1	Structure and functioning of the acid-base system in the Baltic Sea.
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10	constants, biomass production, mineralization

12 Abstract

The marine acid-base system is relatively well understood for oceanic waters. Its structure and 13 14 functioning is less obvious for the coastal and shelf seas due to a number of regionally specific anomalies. In this review article we collect and integrate existing knowledge on the acid-base 15 system in the Baltic Sea. Hydrographical and biogeochemical characteristics of the Baltic Sea, 16 17 as manifested in horizontal and vertical salinity gradients, permanent stratification of the water column, eutrophication, high organic matter concentrations and high anthropogenic pressure, 18 makes the acid-base system complex. We summarize in this study the general knowledge on 19 the marine acid-base system as well as describe the peculiarities identified and reported for the 20 21 Baltic Sea specifically. In this context we discuss issues such as: dissociation constants in brackish water, different chemical alkalinity models including contributions by organic acid-22 base systems, long term changes of total alkalinity, anomalies of borate alkalinity and the acid-23 base effects of biomass production and mineralization. Finally, we identify research gaps and 24 25 specify bottlenecks concerning the Baltic Sea acid-base system.

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1 **1. Introduction**

2 The acid-base system controlling the seawater pH is formed by a number of chemical substances having acid-base properties (Dickson et al., 2007). The importance of each of these 3 4 substances (structure of the acid-base system) depends on both concentrations of individual constituents and their dissociation constants. The control by physical conditions (temperature, 5 salinity) and biogeochemical processes (e.g. biomass production and mineralization) as well as 6 interrelationships between individual components determine the functioning of the acid-base 7 system. The understanding of the structure and functioning of the acid-base system is necessary 8 to investigate important issues that shape the Baltic Sea ecosystem and that are of interest in 9 present-day chemical oceanography like: ocean acidification, calcium carbonate (CaCO₃) 10 formation/dissolution and carbon dioxide (CO₂) exchange through the air-sea interface. 11

12 The steady increase of atmospheric CO₂ concentrations leads to enhanced dissolution of CO₂ in the ocean. Since CO₂ dissolved in seawater forms the diprotic carbonic acid, hydrogen ions 13 14 are released. Although the major fraction of the hydrogen ions are taken up by carbonate ions (a buffering reaction), a significant fraction stays in the water column and thus causes a 15 16 decrease in pH. This phenomenon is known in the scientific literature as "ocean acidification" although seawater does not really become acidic but only moves from its alkaline character 17 18 towards the acidic regime (Riebesell et al., 2010). Ocean acidification has been recognized as 19 one of the greatest threats for marine ecosystems not only by the scientific community (e.g. Bates et al., 2014; Zeebe, 2012), but also in European Union (EU) legislation. EU Marine 20 Strategy Framework Directive (MSFD, 2008) explicitly points out that the EU Member States 21 should put more attention to ocean acidification, and emphasizes the necessity to include 22 measurements of pH and of the CO₂ partial pressure (pCO₂) as descriptors for the 23 environmental status of marine regions. 24

There is a general belief that the magnitude of ocean acidification can sufficiently be quantified 25 26 from the atmospheric pCO₂ levels and the CO₂ exchange between seawater and the atmosphere (Zeebe, 2012; Riebesell et al., 2010; Caldeira and Wicket, 2003). This is approximately true 27 28 for oceanic waters. However, it is not the case for coastal seas because several other processes 29 are influencing the seawater pH such as: CaCO₃ formation and/or dissolution, euthrophication 30 or oligotrophication, total alkalinity (A_T) consumption and production, weathering, contribution by organic substances and terrestrial inputs. The general structure and functioning 31 32 of the acid-base system in seawater are relatively well identified, however still some aspects contain a lot of gaps and/or shortcomings, which can lead to wrong conclusions and 33

questionable predictions of the future pH development (Orr et al., 2015; Koeve and Oschlies,
 2012; Hunt et al., 2011; Dickson et al., 2007; Cai et al; 1998).

3 Since the CO_2 system is an integral part of the ocean acid-base system, it is impossible to 4 understand the CO₂ system and to assess processes such as the CO₂ gas exchange or CaCO₃ dissolution/formation, without a clear idea about the structure and functioning of the whole 5 acid-base system. The saturation states of calcite and aragonite are of crucial importance for 6 organisms forming their exoskeletons from CaCO₃. Although pelagic calcifying organisms do 7 not occur in the Baltic Sea this aspect is still relevant as some benthic organisms can also build 8 CaCO₃ into their shells (Tyrrell et al., 2008). Moreover, calcifying processes not only depend 9 on the acid-base system structure but in turn can modify it by decreasing the concentration of 10 carbonate ions (CO_3^{2-}) and thus the alkalinity. 11

Carbon dioxide is a major component of the global carbon cycle and transport of CO₂ links all 12 Earth's compartments. It is believed that the world oceans absorb about 25 % of anthropogenic 13 14 CO₂ emissions. However, there is still a debate on the role that the shelf seas play in this context (Le Quéré, 2016). The direction and magnitude of the CO₂ exchange through the air-sea 15 16 interface depends largely on the pCO_2 difference between seawater and the atmosphere. The level of seawater pCO₂ is mainly controlled by the structure of the acid-base system that is 17 18 influenced by the combined effect of biological activity (biomass production vs. 19 mineralization), CO₂ exchange with the atmosphere and temperature (Emerson and Hedges, 2008). 20

21 In this context the Baltic Sea can be considered as a very complex ecosystem. On one hand the 22 low buffer capacity (Beldowski et al., 2010) makes the seawater vulnerable to acidification in 23 most regions of the Baltic Sea, on the other hand the sea is exposed to various anthropogenic 24 influences which have the potential to change the acid-base system and thus also seawater pH and all pH-related processes. This makes the Baltic Sea different from the oceans for which the 25 26 CO₂ system as part of the acid-base system has been well characterized during the last decades (Bates et al., 2014; Riebesell et al., 2010; Dickson et al., 2007). Considerable research effort 27 28 was also undertaken in recent years to investigate the Baltic Sea CO₂ system and its 29 peculiarities. However, this was done from different perspectives and resulted in specific 30 problem-oriented and divers knowledge. Therefore, the goal of this review article is to collect 31 and integrate the existing knowledge on the structure and functioning of the acid-base system 32 in the Baltic Sea, to point out the research gaps and thus also to address challenges for the future research in this field. 33

2 2. The standard acid-base model for ocean water

Four measurable variables exist to describe the acid-base system of seawater. These are: pCO₂,
pH, total inorganic carbon concentration (C_T, known also as DIC) and total alkalinity (A_T). C_T
is the total concentration of all inorganic carbon forms present in seawater (Eq. 1), where
[CO₂*] represents the sum of molecular dissolved CO₂ and undissociated carbonic acid
(H₂CO₃):

8
$$C_T = [CO_2^*] + [HCO_3^-] + [CO_3^{2-}]$$
 (1)

9 Measurements of C_T are usually based on acidification of the sample and coulometric or 10 infrared detection of the extracted CO₂ (Dickson et al., 2007). A_T is defined as the excess of 11 proton acceptors (bases formed from weak acids with a dissociation constant of K \leq 10^{-4.5} at 25 12 °C) over proton donors (acids with K>10^{-4.5}) and expressed as the hydrogen ion equivalent in 13 one kilogram of sample (Dickson, 1981):

14
$$A_T = [HCO_3^-] + 2[CO_3^{2-}] + [B(OH)_4^-] + [OH^-] + [HPO_4^{2-}] + 2[PO_4^{3-}] + [SiO(OH)_3^-] + [NH_3] + [NH_3] + [NH_3^-] + [NH_3^-$$

15 $[HS^-] + [minor bases] - [H^+] - [HSO_4^-] - [HF] - [H_3PO_4] - [minor acids]$ (2)

A_T can be determined by open- or closed-cell acidimetric titration. For both C_T and A_T certified 16 reference materials are provided by the Andrew Dickson laboratory, UC San Diego, USA 17 18 (Dickson et al., 2007). pH measurements in seawater are usually calibrated against TRIS buffer. Its characteristics is well known at salinities of 20-35 (Mosley et al., 2004; DelValls and 19 20 Dickson, 1998). The efforts to improve the TRIS buffer characteristics for lower salinities are 21 ongoing within the BONUS PINBAL project (https://www.io-warnemuende.de/pinbalhome.html). The use of TRIS buffer helps to calibrate the systems for pH measurements but 22 23 does not fully guarantee the quality of the pH results as reference materials in seawater matrix still do not exist for pH measurements. In many cases pH is still measured potentiometrically 24 25 on the NBS scale although spectrophotometric measurements on the total scale (pHtot), which takes into account also HSO4⁻ ions, are currently state of the art in the field of chemical 26 oceanography. The concentration of CO_2^* in seawater is obtained from the partial pressure of 27 CO₂ in air in equilibrium with seawater. According to the standard operating procedure 28 29 (Dickson et al., 2007), measurements of pCO₂ require continuous flow of water, which makes 30 this parameter difficult to measure in discrete samples.

All four variables are interacting and control the pH as described by a set of equilibrium constants and mass balance equations. In general the interrelationships between these four

1 parameters facilitate the calculation of any two variables, when the two others are known, e.g. 2 through measurements, and when the dissociation constants of the involved acid-base reactions are known for the respective temperature and salinity. This fact is used in biogeochemical 3 models for simulations of the marine CO₂ system. All biogeochemical models are based on the 4 5 transport and transformations of A_T and C_T, because these variables are independent of temperature and pressure and behave conservatively with respect to mixing (mass 6 7 conservation). Moreover, straightforward techniques for sampling and analysis for A_T and C_T 8 exist and are supported by the availability of reference materials (Dickson et al., 2007). Hence, 9 these two variables are very well suited for CO₂ system studies. However, the use of A_T and C_T for the CO₂ system studies is not free from limitations. To obtain high accuracy for the 10 calculation of pH and pCO₂ from measured or modelled A_T and C_T, all dissociation constants 11 and total concentrations of all non-CO₂ components of the acid-base system must be known. 12 This requirement is approximately fulfilled with regard to ocean studies where the total 13 concentrations of non-CO₂ acid-base components are either negligible or can be approximated 14 as a function of salinity (Riebesell et al., 2010). However, this issue is more critical for coastal 15 and shelf regions, where the biogeochemical composition of seawater shows regional 16 17 peculiarities (e.g. Kuliński 2014; Hernandez-Ayon et al., 2007; Cai et al., 1998). This is the 18 case for the Baltic Sea, which shows various unique biogeochemical characteristics.

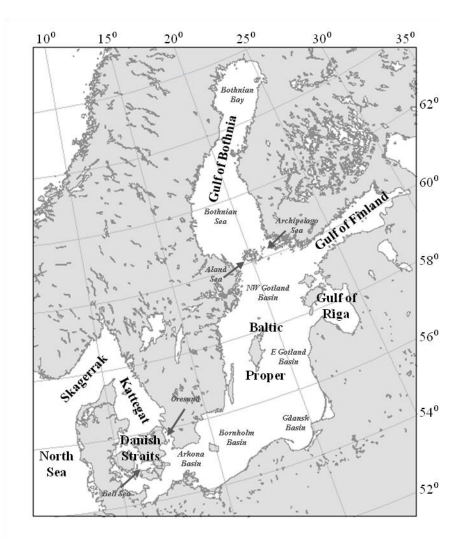
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20 3. Peculiarities of the Baltic Sea acid-base system

21 **3.1. Hydrographic setting**

The Baltic Sea is one of the largest brackish ecosystems in the world. This is caused by both 22 23 limited inflows of saline oceanic water through the shallow and narrow Danish Straits and high river runoff. The drainage basin of the Baltic Sea is almost four times larger than the area of 24 the sea itself, while the mean annual freshwater supply (428 km³) constitutes about 2 % of the 25 Baltic Sea water volume $(22 \cdot 10^3 \text{ km}^3)$. The specific features of the Baltic Sea hydrology cause 26 27 clear horizontal and vertical salinity gradients. Salinity in the surface layer fluctuate from 2 in the northern Bothnian Bay to >20 observed in the Kattegat. Dense and highly saline water, 28 29 which irregularly enters the Baltic Sea, sinks and moves along the sea bottom. This deep water is separated from the brackish surface water layer by a permanent halocline located at a depth 30 of 60-70 m, which is at the same time the maximum mixed layer depth in the Baltic Sea (Lass 31 32 and Matthäus, 2008; Winsor et al., 2001). The stratification limits ventilation of the deep water

masses. Simultaneously, the deeper water layers are supplied with large amounts of particulate
organic matter that either originated from terrestrial sources or derived from the biomass
production in the surface under the influence of high nutrient concentrations (eutrophication).
Some fraction of the sedimentary organic matter mineralizes releasing CO₂ and consuming
oxidants, which leads to hypoxia or even anoxia in the bottom waters (HELCOM, 2009).





7 Fig. 1. Map of the Baltic Sea showing its division into natural basins and sub-basins

8 (modified after Kuliński and Pempkowiak, 2011).

9

10 **3.2. Dissociation constants in brackish water**

11 If any variables of the marine CO_2 system are to be used for biogeochemical studies, it is 12 advisable to measure these directly and not to derive it from the measurements of other 13 variables of the CO_2 system (Dickson et al., 2007). However, sometimes direct measurements 14 may be prevented by technical reasons or routine analytical methods do not exist. The latter refers for example to the determination of CO_3^{2-} ion concentrations which are important for the assessment of the state of the CaCO₃ saturation and thus for the dissolution or formation of CaCO₃ shells. In this case, calculations on the basis of other known variables such as C_T, A_T, pCO₂ or pH are indispensable and require knowledge about the equilibrium constants of the CO₂ system. Here we examine the availability of the necessary constants for low salinity brackish water. Three fundamental equations characterize the equilibria of the marine CO₂ system. The first refers to the solubility of gaseous CO₂ in seawater:

8
$$[CO_2^*] = k_0 * fCO_2$$
 (3)

The solubility constant, k_0 , relates the concentration of CO_2^* in seawater to the CO_2 fugacity, 9 10 fCO_2 . In contrast to the CO_2 partial pressure, pCO_2 , the fugacity accounts for the non-ideal behaviour of CO₂ at atmospheric conditions. It differs only slightly from the pCO₂ and in many 11 12 cases can be replaced by the pCO₂. The solubility and thus k_0 decreases with increasing temperature and salinity and vice versa. Studies by Weiss (1974) that describe k₀ as a function 13 14 of temperature and salinity have received widespread acceptance and are almost exclusively used in chemical oceanography. The situation is somewhat more complicated for the 15 16 determination of the dissociation constants for carbonic acid because it is a diprotic acid. Using again CO₂* as variable, the two dissociation equilibria are given by: 17

18
$$K_1 = [H^+][HCO_3^-]/[CO_2^*]$$
 (4)

19
$$K_2 = [H^+][CO_3^{2-}]/[HCO_3^{-}]$$
 (5)

In these formulas the terms for the hydrogen ions are given in concentration units which include
HSO₄⁻ ion concentration (Dickson, 1984).

22 Systematic studies concerning the dissociation constants of carbonic acid in seawater were already performed during the twenties and thirties of the last century. One of the leading 23 24 scientists in this field was Kurt Buch from the former Finnish Institute for Marine Research, whose research was mainly related to the Baltic Sea, and whose laboratory studies on the CO₂ 25 system focused on brackish water with salinities down to zero. The results of his lab work and 26 of field studies performed during 1927 to 1936 are summarized in Buch (1945). The presented 27 28 dissociation constants must be considered as "hybrid" constants because the hydrogen ions are represented by the hydrogen ion activity whereas the CO₂ constituents refer to concentrations. 29 30 With the increasing awareness of the importance of the oceans for the uptake of anthropogenic

31 CO₂ and thus for climate change scenarios, large-scale measurement programmes concerning

1 the state of the marine CO_2 system on an oceanic scale were performed. These efforts were 2 accompanied by numerous laboratory studies aimed at the determination of improved dissociation constants. All these constants referred to the total hydrogen concentration scale 3 that includes the contribution by HSO4⁻ (Dickson, 1984). However, the validity of most of the 4 5 constants was confined to salinities that are encountered in ocean water. Exceptions are the constants suggested by Roy et al. (1993), which were determined for salinities down to values 6 7 of 5 and could be used accordingly for brackish water. However, with regard to research in the Baltic Sea where large areas, e.g. in the Gulf of Bothnia, have surface water salinities less than 8 9 5, the situation was unsatisfactory. It took until 2006 when Millero et al. (2006) published dissociation constants that covered the salinity range from 0 to 50 and that were consistent with 10 the constants for fresh water. An update of these constants was performed in 2010 (Millero, 11 2010) and since then this set of dissociation constants is state of the art for CO₂ research in 12 brackish waters. The salinity and the temperature dependency of K₁ and K₂ are presented in 13 14 Fig. 2a and 2b, respectively.

A direct comparison between the dissociation constants that were used in the past for brackish water is difficult because the constants by Buch (1945) are based on a pH scale that is based on the hydrogen ion activity. Since it is difficult to convert the activities to total hydrogen ion concentrations, we use the ratio K_1/K_2 for a comparison. This ratio is obtained by dividing the equilibrium equations for the first dissociation step (Eq. 4) through that of the second dissociation step (Eq. 5):

21
$$\frac{K_1}{K_2} = \frac{[HCO_3^-]^2}{[CO_2^*]*[CO_3^{2-}]}$$
 (6)

Eq. 6 shows that K_1/K_2 is not connected to the hydrogen ion concentration and is thus not affected by the choice of the pH scale. Hence it is suited as an indicator for the quality of the early dissociation constants determined by Buch (1945). We used the logarithmic notation, $log(K_1/K_2)$ which is equivalent to $pK_1 - pK_2$, for a comparison of the dissociation constants suggested by Buch (1945), Millero (2010) and Roy et al. (1993). The differences $\Delta log(K_1/K_2)$ referring to Millero (2010) and Buch (1945), and Millero (2010) and Roy et al. (1993) indicated as M-B and M-R, respectively, are shown as a function of salinity in Fig. 2c.

At salinities >7, M-R is less than 0.03, but the differences increase rapidly at lower salinities and amount to 0.06 and 0.10 already at S = 5.4 and S = 3.6, respectively. Such differences in $\log(K_1/K_2)$ are equivalent to a temperature change of 1 °C. The deviations of the Buch (1945) data from those of Millero (2010) (M-B in Fig. 2c) ranged below 0.10 at salinities below 20.

- 1 This is a surprisingly small difference in view of the limited technical capabilities that were
- 2 available to Buch and his co-workers in the twenties and thirties of the last century.
- 3

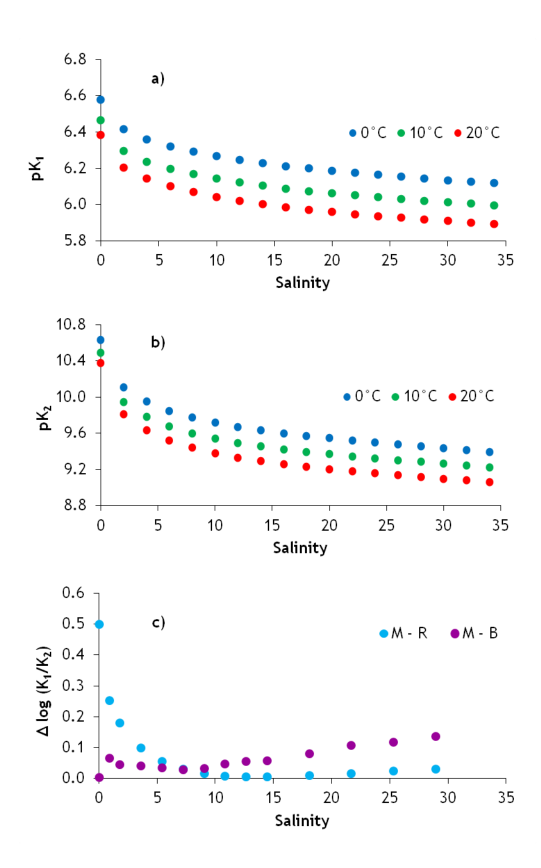


Fig. 2. Dissociation constants (a) pK₁ and (b) pK₂ as a function of salinity calculated according
to Millero (2010) for three different temperatures: 0, 10 and 20 °C; (c) differences in log(K₁/K₂)
between the dissociation constants obtained from Millero (2010) and Roy et al. (1993) (M-R,
blue dots) and between Millero (2010) and Buch (1945) (M-B, purple dots) at different
salinities.

6

7 When alkalinity is also used for the calculation of any variables of the CO₂ system, the contributions of non-CO₂ acid-base components to A_T (Eq. 2), which are generally low, must 8 9 be taken into account. This requires knowledge about the concentration of the individual acid-10 base pairs and about the corresponding dissociation constants. In the deep basins of the central Baltic Sea, ammonia and sulphide anions are released during organic matter mineralization at 11 12 anoxic conditions (see Section 3.4.2). Both these constituents are strong proton acceptors forming mainly HS^- and NH_4^+ . Since their concentrations may be on the order of several tens 13 and up to a few hundred μ mol L⁻¹ (Nausch et al., 2008), respectively, they have a strong effect 14 on alkalinity and pH. Therefore, any calculations concerning the CO₂ system at anoxic 15 16 conditions must account for the dissociation equilibria of H₂S-HS⁻ (Millero et al., 1988) and NH4⁺-NH₃ (Clegg and Whitfield, 1995; Johansson and Wedborg, 1980). The concentrations of 17 S^{2-} at the pH of anoxic water are extremely small and can be neglected. 18

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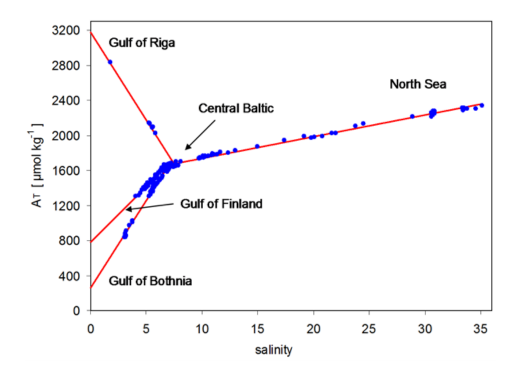
20 **3.3.** The central role of alkalinity

21 3.3.1. Regional A_T vs. S distribution, consequences for pH distribution pattern

There are several different A_T vs. S regimes in the Baltic Sea (Fig. 3, Beldowski et al., 2010). 22 They reflect different A_T concentrations in the respective rivers, which can be deduced from 23 extrapolation of regional A_T vs. S relationships to zero salinity. Low alkalinities are observed 24 in rivers entering the Gulf of Bothnia whereas rivers from south-eastern part of the Baltic 25 catchment, as shown by the A_T vs. S plot for the Gulf of Riga (Fig. 3), are rich in alkalinity. 26 These differences are a consequence of the geological conditions and weathering processes in 27 28 the respective catchment areas. As a result lower alkalinities and lower mean pH are observed in the Gulf of Bothnia and Finland, whereas higher mean alkalinities and thus somewhat higher 29 pH are found in the Gulf of Riga and also in the Gdansk Bay (Kulinski et al., 2014; Beldowski 30 et al., 2010; Hjalmarsson et al., 2008). 31

1 The central Baltic Sea acts as a mixing chamber for the different water masses, including water

originating from the North Sea. This results in alkalinity in the surface water of the Baltic
Proper (salinity around 7) of about 1600-1700 µmol kg⁻¹.



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Fig. 3. Different A_T vs. S regimes observed in the Baltic Sea (modified after Beldowski et al.,
2010)

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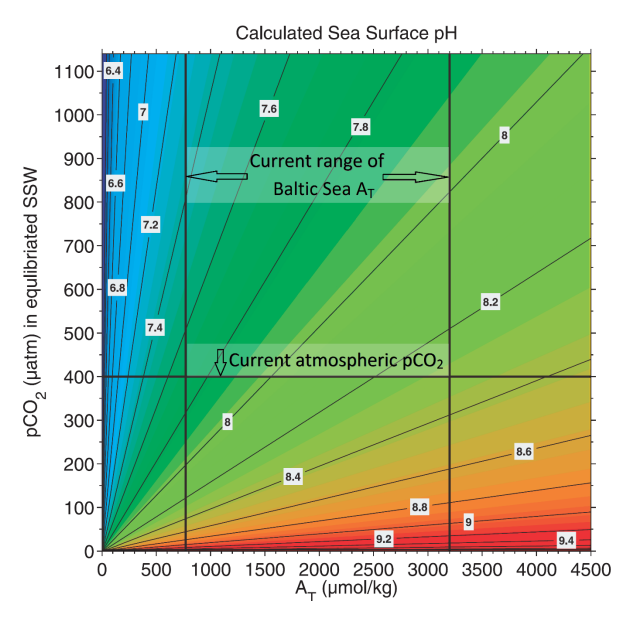
An important, but still not well investigated, aspect of the A_T supply to the Baltic Sea is the A_T 8 seasonality in river water. As A_T is one of the variables used in biogeochemical models to study 9 the CO₂ system, well described spatial and temporal variability of A_T loads could improve the 10 characterization of the A_T distribution in the Baltic Sea and thus increase the accuracy of the 11 12 modelled pH and pCO₂. This could be achieved by evaluation of the A_T monitoring data which are regularly collected by some Baltic Sea states within the framework of the HELCOM 13 monitoring program. In some countries such as Sweden the monitoring data are publicly 14 accessible, in others still some limitations exist. 15

16 A_T can also enter the Baltic Sea by submarine groundwater discharge (SGD). The chemical 17 composition of groundwater, as in river water, depends to some extent on the geological 18 structure of the catchment through which the water flows. Thus, SGD entering the Baltic Sea 19 along the continental part of the coast can be rich in A_T . Szymczycha et al. (2014) noticed 20 significant C_T concentrations (5400 µmol kg⁻¹ on average) in SGD seeping to the southern Baltic along the Polish coast. Although there might be some contribution from CO₂ to this high
 C_T, it is very likely that SGD can play, at least locally, an important role for the A_T budget.
 However, direct measurements of A_T concentrations and loads in SGD have not been reported
 in the Baltic Sea so far.

At equilibrium with the atmospheric CO₂, A_T controls C_T and thus pH. Hence, pH may be 5 depicted as a function of A_T and pCO₂. A sensitivity study performed by Omstedt et al., (2010) 6 7 indicates that the pH of the Baltic Sea surface water that is at equilibrium with the atmosphere (pCO₂ of ca. 400 µatm at 0 °C) can vary between 7.7 and 8.3 depending on the A_T (Fig. 4). 8 9 This range is significantly higher than that observed in the open ocean, where A_T oscillates only in a narrow range: 2170-2460 µmol kg⁻¹ (Riebesell et al., 2010). The reason for that is the 10 high spatial variability of A_T in the surface Baltic Sea waters, from low A_T (below 1000 µmol 11 kg⁻¹) observed in the Bothnian Bay to the A_T-rich (more than 3000 μ mol kg⁻¹) estuaries of the 12 large continental rivers. The diagram presented in Fig. 4 also shows that higher A_T reduces 13 shifts in seawater pH caused by changes in pCO₂. 14

15

16



1

Fig. 4. Distribution of pH_{tot} as a function of A_T and pCO₂ at salinity of 7.5 and temperature of
0 °C (modified after Omstedt et al., 2010). The A_T range represents findings by Beldowski et
al. (2010). SSW is an abbreviation for surface seawater.

6 3.3.2. Long-term A_T changes, consequences for ocean acidification

Long-term high resolution pH measurements at station BATS (Bermuda Atlantic Time Series)
in the sub-tropical North Atlantic have shown that ocean acidification occurs at a rate that is
thermodynamically consistent with the increase of the atmospheric CO₂ (Bates, 2014). This
gives reason to expect that the progression of ocean acidification can be assessed on the basis
of CO₂ emission scenarios and the modelling of the effect on the atmospheric CO₂. It also
implies that changes in alkalinity will not occur in the future. On time scales of decades this

1 may be a reasonable assumption for the oceans which react very slowly to internal or external 2 changes. However, the situation is different for the Baltic Sea that is under the immediate influence of natural and anthropogenic processes on land. The first hints of increasing alkalinity 3 in the surface water of the central Baltic Sea were reported by Schneider et al. (2015) who 4 5 argued that this may have considerably mitigated the acidification due to increasing atmospheric CO₂. Müller et al. (2016) took up this idea and performed a thorough and 6 7 comprehensive statistical analysis of all alkalinity data that were available since the start of CO₂ research in the Baltic Sea at the beginning of the last century. The authors focused on 8 high-quality data measured after 1995 and detected a distinct AT trend that showed a clear 9 regional gradient. The highest trend was found in the Gulf of Bothnia (7.0 µmol kg⁻¹ yr⁻¹), 10 followed by 3.4 µmol kg⁻¹ yr⁻¹ in the central Baltic, whereas no trend could be detected in the 11 Kattegat. Müller et al. (2016) estimated that the changes in A_T have reduced the acidification 12 effect in the central Baltic Sea by about 50 % and that the Gulf of Bothnia was not subjected 13 at all to acidification during the last 20 years. Several reasons for the increasing AT are 14 discussed by Müller et al. (2016), but a major player could not be identified and, hence, it 15 remains unclear whether the trend will continue in the future. 16

17

18 **3.3.3. Effect of organic alkalinity**

The Baltic Sea water contains 3-5 times more organic matter than open ocean waters. In the 19 surface water of the open Baltic Sea, concentrations of dissolved organic carbon (DOC) range 20 from about 260 to about 480 μ mol C L⁻¹, while those in the surface water (top 100 m) of the 21 Atlantic Ocean are much lower and range between 50 and 80 μ mol C L⁻¹ (Hoikkala et al., 2015; 22 Carlson et al., 2010). This is a result of both high inputs of terrestrial organic matter and 23 24 eutrophication driven by nutrient supply from land (Hoikkala et al., 2015; Kuliński and Pempkowiak, 2011; Kuliński et al., 2011). Organic substances contain functional groups, some 25 26 of which (carboxylic, phenolic, amines) have acidic character and can dissociate in seawater releasing protons (H⁺). This contributes to pH decrease. However, as most of these groups are 27 28 believed to act as weak acids (pKa>4.5), their dissociation releases also an equivalent of organic 29 anions being strong bases. This, according to Eq. 2, causes no change in A_T concentration since 30 an equivalent amount of protons have been released. It affects the internal structure of A_T by changing the contributions of different A_T components according to their individual 31 32 dissociation constants (Cai et al., 1998; Hunt et al., 2011; Kuliński et al., 2014). The lower the pK_a of an organic acid added to the system, the greater are the shifts in the internal A_T 33

distribution. As a consequence bases forming the acid-base system are partially protonated and their concentration decreases while the concentration of the corresponding undissociated acid is increasing. In case of the carbonate alkalinity this means a reduction of CO_3^{2-} and an increase of H₂CO₃ and thus of the pCO₂.

Kuliński et al. (2014) estimated the organic alkalinity (Aorg) in the Baltic Sea as an excess 5 alkalinity calculated from the difference between measured A_T and that calculated from C_T and 6 pH or pCO₂. They found A_{org} of 25-35 µmol kg⁻¹ in the surface Baltic Sea water (from the 7 Arkona Basin to the Bothnian Bay) without any clear spatial distribution pattern. This Aorg 8 contribution amounted from 1.5 % to 3.5 % of the measured A_T values. Even higher values of 9 more than 50 µmol kg⁻¹ were found by Hammer et al. (2017) in the Baltic Sea surface mixed 10 layer during the spring phytoplankton bloom. They reported also that Aorg decreases with depth 11 and approaches zero below the permanent halocline due to pH decrease. 12

In sensitivity studies Kuliński et al. (2014) showed for the Baltic Sea that calculations 13 14 concerning the CO₂ system that are using measured A_T may lead to significant errors if A_{org} is ignored. Highest deviations between calculated and observed values were found when AT was 15 used together with C_T for computations of pCO₂ and pH. The pCO₂ values obtained in that way 16 were 27-56 % lower than the measured ones, while pH was overestimated by more than 0.4 17 18 units. These results are especially important as this combination (A_T and C_T) is used in 19 biogeochemical models (Edman and Omstedt, 2013; Kuznetsov and Neumann, 2013; Omstedt et al., 2012 and 2009) because they are conservative variables. This means that they are 20 independent of pressure and temperature and follow the law of conservation of mass during 21 mixing, which are prerequisites for variables transported in models. The first attempts to 22 include an A_{org} parametrisation in biogeochemical models have been made by Gustafsson et 23 al., (2015) and Omstedt et al. (2015). However, the sensitivity of the models with regard to the 24 inclusion/negligence of Aorg has not yet been reported. 25

26 The source of the calculation errors related to ignoring Aorg is due to the fact that measurements include all inorganic and organic contributors, while subsequent calculations using the standard 27 chemical A_T model and respective software, interpret the measured A_T only as inorganic 28 29 alkalinity. Including organic substances into the A_T model is, however, challenging as there are 30 number of organic substances having acidic functional groups and, according to the AT definition, for each of them the information on concentration and dissociation constants must 31 32 be available. Accounting for Aorg in the chemical AT model is less important for oceanic research because the low concentrations of dissolved organic matter (DOM) in the oceans cause 33

1 only minor effects. As a first approximation of the acid-base properties of DOM for the Baltic 2 Sea water, Kuliński et al. (2014) proposed to use one single bulk dissociation constant, pK_{DOM} and relate Aorg to the concentration of dissolved organic carbon (DOC), a commonly measured 3 parameter. According to their studies, slightly corrected by Ulfsbo et al. (2015), pK_{DOM} in the 4 5 Baltic Sea amounts to 7.34. They also estimated experimentally that 12 % of DOC acts as a carrier for acidic functional groups in the Baltic Sea DOM. The model studies by Ulfsbo et al. 6 7 (2015) showed that this experimentally derived share fits to the share estimated from the hypothetical structure of the fulvic acids. They also suggested that the method proposed by 8 9 Kuliński et al. (2014) is the best available approach for representing organic alkalinity in biogeochemical models at the current state of knowledge. They also showed that organic matter 10 (as represented by the fulvic acids) contains also some functional groups having pK_a already 11 below 4.5. According to the definition (Dickson, 1981) they should be considered as strong 12 acids. In experimental studies Hammer et al. (2017) also found that humic and/or fulvic 13 substances are more acidic than the bulk DOM naturally occurring in the Baltic Sea. This may 14 also be a source of uncertainty in the A_T measurements as the determination of the titration 15 16 end-point requires to titrate the samples through this pH region.

17

18 **3.3.4.** Role of borate alkalinity

The carbonate system plays a central role in the marine acid-base system. Thus, studies on 19 20 seawater pH and buffering capacity focus usually on the carbon species. Less attention is paid 21 to boron, though borates are, after bicarbonates and carbonates, the third most abundant 22 constituent of seawater AT (Eq. 2). Boron exists in seawater in form of weak boric acid, B(OH)₃, and the anion, B(OH)₄. The high pK_a (8.60 at salinity 35 and temperature 25 $^{\circ}$ C) 23 means that at seawater pH of about 8 the undissociated boric acid predominates. The boron 24 25 concentration, B_T, in seawater is approximated as a function of salinity or, for the historical data, as a function of chlorinity (S = 1.80655 * Cl). First measurements of boron concentrations 26 in the Baltic Sea were reported by Buch (1945), who found that $B_T [mg kg^{-1}] = 0.133 * S$. This 27 finding was confirmed recently by Lee et al. (2010) for oceanic waters. Another relationship 28 linking B_T and S and often used in acid-base system studies is that by Uppström (1974), who 29 reported that $B_T [mg kg^{-1}] = 0.128 * S$. All these formulas suggest a fixed B_T/S ratio which is 30 only the case if the river water that enters the Baltic Sea contains no boron. However, the 31 studies by Kremling (1970, 1972) indicated that this is not the case and that a B_T concentration 32 anomaly exists in the Baltic Sea, where the experimentally determined B_T vs. S relationship 33

yielded a river water B_T (anomaly term) of $0.15 - 0.20 \text{ mg kg}^{-1}$ at S = 0. This anomaly is not included in the chemical A_T model commonly used for numerical simulations of the CO₂ system and, similar to the effects of ignoring A_{org} , may lead to wrong conclusions. This can be especially critical at low salinities, where the effect of the anomaly is the largest.

5

3.4. Modulation of the acid-base system by organic matter production/mineralization (concurrent A_T and C_T changes)

8 **3.4.1.** Biomass production

9 In the case that pH changes refer to equilibrium with the atmosphere, it may be meaningful to 10 consider the pH at a given alkalinity as a function of pCO_2 , because the atmospheric CO_2 is then the only driver for any pH changes. However, when assessing pH changes as the 11 12 consequence of internal biogeochemical transformations, it is more appropriate to consider A_{T} and C_T as the controls for pH and pCO₂. Biomass production primarily alters C_T because 13 phytoplankton (but also other plants) consume CO₂ in the course of photosynthesis (Fig. 5). As 14 15 a consequence it increases the pH and lowers the pCO₂ in the upper water layers and causes a pCO₂ disequilibrium between seawater and the atmosphere. This leads to CO₂ uptake by the 16 seawater and thus counteracts the effect of biomass production and dampens the pH increase. 17 However, the CO₂ exchange through the air-sea interface is much slower than the effects of 18 biological processes. Therefore, the Baltic Sea water is in almost permanent pCO₂ 19 20 disequilibrium with the atmosphere throughout the year. In spring and summer seawater is 21 undersaturated with respect to atmospheric CO₂ with two characteristic pCO₂ minima and two pH maxima (Fig. 5) which reflect the spring bloom and the mid-summer nitrogen fixation 22 23 period. In autumn and winter the Baltic Sea surface water pCO₂ increases over the atmospheric values as a consequence of less active production in the upper water column and transport of 24 25 deeper CO₂-enriched water to the surface by mixing (Fig. 5; Schneider, 2011).

Another way in which biomass production influences seawater pH and pCO₂ is related to nitrate consumption. As phytoplankton assimilate nitrate for growth an equivalent of H⁺ is also transported to the cells to keep the charge balance neutral. According to Eq. 2 this increases the seawater A_T and thus raises the pH during biomass production (Brewer and Goldman, 1976). This assimilation also decreases the pCO₂ and therefore reinforces the drop in pCO₂ by biomass production. Some small A_T changes can be also induced by phosphate consumption. However, due to low phosphate concentration the effect is negligible (Wolf-Gladrow et al.,
 2007).

Biological production also may consume CO_3^{2-} ions in the calcification processes. This causes A_T reduction and in consequence pCO₂ increase and also pH decrease. However, this mechanism has been recognized as less important in the Baltic Sea. According to studies by Tyrrell et al. (2008) the Baltic Sea surface water is undersaturated in winter with respect to both aragonite and calcite which are biogenic modifications of CaCO₃. This possibly prevents the growth of calcifying plankton and is the reason for the absence of coccolithophores in the Baltic Sea.

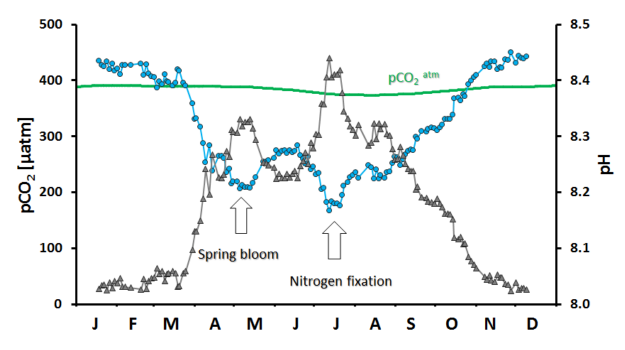


Fig. 5. Typical seasonality of pCO₂ in surface water (blue line) and the atmosphere (green line)
in the Eastern Gotland Sea modified after Schneider (2011) and presented together with pH
calculated from the shown pCO₂ data and the mean A_T value.

14

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15 **3.4.2. Remineralization**

In contrast to biomass production, CO_2 is released during the remineralization processes and causes an increase in C_T . Consequently, this leads to a pH decrease and pCO₂ increase. Some fraction of organic matter produced in situ undergoes remineralization in the upper water layers. Particulate organic matter (POM) may be mineralized directly or via prior release of dissolved organic matter (DOM). In the Baltic Sea terrigenous organic matter entering the sea from rivers constitutes an important part of the organic matter. Kuliński and Pempkowiak 1 (2011) reported that the Baltic Sea receives annually 340 Gmol of organic carbon from land. 2 The model studies by Gustafsson et al. (2014a) suggested that as much as 39.5% of terrigenous organic carbon that enters the Baltic Sea undergoes mineralization without distinguishing 3 between mineralization pathways. This makes that it is difficult to estimate all the 4 5 consequences of that terrestrial organic carbon loss for the Baltic Sea acid-base system. The incubation experiments performed by Kuliński et al. (2016) indicated that ca. 20 % of terrestrial 6 7 dissolved organic carbon and 34 % of DOC present in the Baltic Sea is bioavailable. It is 8 important to mention here that the latter result refers likely to the mixture of terrestrial and 9 marine DOC.

10 Whereas the A_T in surface water is mainly controlled by mixing of different water masses, the deep water A_T distribution depends additionally on the organic matter transformations by 11 12 various redox processes (Brenner et al., 2016; Krumins et al., 2013; Thomas et al., 2009; Schulz and Zabel, 2006). A certain fraction of the organic matter produced in the euphotic zone is 13 14 exported to deeper water layers and to surface sediments, where it undergoes mineralization, 15 produces CO₂, and changes the alkalinity. The change in A_T depends on the oxidant that is required for the mineralization and may cover a wide range. Therefore, it is not possible to 16 17 predict the change of pH or pCO₂ during the mineralization process without knowledge of the oxidant. In the presence of oxygen, mineralization takes place according to Eq. 7 which is 18 19 reversing the bulk photosynthesis reaction. However, in sediments and in deep water layers of some basins of the central Baltic, where longer periods (years) of stagnation occur, oxygen 20 21 may be entirely depleted. Organic matter can then be mineralized in a certain 22 thermodynamically controlled sequence by other oxidants (Schulz and Zabel, 2006). First, 23 manganese dioxide takes over the role of oxygen (Eq. 8), followed by denitrification where 24 nitrate acts as oxidant (Eq. 9). After nitrate is consumed, the mineralization process may be continued by iron (III) oxide (Eq. 10) before sulphate oxidizes the organic matter and generates 25 hydrogen sulphide (Eq. 11). In the Baltic Sea these processes may take place in the water 26 column where a pelagic redoxcline (an interface between oxic and anoxic conditions) can 27 develop during longer periods of stagnation. Only the final mineralization, that is an internal 28 29 oxidation and generates methane (methanogenesis, Eq. 12) after the sulphate concentration has 30 approached zero, is confined to deeper sediment layers only. Except oxic mineralization, all other processes are connected with a simultaneous A_T increase caused either by release of 31 strong bases (S^{2-}) or by consumption of H⁺ (see Eq. 2) (Ulfsbo et al., 2011; Schneider et al., 32 33 2015; Lukawska-Matuszewska, 2016). The change in A_T (ΔA_T) for different mineralization

1 Generally, benthic processes linked to early diagenesis of organic matter can be an important 2 source for A_T to the water column (Brenner et al., 2016). This corresponds to the findings by Gustafsson et al. (2014b), who found in a model study that external sinks and sources of A_T in 3 the Baltic Sea are imbalanced and cannot reproduce the observed A_T inventory, and that an 4 5 internal A_T source must exist in the Baltic Sea. However, it must be taken into account that 6 alkalinity released from sediments as sulphide or ammonia only influence the acid-base system 7 locally. Upon contact with oxic water most of the A_T generating processes – except 8 denitrification - are reversed and do not constitute a permanent A_T source. Some minor increase 9 of A_T can be also expected due to the burial of iron (II) minerals like pyrite or vivianite (Reed 10 et al., 2016).

11

12 **4. Challenges for future research.**

The general knowledge of the marine acid-base system was gained mainly from the oceanic 13 14 research. It has been integrated and documented in the form of a guide (Dickson et al., 2007) and included in different software packages (Orr et al., 2015), that allow for computations of 15 16 two out of the four acid-base system parameters (pCO_2 , pH, A_T , C_T), when the other two are known (e.g. by measurements). Although this works nicely for the open ocean waters, problems 17 appear when that knowledge is used for studying the acid-base system related processes in 18 19 coastal and shelf seas. This uncovers research gaps and bottlenecks but also define challenges for marine biogeochemists working in these regions. 20

21 In this context the Baltic Sea can be considered as a model experimental field, where strong 22 horizontal and vertical salinity gradients, permanent stratification of the water column, 23 eutrophication, high organic matter concentrations and high anthropogenic pressure make the 24 acid-base system even more tangled. The challenges are related to analytical methods, interpretation of the data and parametrization of the acid-base system related processes for the 25 26 use in numerical models. Great effort has recently been made to adapt spectrophotometric pH measurements based on m-cresol purple to the Baltic Sea conditions. This was done by 27 28 Hammer et al. (2014) and was recently improved within a framework of the BONUS PINBAL 29 project. Further progress can be expected if pH reference materials for brackish water are made 30 available in the future. For the pCO₂ measurements the state-of-the-art method that is based on measuring the CO_2 content in air equilibrated with the seawater is commonly used (e.g. 31 32 Schneider et al., 2014). Due to a relatively long response time, this method may, however, not be fast enough to resolve steep horizontal pCO₂ gradients in the vicinities of river mouths. The 33

1 development of accurate and precise sensors for pCO₂, like optodes, could improve our 2 understanding of the CO₂ system in the transition area from the coast to the open sea. Additionally, it would facilitate pCO₂ measurements in discrete samples, which are highly 3 desired for investigations of the organic matter mineralization processes in the water column, 4 5 but difficult to obtain with common underway measurement techniques. The determination of A_T by titration with an acid (HCl) allows a high accuracy (+/-2 µmol kg⁻¹) for ocean water. 6 7 This is difficult to achieve for Baltic Sea water because the detection of the endpoint of the AT titration is more difficult due to the influence of organic acid-base constituents. The studies by 8 9 Ulfsbo et al. (2015) showed that fulvic and humic acids, substances widely spread in the Baltic Sea, have relatively high concentrations of acid-base constituents, which have pKa values close 10 to the pK_a of carbonic acid and may thus impede the correct detection of the titration endpoint. 11 C_T measurements are less critical since reference materials exist and the analytical procedure 12 is relatively straightforward. However, development of high accuracy underway methods for 13 C_T (but also A_T) would improve the horizontal coverage of the data. 14

15 Interpretations of the acid-base system data require appropriate parameterization of the related processes and chemical reactions. With regard to the Baltic Sea some peculiarities of the 16 17 composition of the acid-base system must be taken into account which are neglectable in ocean waters. An example is the influence of DOM on the acid-base system. In the oceans it is ignored 18 19 due to low DOM concentrations but in the Baltic Sea Aorg becomes a noticeable component of A_T (Kuliński et al., 2014). The challenge remains to include the contribution by DOM to the 20 21 chemical A_T model. The approach suggested by Kuliński et al. (2014) to use the so called bulk 22 dissociation constant, K_{DOM}, is only a first approximation, as K_{DOM} has no real thermodynamic 23 meaning. It also does not allow for distinguishing the influence of individual substances. Respective improvements would require substantial progress in analytical techniques, as the 24 characterization of the structure and composition of DOM is still far from being satisfactory 25 (Nebbioso and Piccolo, 2013). Problems also arise due to the ion anomalies observed in the 26 Baltic Sea (Kremling 1970, 1972) that are not taken into account in the A_T model. An example 27 is boron, whose concentration is usually approximated by the assumption of a constant 28 boron/salinity ratio. However, the function linking both these parameters does not account for 29 30 river water as a source of boron. Furthermore, common software for CO₂ system calculation also omits seawater constituents that are generated at anoxic conditions. These are mainly the 31 acid-base systems: H₂S-HS⁻-S²⁻ and NH₄⁺-NH₃ which require consideration in computational 32 models. All these shortcomings may cause uncertainties in any calculations of the CO₂ system 33

1 including those in biogeochemical models when A_T is used as a controlling variable. It is also

recommended to perform sensitivity studies in order to estimate the quantitative importance of
inadequate chemical characterization of A_T.

4 Furthermore, we promote the implementation of the acid-base system (or marine CO₂ system) into biogeochemical models. This includes the simulation of surface water CO₂ partial 5 pressure, pCO_2 . The pCO_2 is an ideal validation variable because on the one hand it can easily 6 be measured with high spatiotemporal resolution (e.g. on Voluntary Observing Ships, VOS), 7 8 and on the other hand it reflects the biogeochemical transformations such as biomass production. Hence, agreement between simulated and measured pCO₂ data give an indication 9 10 of the quality of the biogeochemical process parameterization. Or, vice versa, unsatisfactory 11 agreement gives reason to rethink the description of the respective processes in the model.

12 The latter aspect was the focus of several model studies which initially failed to reproduce the observed pCO₂ draw down during spring and summer (Fig. 5). This indicated that the 13 14 traditional parameterization of the biomass production that is based on the Redfield (1963) C/N/P ratios and on nutrient concentrations at the start of the productive period, did not reflect 15 16 the reality. Omstedt et al. (2009) solved the problem by including a fractional nutrient release into their model. This meant that nutrients that have been used for production are partly released 17 18 from the POM and reused for new production. In another study Kuznetsov et al. (2011) 19 introduced an additional cyanobacteria group and flexible C/N/P ratios into their model in order to achieve approximate agreement between the simulated and measured pCO_2 . A further study 20 by Gustafsson et al. (2014a) investigated the changes in the simulated CO₂ air-sea flux after 21 including organic alkalinity and changes in external inputs of carbon, alkalinity and nutrients. 22 Modelling the CO₂ system was also used to simulate the vertical distribution of total CO₂ and 23 24 alkalinity in stagnant waters of the deep basins (Edman and Omstedt, 2013) and to assess the 25 importance of internal alkalinity generation (Gustafsson et al., 2014b). Finally, model calculations were used to estimate future changes in the Baltic Sea CO₂ system and in particular 26 in the development of the ocean acidification effect (Omstedt et al., 2012; Kuznetzov and 27 28 Neumann, 2013). Both simulations indicated that the increasing atmospheric CO₂ will mainly 29 control long-term changes in pH, of course, not taking into account the recently reported current increase in alkalinity (see Chapter 3.3.2; Müller et al., 2016). 30

31

32 **5. Summary and conclusions**

1 With our review we have proposed that the Baltic Sea acid-base system cannot be explained 2 by dissolved CO₂ and a minor contribution of boric acid. Whereas acid-base components other than CO_2 and borate may be ignored in ocean waters, this is not the case in the Baltic Sea and, 3 presumably, other marginal and semi-enclosed seas, which in a similar way are strongly 4 5 impacted by high biomass production and natural and anthropogenic processes in connected catchment areas as well. The complexity of the Baltic Sea acid-base system is displayed in the 6 7 formulation of the alkalinity (Eq. 2), which is the central variable for the characterization of 8 the acid-base properties of seawater. The peculiarities of the Baltic Sea alkalinity system are 9 either caused by specific internal processes related to intense production/mineralization of organic matter, or to the immediate effect of riverine input of acidic substances. This riverine 10 input refers mostly to dissolved organic matter which may strongly affect the composition of 11 the alkalinity in the Baltic Sea surface water (Kuliński et al., 2014), but also to boric acid 12 (borate) which in ocean water is linked to salinity by a constant ratio, but may show distinct 13 deviations from this ratio in some of the Baltic Sea estuaries because of boron input by river 14 15 water.

Internal biogeochemical processes such as organic matter production or mineralization also 16 17 have the potential to affect the acid-base system in the Baltic Sea. Alkalinity changes by organic matter production caused by the removal of H⁺ during the uptake of nitrate are of minor 18 19 importance. This may be different if the production is associated with calcification, which reduces the alkalinity, but the abundance of calcifying plankton in the Baltic Sea is restricted 20 21 to the high-salinity Kattegat region. Although the effect of organic matter production on 22 alkalinity is minor, it is the major control for the seasonal modulation of the acid-base 23 properties such as pH and pCO₂.

24 In contrast, organic matter mineralization taking place at anoxic conditions in the Baltic Sea 25 deep basins causes changes in the acid base system which are specific for marine systems with a pelagic redoxcline. Anoxic mineralization of organic matter generates large amounts of 26 27 alkalinity by the formation of sulphide ions and the release of ammonia. This reduces the 28 increase of the pCO₂ in the deep water and stabilizes the pH at a value close to 7. Although 29 these processes are reversed upon re-oxidation during a deep water renewal event and do not affect the acid-base system of the Baltic Sea as a whole, they do influence the deep water redox 30 31 chemistry which is partly controlled by the pH. On the other hand the existence of a pelagic 32 redoxcline has another more far-reaching importance. It is the medium for intense

denitrification, which increases the alkalinity not only locally, but may affect the alkalinity
 budget of the entire Baltic Sea proper (Gustafsson et al., 2014).

Our compilation of the major features of the Baltic Sea acid-base system indicates that an exact 3 4 quantitative treatment of its properties is difficult to achieve. This refers to both the biogeochemical modelling and the interpretation of measured alkalinity in terms of the 5 6 calculation of individual alkalinity contributions. An example is the calculation of the carbonate alkalinity that is necessary for the full characterization of the CO₂ system based on 7 8 measurements of alkalinity together with another variable. This requires knowledge of the dissociation constants and the determination of the total concentrations of the individual 9 10 compounds with acid-base properties. Regarding inorganic alkalinity contributions, it may be realistic to obtain this information. But in view of our limited knowledge concerning the 11 12 composition of the dissolved organic matter, it is currently imposible to specify and characterize the contributions of individual organic acid-base compounds to the alkalinity. 13 14 Hence we conclude that this is one of the greatest challenges for more adequate comprehensive 15 physico-chemical characterization of the acid-base system in the Baltic Sea and likely in other coastal seas. Another important bottleneck, especially for the biogeochemical modelling, is 16 17 also an insufficient knowledge on short and long term development of alkalinity loads from land caused by processes occurring in the catchment. 18

19

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