different ecosystems 714 715 which are linked together by spatial flows of energy and matter. The rise of the tide and the chemical/physical/biological processes, which are associated with the increase of the water 716 717 level on the mudflat can be considered as spatial flows. Here, we considered two different 718 events according to the hydrodynamical conditions: 1) the massive resuspension of benthic 719 matter in the water column 2) the sedimentation of pelagic matter on the mudflat sediments associated to a biological resuspension induced by macrofauna activities. As described above, 720 721 their respective impact on the functioning of the benthic and the pelagic food webs strongly differs. The massive resuspension tends to disturb the Brouage meta-system while the 722 sedimentation stabilizes it. These opposite consequences can be explained by the difference in 723 the intensity of the flows. When the massive resuspension occurs, the sum of flows from the 724 sediments to the water column was strong (about 3654 mgC.m⁻² per immersion) while during 725 726 the sedimentation, it was only 10% of the flow during resuspension. The interaction between 727 the benthic and the pelagic compartments also appeared weaker during sedimentation than during massive resuspension. As suggested by Levin (1999), a highly modular system 728 729 (composed of strongly connected sub-systems which are connected by weak links) is a stable system. This concept could be transposed to the meta-ecosystem. We observed that the 730 sedimentation constitutes a weak link between the two subsystems benthos and pelagos. In 731 732 contrast the massive resuspension constitutes a strong link between benthos and pelagos. To conclude the stabilizing pattern of Levin is observed when sedimentation occurs and not 733 during massive resuspension event. This conceptual step appears essential for the better 734 understanding of (meta-)ecosystem structure and functioning in order to improve our 735 prediction for their sustainability. 736

Despite its visible destabilizing effect, the massive resuspension brought some benefits to the 737 Brouage meta-system. It stimulated the pelagic microbial food web by increasing both 738 phytoplanktonic and bacterial production, and by stimulating bacterivory. Because of the 739 coupling of beneficial and destabilizing effects, massive resuspension show features 740 characteristic of an intermediate disturbance (reviewed in Shea et al., 2004). An intermediate 741 disturbance can be defined as an event that alters the specific niche availability, for instance 742 743 by removing the biomass or changing the nutrient availability, while it maintains the general biodiversity (Shea et al., 2004). A complementary and extensive study of the long term 744 massive resuspension frequency and its consequences on the Brouage meta-system would 745 allow to confirm the intermediate disturbance hypothesis. 746

747

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Benthos	
Microphytobenthos	mpb
Benthic bacteria	bcb
Meiofauna	mfb
Nematodes	nem
Deposit-feeders	dep
Suspension-feeders	sus
Facultative suspension-feeders	suf
Omnivorous species	omn
Carnivorous species	car
Benthic viruses	vrb
Benthic particulate carbon	bpc
Benthic dissolved carbon	bdc
Pelagos	
Phytoplankton	phy
pelagic bacteria	bcp
Heterotrophic nanoflagellates	hnf
Cilliates	cil
Mesozooplankton	mes
Grazing fishes	gfi
Pelagic viruses	vrp
Pelagic particulate carbon	ppc
Pelagic dissolved carbon	pdc

Processes	Compartments	Lower limit	Upper limit	References
Gross Growth Efficiency	HNF, CIL, MES	10%	40%	Straile, 1997
Net Growth Efficiency	BCB, BCP	11%	61%	DelGiorgio and Cole, 1998
	MFB	30%	50%	van Oevelen et al. 2006
	NEM	60%	90%	
	MAC	50%	70%	
Assimilation Efficiency (loss to the det)	HNF,CIL,MES	50%	90%	Vézina and Platt, 1988
	MFB	57%	97%	van Oevelen et al. 2006
	NEM	6%	30%	
	MAC	40%	75%	
	GFI	50%	90%	Leguerrier et al., 2004
Excretion (loss to doc)	HNF,CIL,MES	10% of ingestion	100% of respiration	min: Vézina and Pace, 1994 max: Vézina and Platt, 1988
	PHY	10%NPP	55%NPP	Breed et al., 2004
		5%GPP	50%GPP	Vézina and Platt, 1988
Respiration	HNF,CIL,MES	20% of ingestion	-	Breed et al., 2004
	PHY, MPB	5% GPP	30%GPP	Vézina and Platt, 1988
	MES	biomass*4.8*W ^{-0.25}	biomass*14*W ^{-0.25}	min: Hemmingsen, 1960
	CIL, HNF	biomass*0.6*W-0.25	biomass*1.7*W ⁻⁰²⁵	max: Moloney and Field, 1989
Consumption / Biomass	GFI	3%	8%	Bruslé, 1981
Gross primary production (mgC.m ⁻² .h ⁻¹)	РНҮ	10	50	Struski and Bacher, 2006
Loss of doc for pelagic bacteria (mgC.m-2.h-1)	BCP	0.012	-	in this study(with suspension)
		0.005	-	
Bacterivory by HNF	HNF	-	49% of the bacterial production	in this study (with suspension)
	HNF	-	45% of the bacterial production	in this study (without suspension)
Respiration (mgC.m-2.h-1)	GFI	0.226	3.628	min: derived from Killen et al., 2010 max: derived from Brett, 1965

Table 2: List of biologial constraints used for the food web model. NPP: Net Primary
Production, GPP: Gross Primary Production. W: body mass in pgC. Net Growth Efficiency =
(consumption-detritus production-respiration)/ (consumption-detritus production), Gross
Growth Efficiency=(consumption-loss to det-loss to doc-respiration)/ (consumption-loss to det-loss to doc- production).

	Benthic	Pelagic			
	With	Without	With	Without	
	resuspension	resuspension	resuspension	resuspension	
Herbivory (mgC.m-2.HT-1)	20.4 ± 0.1	39.7 ± 7.3	28.7 ± 11.2	33.8 ± 9.8	
Bacterivory (mgC.m-2.HT-1)	85.9 ± 5.8	123.7 ± 42.7	$14.3\ \pm 5.8$	8.1 ± 3.2	
Ratio herbivory/bacterivory	0.2 ± 0.01	0.35 ± 0.1	2.9 ± 3.1	5.6 ± 4.9	

1076	Table 3: Mean values of herbivory and bacterivory in the benthos and the pelagos according
1077	to the resuspension of the biofilm. HT^{-1} = per High Tide. These values correspond to the mean
1078	and the standard deviation calculated from the 500,000 iterations calculated by the inverse
1079	analyses. For each compartment (i.e. benthos and pelagos), values were significantly different
1080	with or without resuspension (Wilcoxon test, p<0.05) according to the condition considered.
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1083 Figures captions

Figure 1: Map of the study site: the Brouage mudflat within the Marennes-Oléron Bay.

Figure 2: Histogram presenting the activity of the compartments composing the food web. 1086 The activity corresponds to the sum of inflows and is expressed in mgC.m⁻² per high tide. A) 1087 corresponds to the sum of the activity of all compartments for the benthos and the pelagos 1088 according to the 2 modeled situations. B) and C) refer to the activity of each compartment of 1089 the benthos and of the pelagos. Compartments are sorted by top down activity in order to 1090 determine which ones of the compartments have higher contribution to the activity of the 1091 whole ecosystem according to the situation considered (i.e. sedimentation or resuspension). 1092 1093 See table 1 for the abbreviations of the compartments.

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Figure 3: Diet of the meiofauna and macrofauna. A) Top diagrams refer to the case without resuspension and B) down diagrams refer to the case where the resuspension occurred. The contribution corresponds to the fraction that represents the consumption on a prey in comparison to the total consumption. The contribution of each species to the compartment consumption was estimated from the mean value of each flow.

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Figure 4: Diet of heterotrophic nanoflagellates (hnf), ciliates (cil) and mesozooplankton (mes). A) top diagrams refer to the simulation without resuspension and B) down diagrams refer to the simulation with resuspension. The contribution corresponds to the fraction that represents the consumption on a specific compartment in comparison to the total diet. The contribution of each species to the compartment consumption was estimated from the mean value of each flow.

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Figure 5: Boxplots displaying the values of different ENA indices: the total System Throughput (TST), the Ascendency, the overheads, the relative Ascendency (A/DC), the Average mutual Information (AMI), the internal relative Ascendency (Ai/DCi) and the Finn Cycling index (FCI). The indices were calculated from the 500,000 solutions coming from the MCMC-LIM method. Red crosses correspond to outliers. Medians of all these indices were significantly different for the two seasons (Wilcoxon test, H₀ was rejected, p-value < 0.01).

1115 Figure 1















