

PDF issue: 2024-09-08

Modeling dissolved inorganic carbon considering submerged aquatic vegetation

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(Citation) Ecological Modelling,431:109188

(Issue Date) 2020-09-01

(Resource Type) journal article

(Version) Accepted Manuscript

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https://hdl.handle.net/20.500.14094/90007400



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4	Modelling dissolved inorganic carbon considering submerged aquatic vegetation
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29	Hig	hlights
30	•	Aquatic vegetation can contribute to carbon capture in a lagoon system.
31	•	A Seasonal NEP (SNEP) model is presented to estimate the change in DIC.
32	•	The model can be applied to estimate lagoon productivity with limited
33		information.
34	•	Model results highlight the importance of estimating residence time.
35	•	The SNEP model proves benefit of seagrass restoration on effective carbon
36		capture.
37		

38 **ABSTRACT:**

39 Net absorption of CO₂ by vegetated coastal ecosystems has been revealed as a key mechanism 40 to capture and store carbon via the renewal of epigeal stem and rhizome biomass. Submerged 41 aquatic ecosystems, such as seagrass meadows, have been termed "blue carbon" ecosystems 42 because they absorb CO2 for their underwater growth. Irradiance and water temperature are 43 significant factors controlling net ecological production (NEP) by seagrass. As seagrass tends to 44 grow in calm coastal areas subject to water-column stratification, such as lagoons, a new method 45 for evaluating NEP accurately to access blue carbon capture in these enclosed waters is required. 46 This study aimed to develop a model to investigate thermal effects, considering irradiance, on 47 changes in dissolved inorganic carbon dynamics in a lagoon system, and assessment of the model 48 to understand controls on carbon dynamics in Komuke Lagoon, Japan. NEP was successfully 49 modelled by verifying its robustness against field observations. Furthermore, the proposed model 50 can be applied to assess and enhance the effectiveness of blue carbon capture and storage as part 51 revegetation measures to mitigate against global warming. 52 53 Keywords: photosynthesis; respiration; net ecosystem production; lagoon;

54 macrophyte

55

57 1. Introduction

58 Urgent adaptation and mitigation measures are required to address natural disasters, such as 59 floods, droughts, extreme heat, and landslides, which have been increasing due to global warming 60 (IPCC, 2014). Such action depends on effective adaptation and mitigation measures. Vegetated 61 shallow regions have recently been revealed to absorb and capture carbon dioxide by submerged 62 aquatic vegetation - a potential sink of anthropogenic carbon, termed "blue carbon" (Duarte et al., 63 2013). Nellmann (2009) showed the effectiveness of the net absorption of CO_2 by blue carbon 64 ecosystems, and it has been demonstrated that about 55 % of the carbon fixed by photosynthetic 65 activity on the Earth is in the form of blue carbon. Seagrass beds or meadows capture and store 66 autochthonous carbon from primary production and allochthonous carbon loaded from other areas 67 during the renewal of their epigeal stem and rhizome biomass (Kennedy et al., 2010; Fourqurean 68 et al., 2012). Therefore, there is the possibility that CO₂ is captured and stored in large amounts 69 in shallow water areas dominated by submerged aquatic vegetation (Macreadie et al, 2019). For 70 example, Beer et al. (1997) demonstrated that eelgrass plays a great role in carbon fixation and 71 Palacios et al. (2007) also demonstrated that seagrass meadows can be absorb CO_2 from the 72 atmosphere. However, Short (1999) indicated the difficulty in predicting the impact of global 73 climate change effects on seagrass communities, as it remains unclear how photosynthesis and 74 the productivity of aquatic plants will react to changes in the physical conditions of aquatic 75 ecosystems under a changing environment. Tada et al. (2018) demonstrated the potential for 76 application of a three-dimensional hydrodynamic model to evaluate the release and absorption of 77 CO2 by eelgrass (Zostera marina) in Komuke Lagoon of Hokkaido Island in Japan (Shintani and 78 Nakayama, 2010; Nakayama et al., 2012; Nakayama et al., 2014; Nakayama et al., 2016; 79 Nakayama et al., 2019). In this study, an attempt was made to model Dissolved Inorganic Carbon 80 (DIC) by simulating the effect of respiration and photosynthesis by eelgrass, where the CO_2 81 concentrations were estimated from DIC, Total Alkalinity (TA), water temperature and salinity 82 by assuming a chemical equilibrium state (per Zeebe et al., 2001). As a result, while DIC in this 83 lagoon varied in the range 300 µmol-C kg⁻¹ to 1300 µmol-C kg⁻¹, photosynthesis was found to 84 exceed respiration, which resulted in an estimated sink for DIC of 325 µmol-C kg⁻¹ (Tada et al., 85 2018). Additionally, Tada et al. (2018) showed that the influence of water temperature on 86 photosynthesis and respiration should be included in numerical models of DIC dynamics due to 87 its significance controlling eelgrass productivity.

Growth due to photosynthesis in eelgrass has been investigated in many previous studies based
 on photon flux density (often termed "photosynthetically active irradiance", PAR) estimates (see

90 review by Lee et al., 2007). A Jassby type equation was found to fit to growth rates in laboratory 91 and field experiments (Drew et al., 1979; Goodman et al., 1995; Holmer et al., 2001; Marsh et al., 92 1986; Olesen et al., 1993; Touchette, 1999; Zimmerman et al., 1995; Zimmerman et al., 1997). 93 However, the best model resoloving DIC changes in relation to photon flux density remains 94 unresolved. For investigating the effect of eelgrass photosynthesis on DIC dynamics, it is 95 necessary to measure daily changes in photon flux density, since DIC decreases greatly due to 96 photosynthesis during the daytime with a smaller increase in DIC due to respiration during the 97 night. Therefore, the duration of photon flux density has been shown to be more important than 98 the light compensation point and light saturation point in evaluating eelgrass production rates 99 (Dennison et al., 1982; Dennison et al., 1985). Additionally, Tada et al. (2018) revealed that 100 accurate topography must also be resolved for evaluating variability in DIC, based on predictions 101 using a three-dimensional hydrodynamic model. Overall, spatial distributions in underwater 102 photon flux density, nutrients, turbidity, and physical conditions have all been revealed to 103 contribute strongly to the net growth of eelgrass (Dennison et al., 1986). Beyond depth as a 104 primary determinant of growth, turbidity plays a great role in the photosynthesis of submerged 105 aquatic vegetation, as it increases light attenuation and reduces the photon flux density, which 106 results in suppression of the growth of eelgrass (Dennison, 1987; Moore et al., 1997). This 107 turbidity effect on productivity could be due to the effect of algal blooms reducing the effective 108 underwater photon flux density due to a high water column light extinction rate (Borum, 1985), 109 or due to excessive sediments brough about from inflows or resuspension (Adams et al., 2016). 110 Therefore, it is necessary to develop a DIC model which can resolve changes in photon flux 111 density precisely from the viewpoint of the suppression of photon flux density due to high 112 turbidity or algae blooms.

113 The effect of eelgrass photosynthesis on DIC has be shown to create site-specific patterns. For 114 example, in previous studies, the minimum light requirement has been investigated, showing large 115 variability from 5 % to about 40 % of the maximum irradiance (Dennison et al., 1993; Koch et al., 116 1996; Olesen et al., 1993). Additionally, the optimal growth temperature of eelgrass has been 117 shown to range broadly from 13 °C to 24 °C in temperate zones (Boström et al., 2004; Ibarra-118 Obando et al., 1987; Lee et al., 2005; Lee et al., 2006; Moore et al., 1996; Sand-Jensen, 1975; 119 Sfriso et al., 1998; Watanabe et al., 2005). In tropical or subtropical zones, the optimal growth 120 temperature of eelgrass has been shown to range from 16 °C to 30 °C (Biebl et al., 1971; Cabello-121 Pasini et al., 2003; Dennison, 1987; Drew, 1979; Evans et al., 1986; Marsh et al., 1986; Penhale, 122 1977). Therefore, it is necessary to model the effects of eelgrass photosynthesis and respiration 123 on DIC dynamics in order to reflect regional characteristics under different climates.

124 Related to respiration and photosynthesis of eelgrass, it has been revealed that seagrass growth 125 and production can be limited by nutrients even though photon flux density is plentiful (Coleman 126 et al., 1994; Dennison et al., 1987; Iizumi and Hattori, 1982; Murray et al., 1992; Orth, 1977; 127 Orth et al., 1983; Short et al., 1995; Thursby et al., 1982; Van Lent et al., 1995; Williams et al., 128 1993). There are many studies that have related the growth rate of eelgrass to nutrient supply from 129 the roots (Boström et al., 2004; Dennison et al., 1987; Iizumi et al., 1982; Jørgensen, 1982; 130 Mazzella et al., 1986; Moore et al., 1996; Pedersen et al., 1993; Short, 1987). The contribution of 131 sediment nutrients to eelgrass growth rates has been found to be equal or be more than that from 132 the water column (Short et al., 1984; Pedersen et al., 1992). Furthermore, Capone (1982) showed 133 that nutrient concentrations vary greatly due to the presence of eelgrass, and nutrient pools have 134 been shown to have rapid turnover rates. However, Burkholder et al. (1992, 1994) demonstrated 135 that high nutrient enrichment inhibited eelgrass growth due to carbon limitation, phosphorus 136 limitation, or other internal nutrient imbalances leading to physiological effects (Burkholder et 137 al., 1992; Burkholder et al., 1994). Zimmerman et al. (1987) demonstrated the application of a 138 numerical model to evaluate how nutrients affect the ecological functions of eelgrass, including 139 respiration and photosynthesis. Therefore, in the context of nutrient supply, it can again be said 140 that it is necessary to develop a detailed and precise submerged aquatic vegetation model for 141 evaluating CO₂ flux.

142 Aside from light, photosynthesis parameters change seasonally due to differences in growth 143 rate and water temperatures (Dennison, 1987; Orth et al., 1986; Phillips et al., 1983), with water 144 temperature considered the primary factor controlling seasonal growth (Bulthuis, 1987; Setchell, 145 1929). Martin et al. (2006) showed that net CO_2 flux increases with the seasonal increase in water 146 temperatures in the Bay of Brest, Western Brittany, France, with similar patterns in net O₂ flux. 147 However, very limited studies have investigated how this temperature variability affects DIC 148 fluxes. As submerged aquatic vegetation tends to grow best in calm, stratified coastal areas such 149 as enclosed bays and lagoons, methods resolving the interaction between hydrodynamics, 150 vegetation and carbon are required for application to these types of ecosystems. Therefore, this 151 study aims to investigate thermal effects, considering irradiance, on changes in DIC fluxes under 152 nutrient-rich (non-limiting) conditions. We develop a simplified and practical model to 153 understand DIC dynamics, based on easily measurable experimental data, that can be applied to 154 estimate seasonal productivity. The model is verified through application to Komuke Lagoon, 155 Japan, where it is used to estimate carbon uptake by eelgrass.

157 2. Method

158 2.1. Laboratory experiments

159 Komuke Lagoon is connected to the Sea of Okhotsk through a tidal inlet and has a volume of 160 approximately 3,333,000 m³ and a maximum water depth of 3 m (Fig. 1). Although the tidal range 161 of the Sea of Okhotsk is approximately 1 m, the tidal range of Komuke Lagoon is only 0.5 m 162 because of the narrow 15 m width of the tidal inlet. For this reason, calm water conditions are 163 usual in Komuke Lagoon and the eelgrass population is widely distributed. A laboratory 164 experiment evaluating photosynthesis was conducted with lagoon water and eelgrass collected on the 25th of June 2018 at the sampling point (44° 15' 7.3″ E, 143° 30' 38.9″ N) shown in Fig. 165 166 1. The experiment was conducted outdoors in fine weather using a transparent acrylic water tank 167 with a depth of 150 cm, a width of 20 cm, and a length of 20 cm (Fig. 2). Four shoots of eelgrasses 168 taken from the lagoon were deployed in the water tank. The roots of the eelgrass were covered 169 with vinyl sheets to avoid oxygen consumption by oxygen-demanding substances contained in 170 the sediment. The salinity of the water used in the experiment was 22 psu. The dimensions (leaf 171 width, length, and thickness) of eelgrass used in experiment were measured before 172 experimentation (Table 1).

173 Two thermometers (Eijkelkamp Co. Ltd., SERA Diver) and two light quantum loggers (JFE 174 Advantec Co. Ltd., DEFI2-L) were deployed in the water tank, recording every ten minutes. The 175 experiment started at night, when photosynthesis was inactive, and continued for twenty-four 176 hours. To provide vertically uniform conditions for eelgrass in the water tank, the water was well 177 mixed using a peristatic pump with a flow rate of 7 mL min⁻¹ through a silicon tube with a diameter 178 of 2 mm; the water was gently taken from an inlet at a height of 100 cm from the tank bottom 179 through the silicon tube and discharged from an outlet at a height of 0 cm from the tank bottom. 180 The uniformity of water quality profiles in the water tank was confirmed by measuring 181 temperatures periodically.

Approximately 200 mL water samples were taken from the water tank into a Scott Duran bottle by syringe and tube apparatus every hour. After sampling, 200 μL of a saturated aqueous solution of mercuric chloride was quickly added to the sample to prevent biological activity. DIC and TA of the water sample was then measured using a Total Alkalinity meter (Kimoto Electronic Co. Ltd., ATT-15) within 2 hours of sampling. Titrant for volumetric analysis (Kanto Kagaku, Co. Ltd., 0.1 mol L⁻¹ hydrochloric acid) was used for DIC and TA measurement by high accuracy titration. Additionally, a water tank experiment under the same conditions but without eelgrass was conducted as a control experiment. Concentrations of NO₃-N, NO₂-N, NH₄-N, and PO₄-P in
water samples were measured for each experiment using an auto-analyzer (BLTEC Co. Ltd.,
QuAAtro 2HR).

The wet weight, dry weight, and carbon content of four eelgrass used in the laboratory experiment were measured separately for leaves and stems. Dry weight was measured after heating by an electric oven at 105°C for 24 hours. Approximately 2 mg of dried sample was ground using a mortar and pestle and elemental carbon content analyzed using a CHNS analyzer (Parkin Elmer Co. Ltd., 2400II) in CHN mode to estimate eelgrass biomass.

197

198 2.2. Evaluation of coefficients of NEP model

199 Photon flux density and water temperature were measured every 10 minutes. The 10-minute 200 data were converted to 1-hour interval data. Nitrogen and phosphorus could limit eelgrass 201 respiration and photosynthesis when sediment water NH_4^+ , $NO_3^- + NO_2^-$ and PO_4^{3-} concentrations are lower than 0.1, 0.5 and 0.03 μ M, respectively (Lee et al., 2007). We confirmed 202 203 that nitrogen and phosphorus concentrations were sufficient and didn't limit the eelgrass 204 respiration and photosynthesis (Table 2). Therefore, in this study, we decided to apply Eq. (1) for 205 modelling Net Ecosystem Production by eelgrass (NEP) taking into account photon flux density 206 and water temperature. A Jassby type equation and Arrhenius equation were applied to simulate 207 photosynthesis related to photon flux density and water temperature (Staehr et al., 2011; Beca-208 Carretero et al., 2018; Burkholz et al., 2019), where NEPR refers to Net Ecosystem Production 209 due to Respiration by eelgrass and NEPP to Net Ecosystem Production due to Photosynthesis by 210eelgrass:

$$\frac{d}{dt}(\text{DIC}) = \Delta \text{DIC} = R_A \exp\left(-\frac{E_{aR}}{T_w R}\right) - P_{\Psi} \tanh\left(\frac{\alpha_{\Psi}I}{P_{\Psi}}\right) R_P \exp\left(-\frac{E_{aP}}{T_w R}\right)$$

$$= -\text{NEP} = \text{NEPR} - \text{NEPP}$$
(1)

211 where, DIC μ mol L⁻¹ is the dissolved inorganic carbon, $R_A \mu$ mol kg⁻¹ h⁻¹ is the parameter for 212 respiration, E_{aR} m² kg s⁻² is the activation energy for respiration, T_w K is the water temperature, 213 R m² kg s⁻² K⁻¹ is the Boltzmann constant (1.380649 x 10⁻²³), $P_{\psi} \mu$ mol kg⁻¹ h⁻¹ and α_{ψ} m² s kg⁻¹ 214 h⁻¹ are the parameters for photosynthesis, $I \mu$ mol m⁻² s⁻¹ is the photon flux density, R_P is the 215 parameter for photosynthesis, E_{aP} m² kg s⁻² is the activation energy for photosynthesis. 216 Firstly, the parameters for the respiration term in Eq. (1), $R_A \mu$ mol kg⁻¹ h⁻¹ and E_{aR} , were

217 obtained using the fact that photosynthesis activity is negligible from sunset to sunrise when 218 photon flux density is zero. We calculated Δ DIC under three water temperatures: 12 °C, 20 °C and 24 °C in order to evaluate the effect of the change in water temperature on eelgrass respiration in dark conditions. Secondly, the parameters for photosynthesis was obtained by considering photon flux density and using the R_A and E_{aR} for NEPR. Thirdly, we proposed an eelgrass Seasonal Net Ecosystem Production (SNEP) model using the photon flux density duration recorded in the field. Finally, we applied the SNEP model to estimate changes in DIC due to eelgrass in Komuke Lagoon.

225

226 3. Results

227 3.1. Biological characteristics of eelgrass

228 The dry weight and biomass (carbon content) of eelgrass used in the laboratory experiment are 229 shown in Table 1 and nutrient concentrations in the lagoon water are shown in Table 2. The 230 biomass shown in Table 1 was based on the carbon content per unit leaf area calculated from the 231 elemental carbon content and measured size of eelgrass. It should be noted that one data point 232 was not available due to a measurement error. Mean leaf biomass was about 288 g C using the leaf area of both sides of the leaf m². Although the concentration of NO₃-N, NO₂-N, NH₄-N, and 233 234 PO₄-P decreased during the experiment, photosynthetic rates were not limited by nutrients 235 because concentrations remained sufficiently high (Lee et al., 2007).

236

237 3.2. Thermal effect on NEP

238 Partial pressures of carbon dioxide in water (fCO₂) started increasing at 8:00 pm on the 25th of 239 June, and fCO_2 decreased from 5:00 am on the 26th. DIC changed in a pattern similar to fCO_2 (Fig. 240 3). In contrast to the eelgrass experiments, in the control tank, DIC was almost constant, which 241 suggests that water column photosynthesis and respiration due to organisms other than eelgrass, 242 such as phytoplankton, were negligible in terms of changes in DIC. Also, since TA changed only 243 slightly, we found that the eelgrass predominantly affected DIC rather than TA. The maximum 244 photon flux density occurred at 11:00 am on the 26th, when water temperature had increased from 245 12 to 27 °C. ΔDIC (=-NEP) µmol kg⁻¹ h⁻¹ was negative from 5:00 am on the 25th to 5:00 pm on 246 the 26th, which demonstrated positive net ecosystem production (Fig. 3f).

When NEP was plotted against photon flux density, NEP values showed different tendencies
below and above 250 μmol m⁻² s⁻¹ (Fig. 4). Red circles in Fig. 4 indicate 5 samples from 7:00 am
to 11:00 am on the 26th, and green circles indicate 5 samples from 0:00 pm to 4:00 pm on the 26th.
Red and green circle samples were sampled under mean water temperature of 16 °C and 25 °C,
respectively. Therefore, it is noted that mean water temperature is a key factor controlling the

tendencies of NEP, which suggests the development of a NEPP parameterization needs to includethe effect of water temperature in Eq. (1).

254

255 3.3. Estimation of coefficients of NEPR

256 Values for R_A and E_{aR} were obtained by assuming that Δ DIC represents NEPR from 7:00 pm of the 25th to 3:00 am of the 26th and from 6:00 pm to 7:00 pm of the 26th when photosynthetic 257 258 activity is negligible (Fig. 5). It should be noted that only data with the condition $\Delta DIC>0$ was 259 applied because the respiration term, NEPR, should be higher than zero. Since ADIC from the 260 25th to 26th of June in 2018 was obtained when the water temperature was about 20 °C, we conducted separate laboratory experiments on the 10th of September 2018 with different water 261 262 temperature conditions. Finally, 9 validated samples were obtained, in which a significant 263 difference was confirmed to exist at the 5 % level using a Kruskal Wallis test. Although there are 264 some fluctuations evident in Figure 5, parameters for respiration could be obtained based on strong relationships with water temperature: R_A =1.04 x 10¹⁷ µmol kg⁻¹ h⁻¹ and E_{aR} =1.52 x 10⁻¹⁹ 265 $m^2 kg s^{-2}$ with $R^2 = 0.68$. 266

267

268 3.4. Estimation of coefficients of NEPP

By applying R_A =1.04 x 10¹⁷ μ mol kg⁻¹ h⁻¹ and E_{aR} =1.52 x 10⁻¹⁹ m² kg s⁻² into NEPR of Eq. (1), NEPP was obtained using Eq. (2):

$$NEPP = -\Delta DIC + NEPR = -\Delta DIC + R_A \exp\left(-\frac{E_{aR}}{T_w R}\right)$$

and the relationship between photon flux density and NEPP is shown in Fig. 6.

To investigate thermal dependence of eelgrass photosynthesis, a value that ignores thermal effects, NEPP_{*noT*}, was plotted against photon flux density using Eq. (3) (solid lines in Fig. 6):

$$NEPP_{noT} = P_{\psi} \tanh\left(\frac{\alpha_{\psi}I}{P_{\psi}}\right)$$
(3)

(2)

when the parameters are given as $P_{\psi noT}$ =21.5 µmol kg⁻¹ h⁻¹ and $\alpha_{\psi noT}$ =21.5/200 m² s kg⁻¹ h⁻¹, comparison between NEPP and NEPP_{noT} revealed a large difference in the tendencies between afternoon (green) and morning (red) conditions; red circles represented water samples from temperatures ranging from 12 to 20 °C, and green circles from 22 to 27 °C. The higher the water temperature, the larger the NEPP and vice versa, which is similar to the differences in the tendencies of NEPR. Therefore, an attempt was made to correct for the effect of temperature on the parameters for photosynthetic activity, R_P and E_{aP} , using NEPP / NEPP_{noT}, (Fig. 7):

$$\frac{\text{NEPP}}{\text{NEPP}_{noT}} = R_P \exp\left(-\frac{E_{aP}}{T_w R}\right) \tag{4}$$

where $R_P=2.30 \ge 10^7$ and $E_{aP}=0.69 \ge 10^{-19} \le 10^{-19} \le 10^{-2} \le 10^{-19} \le 10^{-2} \le 10^{-19} \le 10$

287

288 3.5. Proposal of the SNEP model

Eq. (1) can not only be applied to analyze local-scale NEP, for example though coupling with a three-dimensional hydrodynamic model, but it can also be applied to estimate the long-term (seasonal) change in DIC in enclosed waterbodies. To this end, we propose a method to estimate Seasonal Net Ecosystem Production (SNEP) by adapting Eq. (1). To estimate SNEP, it is necessary to include the hourly change in DIC (Dennison et al., 1982; Dennison et al., 1985), and SNEP is defined by integrating Eq. (1) for one day:

$$SNEP = -R_A \frac{1}{t_{1day}} \int \exp\left(-\frac{E_{aR}}{T_w R}\right) dt + P_{\Psi} \frac{1}{t_{1day}} R_P \int \tanh\left(\frac{\alpha_{\Psi} \beta_I I_S}{P_{\Psi}}\right) \exp\left(-\frac{E_{aP}}{T_w R}\right) dt$$
(5)

295 where, t_{1day} is the integration period (= 24 h), β_I is the effectiveness coefficient (0: no photon flux

density to 1: maximum photon flux density), and I_s is the photon flux density at water surface.

297 By assuming that water temperature changes sinusoidally:

$$T_w = T_m + T_h = T_m + T_{h0} \sin\left(\frac{2\pi}{24}t - \frac{\pi}{2}\right)$$
(6)

Eq. (5) can be simplified to:

$$\text{SNEP} \approx -R_A \exp\left(-\frac{E_{aR}}{T_m R}\right) \gamma_R + P_{\Psi} \exp\left(-\frac{E_{aP}}{T_m R}\right) \gamma_P \tag{7}$$

where

$$\gamma_R = \frac{1}{t_{1day}} \int \exp\left(\frac{E_{aR}T_h}{T_m^2 R}\right) dt$$
(8)

$$\gamma_P = \frac{1}{t_{1day}} \int \tanh\left(\frac{\alpha_{\Psi}\beta_I I_S}{P_{\Psi}}\right) \exp\left(\frac{E_{aP}T_h}{T_m^2 R}\right) dt$$
(9)

where, T_m Celsius degree is the daily mean water temperature, $T_h = T_w - T_m$, T_{h0} is the amplitude of water temperature in a day, and *t* h is the time from 0 to 24 h. It should be noted that it is necessary to include the influence of topographical and tidal effects on a specific location because the photon flux density reaching the canopy may change significantly due to changes in bathymetry and tides. These effects can be included by changing the value of β_l , which is the parameter for attenuation of photon flux density and is a function of the extinction coefficient for light attenuation (Dennison et al., 1986).

To estimate and project dissolved oxygen (DO) over the bottom in an enclosed bay, Nakayama et al., (2010) demonstrated the suitability of a conceptual DO model by taking into account residence time between the inside of the bay and the outer ocean (Okada and Nakayama, 2007; Okada et al., 2011; Sato et al., 2012). This study thus makes an attempt to propose a SNEP model to estimate DIC using the SNEP approach and accounting for water exchange (Fig. 8):

$$V_0 \frac{d}{dt} (\text{DIC}_S) = -V_0 \text{SNEP} - (Q_E + Q_R) \text{DIC}_S + Q_E \text{DIC}_{out} + Q_R \text{DIC}_R$$
(10)

310 where, $DIC_S \mu mol kg^{-1}$ is the monthly mean DIC, $V_0 m^3$ is the volume of the lagoon, $Q_E m^3 s^{-1}$ is 311 the exchange flux between a lake and the ocean, $Q_R m^3 s^{-1}$ is the river discharge, $DIC_R \mu mol kg^{-1}$ 312 is DIC of a river, and $DIC_{out} \mu mol kg^{-1}$ is the typical DIC concentration of the ocean. We note that 313 this model assumes the net atmospheric exchange of CO₂ over the period of the calculation is 314 negligible, which is supported by estimates from in situ *f*CO₂ values.

By assuming a steady state in Eq. (10) (i.e. the left term is zero), the change in DIC_s can be obtained as:

$$\Delta \text{DIC}_S = \text{DIC}_{stable} - \text{DIC}_S = t_S \text{SNEP}$$
(11)

$$t_S = \frac{V_0}{Q_E + Q_R} \tag{12}$$

$$DIC_{stable} = \frac{Q_E DIC_{out} + Q_R DIC_R}{Q_E + Q_R}$$
(13)

317 where, $\Delta DIC_s \ \mu mol \ kg^{-1}$ is the change in DIC with NEP, $DIC_{stable} \ \mu mol \ kg^{-1}$ is DIC in a steady 318 state without NEP (NEP = 0), and t_s h is the residence time of the target domain. In other words, 319 if we obtain residence time, t_s , and SNEP, the change in DIC due to eelgrass can be estimated 320 using Eq. (11), and used as a measure of system scale carbon uptake.

322 4. Discussion

323 In this study, we obtained parameters needed to simulate the response of photosynthesis to 324 photon flux density and water temperature based on a Jassby type P-I equation and an Arrhenius 325 equation (Staehr et al., 2011; Beca-Carretero et al., 2018; Burkholz et al., 2019). Lee et al. (2007) 326 found that laboratory experiments tend to underestimate saturation irradiance because of the usage 327 of leaf segments. Since we did not remove the roots in our laboratory experiments, the saturation irradiance was estimated as 200 μ mol m⁻² s⁻¹ by $P_{\psi noT} / \alpha_{\psi noT}$, where $P_{\psi noT}$ =21.5 μ mol kg⁻¹ h⁻¹ 328 329 and $\alpha_{\psi no7} = 21.5/200 \text{ m}^2 \text{ s kg}^{-1} \text{ h}^{-1}$. The value is larger than the average level of saturation irradiance 330 for leaf segments of Zostera marina, 116 µmol m⁻² s⁻¹, while the average value of saturation 331 irradiance for whole plant is 450 μ mol m⁻² s⁻¹ (Lee et al., 2007).

332 The rate of change in NEP due to photosynthesis and respiration has been shown to increase 333 with increasing water temperature (Dennison, 1987; Orth et al., 1986; Phillips et al., 1983; 334 Bulthuis, 1987; Setchell, 1929). To understand thermal dependencies, NEPP, NEPR and NEP 335 were computed by changing water temperatures from 5 °C to 30 °C using Eq. (1) (Fig. 9). NEPP 336 and NEPR increased with the increasing water temperature (Fig. 9a and Fig. 9b). When the photon 337 flux density was more than about 250 μ mol m⁻² s⁻¹, NEPP varied from 8 μ mol kg⁻¹ h⁻¹ to 34 μ mol kg⁻¹ h⁻¹, an increase of 26 µ mol kg⁻¹ h⁻¹ (Fig. 9a). In comparison, NEP under a thermal 338 increase from 5 to 30 °C varied from 7 μ mol kg⁻¹ h⁻¹ to 17 μ mol kg⁻¹ h⁻¹, an increase of 10 μ mol 339 340 kg⁻¹ h⁻¹ (Fig. 9c). Furthermore, when photon flux density was more than 250 μ mol m⁻² s⁻¹, NEP 341 was almost constant under a thermal increase from 25 to 30 °C because the increase in NEPR was 342 almost equal to the increase in NEPP (Fig. 9c). Lee et al. (2007) revealed that the average value 343 of optimal temperature of Zostera marina is about 15 °C for growth and 23 °C for photosynthesis, 344 which is in good agreement with our study with optimal temperatures of about 25 °C for 345 photosynthesis.

346 We applied the SNEP model to Komuke Lagoon to explore the applicability for the evaluation 347 of seasonal changes in DIC. We referred to the field observations which were conducted in 348 Komuke Lagoon in May 2013 (Tada et al., 2018) and August 2018. May and August were chosen 349 as target months in order to clarify the seasonal effect of the difference in water temperature 350 between spring and summer. Firstly, γ_R and γ_P for SNEP were estimated from Eqs. (8) and (9) 351 by giving 5 different β_l : 100 %, 75%, 50 %, 25 % and 10 % (Table 3). This showed that γ_P 352 varied greatly with the change in β_I . In contrast to β_I , the difference in photon flux density 353 between May and August was negligible and the values for γ_P were the same between the two 354 seasons. Since eelgrass meadows exist from the bottom to a water depth of about 1.0 m in Komuke

Lagoon, and the extinction coefficient was about 2.0 m⁻¹ including the effect of turbidity with a mean water depth of 2.0 m, β_I was obtained as 25 % using Eq. (14):

$$\beta_{I} = \frac{1}{h_{m}} \int_{0}^{l_{E}} \exp[-k_{E}(h_{m} - z)] dz$$
(14)

357 where, h_m m is the mean water depth, k_E m⁻¹ is the extinction coefficient, and l_E m is the deflected 358 vegetation height.

SNEP was computed against water temperature, T_w , from $\beta_I = 0\%$ to $\beta_I = 100\%$ (Fig. 10). 359 Although the SNEP did not change much from $\beta_I = 50$ % to $\beta_I = 100$ %, SNEP decreases 360 greatly from $\beta_I = 50$ % to $\beta_I = 0$ %. When β_I was 25 %, a peak in SNEP appeared at a water 361 362 temperature of 15 °C and SNEP became negative when the water temperature was more than 363 28 °C. Watanabe et al. (2005) demonstrated that optimal temperature for Zostera marina is 364 16.1 °C in Akkeshi Bay, which is located close to Komuke Lagoon. This may suggest the practical 365 applicability of the relatively simple SNEP model for real field environments. Tada et al. (2018) 366 revealed that DIC at the water surface, DIC_s, was smaller than the no eelgrass stable condition, DIC_{stabl}, in Komuke Lagoon on the 15th of May 2013 using a three-dimensional hydrodynamic 367 368 model, which shows $\Delta DIC_S = DIC_{stabl}$ - $DIC_S = 325 \mu$ mol kg⁻¹ (Table 4). The residence time of 369 Komuke Lagoon was $t_S = 110$ h and the mean water temperature was 7.1 °C. Since SNEP is 2.82 370 μ mol kg⁻¹ h⁻¹ from Fig. 10, t_s SNEP can be obtained as 310 μ mol kg⁻¹. Therefore, the estimated 371 t_s SNEP = 310 μ mol kg⁻¹ approximately agreed with Δ DIC_s = 325 μ mol kg⁻¹, which suggests 372 the applicability of the SNEP model to an enclosed waterbody, when the residence time is known. 373 In addition to the estimation of t_s SNEP in May 2013, we conducted water sampling at the 374 water surface in Komuke Lagoon and measured water temperature, salinity and DIC in 2018 in 375 order to investigate the applicability of the SNEP model Eq. (11) (sampling stations are shown in 376 Fig. 1). It should be noted that we found no significant differences in water quality and flow fields 377 between 2013 and 2018 since Komuke Lagoon has a very narrow inlet and the Okhotsk Sea 378 provides stable periodic annual changes in water quality. The mean water temperature was 23.5 °C, 379 which corresponds to SNEP = 2.23 μ mol kg⁻¹ h⁻¹ using β_I = 25 % from Fig. 10 (Table 4). As the 380 residence time is about 110 h, t_s SNEP was found to be 245 μ mol kg⁻¹. Using the low-salinity 381 endmember of rivers and the high-salinity endmember of the ocean, ΔDIC_S was estimated as 255 382 μ mol kg⁻¹ from field observations following Zeebe et al. (2001). Again, the estimated t_S SNEP = 383 245 μ mol kg⁻¹ agreed well with Δ DIC_S = 255 μ mol kg⁻¹ from the field observations. The proposed 384 SNEP model is specific to enclosed or semi-enclosed waterbodies and not for suitable for the 385 open ocean because of the requirement to evaluate residence times. Since eelgrass appears to be

- adapted to calm conditions, and there is the possibility to obtain residence times accurately for
 these waterbodies, this suggests the SNEP model can be useful to improve predictions of carbon
 capture and storage by eelgrass populations in protected waters.
- 389

390 5. Conclusion

391 Aiming to understand the NEP of eelgrass in Komuke Lagoon, which is located in the east of Hokkaido, parameters for NEPR were obtained as $R_A=1.04 \times 10^{17} \mu \text{ mol kg}^{-1} \text{ h}^{-1}$ and $E_{aR}=1.52 \text{ x}$ 392 10⁻¹⁹ m² kg s⁻² using the hourly change in DIC without photon flux density in laboratory 393 394 experiments. The photosynthetic activity of eelgrass was confirmed to be affected by water 395 temperature from laboratory experiments, and parameters for NEPP were obtained as $P_{\psi}=21.5 \mu$ mol kg⁻¹ h⁻¹, α_{ψ} =21.5/200 m² s kg⁻¹ h⁻¹, R_P =2.30 x 10⁷ and E_{aP} =0.69 x 10⁻¹⁹ m² kg s⁻². The SNEP 396 397 model was proposed, in which the seasonal change in DIC, ΔDIC_s , can be estimated as t_s SNEP 398 by using the residence time, t_s . The SNEP model was applied to two sets of field observations 399 from Komuke Lagoon, May 2013 and August 2018, and the estimated seasonal change in DIC, t_s 400 SNEP, was confirmed to agree with ΔDIC_s from field observations. In other words, the seasonal 401 change in NEP can be obtained as $\Delta DIC_S / t_S$ using the residence time, t_S , and the seasonal change 402 in DIC, Δ DIC₈, from endmember analysis using field observations. Since the development and 403 implementation of mitigation measures against global warming is an urgent issue, the proposed 404 SNEP model in this study may be applied to assist efforts to enhance effective capture and storage 405 of carbon dioxide through seagrass restoration. Also, the SNEP model may be useful for assessing 406 changes due to climate change, e.g. as a screening tool to investigate whether the warming trend 407 improves net productivity or not with limited information.

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- 412 Acknowledgments
- 413 This work was supported by the Japan Society for the Promotion of Science under grant
- 414 18H01545 and 18KK0119.
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608	Figure
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610	Fig. 1 Sampling stations in Komuke Lagoon
611	
612	Fig. 2. Schematic diagram of the laboratory experiment tank containing eelgrass
613	
614	Fig. 3. Laboratory experiment results from $21:00$ on the 25^{th} to $19:00$ on the 26^{th} of June 2018.
615	(a) fCO_2 with and without eelgrass. (b) DIC with and without eelgrass. (c) TA with and without
616	eelgrass. (d) photon flux density. (e) water temperature. (f) Δ DIC.
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618	Fig. 4. Photon flux density and NEP.
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620	Fig. 5. Water temperature and Δ DIC. Solid line is the first term of equation (1).
621	
622	Fig. 6. Photon flux density and NEPP by equation (2). Solid line is equation (3).
623	
624	Fig. 7. Water temperature and the ratio of NEPP and NEPP _{noT} . Solid line is equation (4).
625	
626	Fig. 8. Schematic diagram of SNEP model.
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628	Fig. 9. Effect of water temperature on (a) NEPP, (b) NEPR and (c) NEP.
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630	Fig. 10. Water temperature and DNEP in Komuke Lagoon.
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634	Table
635	
636	Table 1. Specimens in laboratory experiments.
637	
638	Table 2. Nutrient concentrations in experimental water.
639	
640	Table 3. γ_R and γ_P in May 2013 and August 2018.
641	
642	Table 4. Field observations in Lake Komuke.
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644	



Fig. 1 Sampling stations in Komuke Lagoon





649 Fig. 2. Schematic diagram of the laboratory experiment tank containing eelgrass









Fig. 5. Water temperature and Δ DIC. Solid line is the first term of equation (1).





Fig. 6. Photon flux density and NEPP by equation (2). Solid line is equation (3).





670 Fig. 7. Water temperature and the ratio of NEPP and NEPP_{noT}. Solid line is equation (4).







Fig. 10. Water temperature and DNEP in Komuke Lagoon.

Eelgrass	Dry weight	Leaf biomass
No.	(leaf) (g)	$(g C/m^2)$
1	379.94	253.15
2	419.07	284.27
3	490.96	325.87

Table 1. Specimens in laboratory experiments.

Water sample	NO ₃ -N	NO ₂ -N	NH4-N	PO ₄ -P
	(µM)	(µM)	(µM)	(µM)
Before experiment	1.642	0.100	2.127	0.804
After experiment	0.899	0.079	0.671	0.565

Table 2. Nutrient concentrations in experimental water.

Table 3. γ_R and γ_P in May 2013 and August 2018.

βι	100%	75%	50%	25%	10%	0%
γ_R	1.01					
γ_P (May in 2013)	0.56	0.54	0.51	0.42	0.28	0
γ_P (August in 2018)	0.54	0.52	0.49	0.42	0.28	0
γ_P (mean)	0.55	0.53	0.50	0.42	0.28	0

Table 4. Field observations in Lake Komuke.

	Mean water	$t_{S}(h)$	ΔDIC_S	SNEP	t _S SNEP
	temperature		(µ mol kg ⁻¹)	$(\mu \text{ mol } \text{kg}^{-1} \text{ h}^{-1})$	$(\mu \text{ mol kg}^{-1})$
	(Celsius		from observation	from (7)	
	degree)				
May 2013	7.1	110	325	2.82	310
August 2018	23.5	110	255	2.23	245