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Seasonal metabolism and carbon export potential of a key coastal habitat: The perennial canopy-forming macroalga *Fucus vesiculosus*

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**Abstract**

The important role of macroalgal canopies in the oceanic carbon (C) cycle is increasingly being recognized, but direct assessments of community productivity remain scarce. We conducted a seasonal study on a sublittoral Baltic Sea canopy of the brown alga *Fucus vesiculosus*, a prominent species in temperate and Arctic waters. We investigated community production on hourly, daily, and seasonal timescales. Aquatic eddy covariance (AEC) oxygen flux measurements integrated ~ 40 m² of the seabed surface area and documented considerable oxygen production by the canopy year-round. High net oxygen production rates of up to 35 ± 9 mmol m⁻² h⁻¹ were measured under peak irradiance of ~ 1200 μmol photosynthetically active radiation (PAR) m⁻² s⁻¹ in summer. However, high rates > 15 mmol m⁻² h⁻¹ were also measured in late winter (March) under low light intensities < 250 μmol PAR m⁻² s⁻¹ and water temperatures of ~ 1 °C. In some cases, hourly AEC fluxes documented an apparent release of oxygen by the canopy under dark conditions, which may be due to gas storage dynamics within internal air spaces of *F. vesiculosus*. Daily net ecosystem metabolism (NEM) was positive (net autotrophic) in all but one of the five measurement campaigns (December). A simple regression model predicted a net autotrophic canopy for two-thirds of the year, and annual canopy NEM amounted to 25 mol O₂ m⁻² yr⁻¹, approximately six-fold higher than net phytoplankton production. Canopy C export was ~ 0.3 kg C m⁻² yr⁻¹, comparable to canopy standing biomass in summer. Macroalgal canopies thus represent regions of intensified C assimilation and export in coastal waters.

Macroalgae form extensive canopies within near-shore areas that are among the most productive and biodiverse habitats on Earth (Mann 1973). Despite covering just ~ 7 million km², or < 2% of the total ocean area (Charpy-Roubad and Soumia 1990; Gattuso et al. 2006; Krause-Jensen and Duarte 2014), and accounting for < 5% of total oceanic net primary production (Field et al. 1998; Duarte 2017), macroalgal canopies represent “hotspots” of photosynthetic biomass. On a local scale, the biomass per unit area occupied by macroalgae can be approximately 400-fold greater than that of phytoplankton, and on a global scale, macrophyte biomass represents about two-thirds of total oceanic photosynthetic biomass (Smith 1981). Macroalgal beds typically produce organic carbon well in excess of their ecosystem requirements (net autotrophic), and it is estimated that a large proportion of the organic material (~ 40%) is exported to the surroundings as particulate and dissolved matter (Duarte and Cebrian 1996). Drift macrophytes constitute significant enrichment of organic C to the benthos and provide important spatial subsidies that support high levels of secondary production in adjacent communities (Vetter 1994; Harrold et al. 1998; Filbee-Dexter and Scheibling 2014), with episodic events such as high winds and storms facilitating massive C export fluxes (Dierssen et al. 2009). On a global scale, macroalgae have a large C sequestration capacity. Preliminary analyses estimate that around 173 Tg C yr⁻¹ is sequestered, a value that exceeds C sequestration by angiosperm-based aquatic habitats (Krause-Jensen and Duarte 2016).

The important role of marine macroalgae for coastal productivity was realized more than four decades ago (Mann 1973). Despite this, productivity rates of macroalgal canopies and their role in the ocean C budget remains poorly quantified (Krause-Jensen and Duarte 2016; Duarte 2017). Performing detailed in situ studies on the production, degradation, and export potential of organic C within macroalgal canopies is challenging: traditional flux measurement techniques such as benthic chambers are difficult to apply to rocky outcrops...
and coarse sediment deposits, where these habitats often prevail (Glud et al. 2010). Additionally, chambers enclose only a small surface area of the canopy (~ 0.1 m²), incubation times are short (~ 1 h) (Tait and Schiel 2010; Bordeyne et al. 2017), and the enclosure process excludes natural variations in hydrodynamics that may affect photosynthetic rates of the enclosed algae (Mass et al. 2010). Biomass addition techniques (e.g., Ziemann 1974) have been applied extensively to study productivity rates of submerged vegetation. However, this method is time-consuming, destructive, underestimates true productivity rates, and cannot resolve short-term (hourly to daily) dynamics (Long et al. 2015a).

The aquatic eddy covariance (AEC) technique is a recent technological development that overcomes many of these issues. It quantifies benthic oxygen fluxes noninvasively, includes a large surface area of the seabed (tens of square meter) in its measurement and produces benthic fluxes for consecutive 15 min time periods over 24 h or more (Berg et al. 2003; Berg et al. 2007). From this, the daily net ecosystem metabolism (NEM), representing the integrated balance between habitat primary productivity versus respiration in a 24 h period, can be calculated either as the sum of daytime and nighttime fluxes weighted to the hours of day and night, respectively (Long et al. 2013; Attard et al. 2014), or as an integration of the AEC fluxes over 24 h (Hume et al. 2011; Rheuban et al. 2014b; Attard et al. 2015). Daily NEM O₂ fluxes can then be converted to C equivalents. Positive NEM values indicate a surplus of organic C production (net autotrophy), which results in accumulation of vegetation biomass within the habitat and/or export of particulate or dissolved matter to its surroundings. When the habitat is consuming more organic C than is being produced, NEM is negative. Therefore, by quantifying the NEM and the canopy standing biomass over time, it is possible to estimate the organic C export rate, which permits novel assessments on the role of macroalgal canopies within the marine C cycle.

_**Fucus vesiculosus***(Linnaeus, 1753), commonly known as “bladder wrack,” is a prominent species in temperate and Arctic coastal waters and it forms extensive canopies in the central Baltic Sea region (Råberg and Kautsky 2007). _F. vesiculosus_ canopies occur most commonly at moderately exposed sites, with a maximum depth distribution limit located at ~ 6% of surface irradiance levels (Tom et al. 2006). Gas spaces on its fronds not only maintain canopy posture and facilitate spore dispersal but also accumulate considerable oxygen under illumination (Aleem 1969). Fucoid beds harbor greater macrofaunal abundance and biomass of small crustaceans than nonfucoid vegetated sites (Råberg and Kautsky 2007) and are important foraging grounds for commercially important fish such as perch (**Perca fluviatilis** (Lappalainen et al. 2001). Despite the widespread occurrence of the species, _F. vesiculosus_ canopies vary in their spatial extent and depth distribution in response to environmental drivers. In the Baltic Sea, canopies of _F. vesiculosus_ have undergone large changes in extent and depth distribution (Kautsky et al. 1986; Torn et al. 2006). High nutrient loading of coastal waters has led to eutrophication and a decrease in sunlight transmittance, favoring instead proliferation of opportunistic ephemeral macroalgae such as the green alga _Ulva intestinalis_ and the free-living brown alga _Pilayella littoralis_. In contrast, in the Arctic, the species is expected to expand to higher latitudes as the sea-ice cover diminishes (Krause-Jensen and Duarte 2014). The impacts this has had on the biodiversity of seafloor communities, including species’ abundance, community structure, and living-mode of the benthos (mobility, feeding capabilities, and predator–prey interactions) are well documented (Bonsdorff 1992; Norkko and Bonsdorff 1996; Norkko et al. 2000; Råberg and Kautsky 2007). However, little is known about the effects these changes have had on seafloor primary productivity and the C cycle of the Baltic Sea and elsewhere.

The main objective of this study was to understand the role of perennial macroalgal canopies for primary productivity and organic C cycling dynamics in near-shore waters. We quantified rates of _F. vesiculosus_ canopy NEM from benthic oxygen flux measurements using the AEC technique at a single location in the Baltic Sea multiple times within a year. The AEC measurements were complemented with biological surveys, which were performed in parallel with the AEC measurements to monitor _F. vesiculosus_ canopy abundance and biomass. Based on regression analysis of the AEC fluxes and seabed sunlight availability (photosynthetically active radiation, [PAR]), we developed a simple predictive model for estimating daily canopy NEM continuously over 1 yr. This approach enabled us to evaluate canopy productivity on timescales ranging from 1 h to 1 yr, permitting for the first time detailed assessments of the productivity and potential rate of organic C export of a broadly distributed but understudied key coastal habitat.

**Materials and methods**

**Study location and sampling**

Seasonal sampling was performed on a macroalgal canopy situated close to the island archipelago of Spikama (59°48.40N, 23°12.26E), located 4.4 km SW from the Tvärminne Zoological Station in SW Finland (Fig. 1A). This location is typical of the extensive Finnish rocky coastline and Archipelago Sea region. A submerged rocky outcrop at a water depth of 2 m was colonized by a continuous ~ 50 m × 50 m macroalgal bed consisting primarily of the canopy-forming brown alga _F. vesiculosus_ (Fig. 1B, C). Benthic oxygen flux measurements and biological sampling of the _F. vesiculosus_ canopy were carried out at this site on five occasions between August 2016 and June 2017 (August, October, and December 2016 and March and June 2017).

**Benthic oxygen flux measurements**

Benthic oxygen fluxes were quantified in situ using the AEC technique (Berg et al. 2003). Our AEC systems are very
similar to the original design used by Berg and Huettel (2008), consisting of fast-response oxygen microsensors (Clark-type, Revsbech [1989]; $T_{90} \leq 0.3$ s, low stirring sensitivity < 1%, Gundersen et al. [1998]) interfaced with a 6 MHz acoustic velocimeter (Nortek) via submersible amplifiers (McGinnis et al. 2011). Two oxygen microsensors were used simultaneously for redundancy and intercomparison purposes and were attached to the stem of the velocimeter using a polyoxymethylene mounting that allowed positioning the sensors at an angle of ~ 60° relative to the velocimeter, with the ~ 20 μm microsensor tips located 0.5 cm away from the instrument’s 1.5 × 1.5 cm measurement volume (McGinnis et al. 2011). Instrumentation was mounted onto a sturdy aluminum tripod frame and was affixed to the frame so that the measurement volume was located ~ 35 cm above the seabed, well above the canopy height (Fig. 1B, C). A PAR sensor (LI-192, Li-Cor), a dissolved oxygen optode (Dissolved Oxygen Logger U26–001, HOBO), and a saltwater conductivity sensor (Salinity Data Logger U24–002-C, HOBO) located on the AEC frame logged transmitted (seabed) PAR, dissolved oxygen concentration, water temperature, and salinity at 5 min intervals throughout each deployment. AEC instrumentation was deployed by divers. The instrument was oriented within the predominant water flow direction (magnetic N-S), and care was taken to position the instrument level with the seafloor. Once deployed, the velocimeter measured the actual distance between the sensor measurement height and the seabed, before proceeding to log flow velocity and oxygen microsensor output in continuous sampling mode at 32 Hz. Individual deployments lasted 3–4 d.

**Biological sampling**

To relate the seasonal AEC measurements to the macroalgal canopy, abundance and biomass of *F. vesiculosus* were quantified seasonally, in conjunction with the AEC measurements at the end of each deployment. Biological sampling was performed by divers within an 80 m² circular area on the seafloor, with the AEC instrument located in its center. Eight 5 m-long weighted guidelines were used to delimit eight 45° direction sectors within the circle (Rodil et al. unpubl.). Abundance of *F. vesiculosus* (ind. m⁻²) within this area was quantified by randomly placing quadrats (50 cm × 50 cm polyvinylchloride frames) within the eight direction sectors, and then enumerating the individuals in situ. For the *F. vesiculosus* standing biomass estimation, eight intact *F. vesiculosus* individuals, one from each direction sector, were sampled carefully using large mesh bags (0.5 mm mesh size). In the laboratory, *F. vesiculosus* individuals were oven dried to constant weight at 60°C and the dry weight was assessed to the nearest 0.1 g. Macroalgal dry weight was converted to organic C assuming a conversion ratio for Baltic Sea *F. vesiculosus* of 1.00 g dw: 0.35 g C (Ilvessalo and Tuomi 1989).

**Oxygen flux extraction**

Benthic oxygen fluxes were extracted from the measured 32 Hz velocity and oxygen microsensor data streams following established protocols for bin-averaging the measured 32 Hz data to 8 Hz, O₂ sensor calibration, detrending, and quality-checking individual flux intervals for anomalous data (Berg et al. 2003; Lorrai et al. 2010; Berg et al. 2013). Oxygen fluxes for consecutive 15 min intervals were extracted from the 8 Hz data using the software package EddyFlux version 3.00 (P. Berg, unpubl.). The eddy flux is calculated as $w \overline{C}$, where $w$ is the fluctuating vertical velocity around its mean, and $\overline{C}$ is the fluctuating O₂ concentration around its mean, and the overbar symbolizes a time period average of 90 s that is sufficiently large to include the flux-contributing turbulent eddies.

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**Fig. 1.** (A) Location of our study site in Southwest Finland (red circle). (B) AEC instrumentation deployed at the site, and (C) a close-up of the *F. vesiculosus* canopy and the AEC sensors. Note that gas spaces (receptacles) are present on the apical tips. The height of the eddy covariance frame in (B) is 90 cm and the canopy height in (C) is ~ 20 cm. [Color figure can be viewed at wileyonlinelibrary.com]
(Berg et al. 2003). Following careful analysis of mean oxygen concentration dynamics, which were easily distinguishable from shorter term turbulence-driven variations, $w$ and $C$ were isolated from a 90 s running mean (Berg et al. 2003; McGinnis et al. 2008; Berg et al. 2013). Individual 15 min fluxes were then evaluated for their quality by carefully screening the 8 Hz oxygen concentration and cumulative instantaneous flux $wC$ for anomalous variations. Large jumps or spikes in oxygen concentration, typically caused by particles colliding with the sensors, were excluded (Berg et al. 2013). Once quality checked, we assessed the sensitivity of the eddy fluxes to coordinate transformation and data shifting. Coordinate corrections typically are required for measurements in complex flow conditions or due to instrument leveling errors, to exclude horizontal flux projections into the vertical eddy flux (Reimers et al. 2012; Lorke et al. 2013). Sensor separation distances and stirring sensitivity effects (McGinnis et al. 2008; Donis et al. 2012; Lorke et al. 2013). Sensor separation distances and stirring sensitivity effects (McGinnis et al. 2008; Donis et al. 2012; Lorke et al. 2013). Eddy flux footprint characteristics

For each sampling campaign, the size and shape of the sea-floor surface area included in the AEC flux measurements, the “flux footprint,” was investigated from measurements of sea-floor hydraulic roughness ($z_0$) and sensor measurement height, as described by Berg et al. (2007). The mean $z_0$ was computed from the flow velocity data streams using a multiple-step process. First, the complex Reynolds stress was calculated from streamwise ($u$), traverse ($v$), and vertical ($w$) velocity components, that were decomposed into a mean and a fluctuating velocity as $u = U + u'$, $v = v + v'$, and $w = W + w'$. From this, the friction velocity ($u_f$) was calculated as $u_f^2 = \overline{w w}^2 + \overline{v w}^2$ (Inoue et al. 2011). Subsequently, $z_0$ was calculated as $z_0 = z \times \exp\left(-k \times \frac{U}{\nu}\right)$, where $z$ is the AEC sensor measurement height above the seabed, $k$ is the Von Kármán constant (0.41), and $U$ is the mean flow velocity magnitude (Wüest and Lorke 2003). The length, width, and region of maximum contribution ($X_{\text{max}}$) within the AEC flux footprint were calculated for each sampling campaign using the equations provided by Berg et al. (2007). The estimated length and width of the flux footprint area were used to estimate surface area by assuming that the footprint was elliptical in shape.

Annual estimate of canopy NEM

A regression analysis was performed to develop a predictive model for estimating canopy daily NEM continuously over 1 yr. First, daily NEM for the five measurement campaigns was related to the corresponding seabed PAR (daily integrated; mol PAR m$^{-2}$ d$^{-1}$) using nonlinear regression and the asymptotic exponential function $NEM = a - b \times e^{-c \times PAR}$. The fitting function was then applied to year-round measurements of seabed PAR. Year-round PAR at the F. vesiculosus canopy was estimated from measurements of incoming PAR (SMEAR III Station, University of Helsinki) that were corrected for attenuation, by applying attenuation factors that were determined in situ for different seasons during each measurement campaign. Daily canopy NEM was integrated over the year to compute annual canopy NEM in mol O$_2$ m$^{-2}$ yr$^{-1}$ and in kg C m$^{-2}$ yr$^{-1}$ (assuming respiratory and photosynthetic O$_2$ : CO$_2$ of 1.0). This approach is similar to the one used in the studies by Rheuban et al. (2014a) and Attard et al. (2015) to quantify annual NEM in seagrass beds and coralline algal beds.

Oxygen storage within F. vesiculosus gas spaces

We explored whether gas storage within intercellular air spaces of F. vesiculosus could be significant. Earlier studies have documented considerable oxygen accumulation within macroalgal gas spaces under illumination (Damant 1936; Vallance and Coult 1951; Tammes 1954; Aleem 1969). To investigate this potential mechanism, oxygen microsensors (fiber-optode, 300 μm tip diameter, $T_{90} < 10$ s) connected to an optical oxygen meter (FireStingO2 4-channel, PyroScience) were inserted into gas spaces of intact F. vesiculosus plants that were collected from the study area in June, and percentage air saturation was logged within the bladders at 5 s intervals while exposing the submerged plant to light–dark cycles in temperature-controlled (10 °C) laboratory flow-through aquaria, under ~100% air saturated water. A light-emitting diode panel that accurately simulates natural outdoor light was used for illumination (~400 μmol PAR m$^{-2}$ s$^{-1}$), and a dissolved oxygen optode (Dissolved Oxygen Logger U26-001, HOBO) logged dissolved oxygen concentration in the aquarium water at 5 min intervals.

Results

Environmental conditions

The environmental conditions for the five measurement campaigns are summarized in Table 1. Water temperature ranged from 1.3 ± 0.3°C in March to 15.9 ± 0.6°C in August (mean ± SD). Salinity was between 5.6 ± 0.0 and 6.4 ± 0.1 throughout the year (mean ± SD). Daily dissolved oxygen concentration was highest in March (435 ± 21 μmol L$^{-1}$) and lowest in August (277 ± 26 μmol L$^{-1}$) (mean ± SD). Oxygen saturation ranged from 90% ± 2% in December to 103% ± 5% in March (mean ± SD). Diel variations in dissolved oxygen concentration were evident in all seasons. The highest diel amplitudes were observed in June (120 μmol L$^{-1}$) and August (111 μmol L$^{-1}$), with intermediate values observed in March (63 μmol L$^{-1}$) and October (56 μmol L$^{-1}$), and the lowest values in December (16 μmol L$^{-1}$). Daily integrated
Seasonal canopy dynamics

Our seasonal measurements of *F. vesiculosus* abundance indicate that the number of canopy-forming individuals remained more or less constant throughout the year, ranging from 15 ± 4 ind. m−2 in October to 19.5 ± 3 ind. m−2 in August (mean ± SD, n = 8), with no significant differences observed between the five measurement campaigns (one-way ANOVA $F_{5,42} = 1.22; p = 0.32$; Fig. 2). However, the canopy standing biomass and the estimated C standing stock decreased significantly from summer to late winter (from 0.4 ± 0.2 kg C m−2 in August to 0.2 ± 0.1 kg C m−2 in March; one-way ANOVA $F_{5,42} = 3.47; p = 0.01$), and then ammased biomass to values comparable to the previous summer between March and June (0.4 ± 0.2 kg C m−2; Fig. 2). Length of *F. vesiculosus* individuals (mean ± SD, n = 8) ranged from 33 ± 6 cm in December to 40 ± 6 cm in August, and the canopy height was typically 15–20 cm.

Eddy fluxes: Hourly dynamics

Each measurement campaign yielded 1–3 d of continuous, high-quality eddy flux data. The hourly fluxes document considerable temporal dynamics in response to changing environmental conditions such as PAR over diel periods (Fig. 3). High flow velocities of up to 10.8 cm s−1 at this moderately exposed site combined with a rough seabed surface maintained well-developed turbulent conditions throughout the selected datasets (Fig. 3). Linear cumulative instantaneous fluxes evidenced a favorable turbulent flux signal for extracting eddy fluxes (Fig. 3C). The mean ± SD hydraulic roughness ($z_{0}$) ranged from 2.1 ± 1.3 cm (n = 322) in June to 3.6 ± 1.3 cm (n = 193) in December, corresponding to a flux footprint length (area) range between 26 m (35 m²) and 42 m (56 m²) (Berg et al. 2007). The global average footprint length for the five measurement campaigns was 30 ± 7 m (n = 5) and the average area was 40 ± 9 m² (n = 5). The region of maximum flux contribution ($X_{\text{max}}$) within the footprint area was located between 0.6 and 1.5 m upstream from the instrument (global average = 0.8 ± 0.4 m; n = 5).

The hourly eddy fluxes were broadly correlated with the PAR availability in all seasons, where sunlight reaching the canopy stimulated a measurable photosynthetic oxygen production (Fig. 4A, B). The highest hourly net oxygen release rates by the *F. vesiculosus* canopy were observed in August. Here, canopy PAR of up to ~ 900 μmol quanta m−2 s−1 corresponded to net oxygen release rates by the canopy of up to 35.2 ± 9.1 mmol m−2 h−1. As measurements progressed into the autumn, the amount of available sunlight at the canopy surface decreased to a maximum of ~ 150 μmol PAR m−2 s−1, with a corresponding decrease in the maximum observed net oxygen release rates (4.6 ± 0.3 mmol m−2 h−1). Canopy PAR and oxygen release rates were lowest in early winter (December). During this measurement campaign, a maximum of ~ 90 μmol PAR m−2 s−1 was measured at the canopy surface,
and daytime oxygen fluxes were less negative than nighttime values, but were low in magnitude compared to daytime fluxes measured during the other campaigns (up to $0.2 \pm 0.2 \text{ mmol m}^{-2} \text{ h}^{-1}$). As sunlight intensity and the duration of the photic period increased toward the end of winter, remarkably high hourly rates of net oxygen production were measured, despite the cold water temperatures. In March, oxygen fluxes reaching $16.7 \pm 5.2 \text{ mmol m}^{-2} \text{ h}^{-1}$ were measured under maximum PAR of $\sim 250 \mu\text{mol m}^{-2} \text{ s}^{-1}$ and water temperatures of just $1.3 \pm 0.3^\circ\text{C}$. High canopy net oxygen release rates of up to $23.4 \pm 7.7 \text{ mmol m}^{-2} \text{ h}^{-1}$ were maintained through to early summer (June), under the highest measured canopy PAR values for our seasonal measurement campaign of up to $\sim 1350 \mu\text{mol PAR m}^{-2} \text{ s}^{-1}$ (Fig. 4).

The eddy flux datasets for June, August, and October showed interesting hourly dynamics where high oxygen release rates were maintained through to the end of the day under very low light levels (e.g., $3.6 \pm 1.1 \text{ mmol m}^{-2} \text{ h}^{-1}$ under $\sim 4 \mu\text{mol PAR m}^{-2} \text{ s}^{-1}$ in June; Fig. 4). Furthermore, dark rates of oxygen uptake during these months overall were very low. During some nighttime periods with well-developed turbulent conditions in June and in August, nighttime fluxes were around $0 \text{ mmol m}^{-2} \text{ h}^{-1}$ or were even positive, suggesting minimal oxygen uptake or even an apparent oxygen uptake.
release by the canopy under dark conditions (Figs. 3–4). This repeated pattern could not be explained by accounting for uncertainties in eddy flux extraction, namely from (a) changes in water flow velocity and oxygen concentration (Holtappels et al. 2013; Rheuban et al. 2014b), (b) accumulation/depletion dynamics of oxygen within the canopy (Hendriks et al. 2014; Long et al. 2015b), (c) changes in water flow direction, (d) flux sensitivity to instrument coordinate rotation (Reimers et al. 2012; Lorke et al. 2013), and (e) response time of the AEC sensors to changes occurring at the seafloor surface (integration time; Rheuban and Berg 2013). The original fluxes therefore were of high quality and were representative of the true benthic flux. Subsequently, we considered whether gas storage within gas spaces of F. vesiculosus could be significant. Intact specimens of F. vesiculosus that were collected from the field in the morning had high oxygen saturation levels within their gas spaces (~33% oxygen), and this value increased to ~40% following 18 h of illumination (Fig. 5). Oxygen content decreased exponentially with time in the dark, but remained above air saturation levels of the ambient water for ~12 h, much longer than the typical nighttime duration for this time of the year (June, ~5 h). Despite there being lower light levels in the experimental setup relative to typical summer time field conditions (400 µmol PAR m⁻² s⁻¹ vs. ~1200 µmol PAR m⁻² s⁻¹), these results indicate that there is considerable oxygen accumulation and drawdown from within F. vesiculosus gas spaces during day and night periods, respectively. Furthermore, there exists a disequilibrium between internal (gas space) and external (water phase) oxygen concentration, and interestingly, this concentration gradient indicates a potential for oxygen release by the canopy under dark conditions, which is consistent with the dynamics we observed in the eddy covariance fluxes. Quantifying photosynthetic production and respiration rates of gas-storing macroalgae from in situ hourly fluxes or from external changes in oxygen concentration (using, e.g., eddy covariance, chamber incubations, or from microprofiling measurements within boundary layers above fronds) would thus underestimate the true gross metabolic activity. In our study, however, we focus on the daily NEM, which is calculated from continuous flux time series lasting one or more days. This approach would cancel out the gas storage effect and would provide robust daily rate estimates of NEM.

Eddy fluxes: Daily and seasonal canopy productivity

Oxygen production by the F. vesiculosus canopy exceeded oxygen consumption over 24 h in four out of the five measurement campaigns (Fig. 4C). Daily NEM, quantified as the integration of continuous hourly eddy fluxes over 24 h, indicated that the canopy was strongly net autotrophic in summer (June = 166 mmol O₂ m⁻² d⁻¹ and August = 98 mmol O₂ m⁻² d⁻¹). Net autotrophy persisted in autumn (October = 11 mmol O₂ m⁻² d⁻¹) and in late winter (March = 48 mmol O₂ m⁻² d⁻¹), but the canopy was net heterotrophic in early winter (December = −17 mmol O₂ m⁻² d⁻¹; Fig. 4C).

Following a regression analysis of daily canopy NEM and light (PAR) availability, we found that the daily NEM for the five measurement campaigns was highly correlated with the
corresponding daily integrated PAR reaching the *F. vesiculosus* canopy surface. The asymptotic exponential function \( NEM = a - b \times c^{\text{PAR}} \) gave a remarkably good fit \( (R^2 = 0.99) \) to the measured data, providing us with a simple empirical relationship for estimating canopy \( NEM \) (Fig. 6). PAR attenuation factors for the different seasons were derived by relating the measurements of incoming PAR vs. canopy PAR using linear regression. The \( R^2 \) value was 0.84 in March, 0.92 in December, 0.94 in August, 0.95 in June, and 0.96 in October. Applying the established fitting function to year-round canopy PAR (Fig. 7A) yielded continuous daily \( NEM \) throughout the 12 months of our measurement period (Fig. 7C). The predictive model was based on canopy PAR that ranged from 0.1 mol photons m\(^{-2}\) d\(^{-1}\) (model \( NEM = -28 \text{ mmol O}_2 \text{ m}^{-2} \text{ d}^{-1}\) in December to 39.6 photons mol m\(^{-2}\) d\(^{-1}\) (model \( NEM = 183 \text{ mmol O}_2 \text{ m}^{-2} \text{ d}^{-1}\) in June (Fig. 7A, C). This variability range was similar to the range observed in the field during our five measurement campaigns. Here, daily integrated canopy PAR ranged from 1.0 mol m\(^{-2}\) d\(^{-1}\) in December (measured \( NEM = -17 \text{ mmol O}_2 \text{ m}^{-2} \text{ d}^{-1}\) to 30.7 mol m\(^{-2}\) d\(^{-1}\) in June (measured \( NEM = 166 \text{ mmol O}_2 \text{ m}^{-2} \text{ d}^{-1}\); Table 1; Fig. 4), indicating that our measurements covered a large part of the seasonal variability in daily PAR that the *F. vesiculosus* canopy was exposed to in situ. The results from this simple dynamic modeling exercise based on the empirical relationship that we established between daily PAR and \( NEM \) indicate that the canopy was net autotrophic \( (NEM > 0 \text{ mmol O}_2 \text{ m}^{-2} \text{ d}^{-1}) \) for 246 \( \text{ d yr}^{-1} \), with high net autotrophy \( (NEM > 100 \text{ mmol O}_2 \text{ m}^{-2} \text{ d}^{-1}) \) observed during 150 \( \text{ d yr}^{-1} \). The canopy was net heterotrophic \( (NEM < 0 \text{ mmol O}_2 \text{ m}^{-2} \text{ d}^{-1}) \) from mid-November to mid-February, equivalent to 33% (or 119 d) of the year (Fig. 7C). Integrating daily \( NEM \) over the year indicated that the *F. vesiculosus* canopy was highly net autotrophic on an annual basis, with annual canopy \( NEM \) amounting to 25.2 mol O\(_2\) m\(^{-2}\) yr\(^{-1}\) or 0.3 kg C m\(^{-2}\) yr\(^{-1}\) (Fig. 7D).

**Discussion**

**Hourly, daily, and seasonal canopy productivity**

Our in situ AEC measurements integrating ~ 40 m\(^2\) of seabed surface area document considerable oxygen production by the *F. vesiculosus* canopy year-round, with hourly net production rates of up to \( 35 \pm 9 \text{ mmol m}^{-2} \text{ h}^{-1} \) in late summer under maximum water temperature of \( \sim 16^\circ\text{C} \) and canopy PAR of up to \( \sim 900 \mu\text{mol quanta m}^{-2} \text{ s}^{-1} \). Even though it is likely that the hourly rates underestimate the true net production due to gas storage, the rate magnitudes are still high, and are comparable to rates observed in shallower, dense seagrass canopies at lower latitude exposed to higher PAR and warmer water temperature (Rheuban et al. 2014b). Macroalgal canopies in other geographical locations similarly express remarkably high hourly rates of productivity in excess of 100 mmol C m\(^{-2}\) h\(^{-1}\) during peak irradiance in summer (Tait and Schiel 2010; Bordeyne et al. 2017). Interestingly, high hourly rates of net
oxygen production by the *F. vesiculosus* canopy > 15 mmol O₂ m⁻² h⁻¹ were not limited to the summer months but were also observed in late winter (March) under maximum PAR of ~ 250 μmol m⁻² s⁻¹ and water temperatures of just 1.3 ± 0.3°C. It is interesting to note that our seasonal measurements indicate key differences in the biogeochemical functioning of the canopy between early and late winter. In December, under water temperatures of 3.4 ± 0.5°C and daily PAR of 1.0 mol m⁻² d⁻¹, a distinct light-response was evident in the eddy fluxes (Fig. 4), but respiration exceeded primary production over 24 h, and the canopy was net heterotrophic *(NEM = −17 mmol m⁻² d⁻¹)*. By contrast, in March, under colder water temperatures (1.3 ± 0.3°C) but higher light availability (5.4 mol m⁻² d⁻¹) and a longer photic period, the canopy was net autotrophic *(NEM = 48 mmol m⁻² d⁻¹)*. These observations challenge traditional views on high-latitude seafloor ecosystems as being generally unproductive in winter with sluggish rates of activity. While early winter measurements indicate a net heterotrophic habitat relying on external or stored organic material to sustain ~ 75% of the daily C demand, our subsequent winter measurements performed in March indicate that macroalgal canopies are able to efficiently harvest the available light and are able to assimilate large amounts of C, more than is required to sustain the daily respiratory requirements of the canopy habitat. Indeed, positive daily NEM was observed in four of the five measurement campaigns. Comparison of these results to NEM estimates for other habitats worldwide (seagrass, macroalgae, coralline algal beds, bare sediments; considering only studies that computed the daily NEM from continuous diel measurements over 24 h) suggests that macroalgal canopies overall have amongst the highest (i.e., most positive) daily NEM (Fig. 8). While some studies have investigated productivity rates of *F. vesiculosus* in the Baltic Sea (Kairesalo and Leskinen 1986; Leskinen et al. 1992), studies on whole-canopy productivity remain few. Our results indicate that macroalgal canopies effect amongst the highest hourly rates of seafloor primary productivity, and are amongst the most highly net autotrophic habitats. The seasonal and annual NEM values we obtained in our study are high when compared to existing values for the pelagic environment nearby our study site. Annual phytoplankton gross productivity rates (depth-integrated over the ~ 10 m photic zone) quantified using ¹⁴C incubations range from 74–170 g C m⁻² yr⁻¹, with about two-thirds of phytoplankton productivity being respired within the water column (Kuparinen et al. 1984; Lignell et al. 1993). These results document the importance of seafloor primary production on an ecosystem level. The spatial extent and distribution of macroalgal canopies on the seafloor is likely to be an important factor in determining organic C assimilation, CO₂ drawdown and O₂ production rates in coastal waters.

Our laboratory measurements documented substantial oxygen accumulation and light-dependent dynamics within gas spaces of *F. vesiculosus*. Two types of intercellular gas spaces were identified on the thallus branches of the *F. vesiculosus* canopy we investigated: vesicles were located along the fronds at regular intervals on either side of the midrib, and receptacles were present at the apical tips. Earlier studies have documented considerable oxygen accumulation in macroalgae gas spaces under illumination (Damant 1936; Vallance and Coult 1951; Tammes 1954; Aleem 1969). Even though the gas space volume typically is small relative to that of the surrounding water, the accommodation coefficient of oxygen is ~ 40 times higher in air than in water. The degree of storage *(E, in %)* can be assessed on the basis of gas-to-water ratio and solubility of oxygen according to the equation: \[ E = p/\alpha (V_g \times 1000/V_o) \], where *p* is the fractional volume of oxygen in air (0.21), \( \alpha \) is oxygen solubility in water, \( V_g \) is the gas volume, and \( V_o \) is the volume of incubated water (Sorrell and Dromgoole 1986). This equation implies that under conditions of instantaneous oxygen equilibrium between gas space and water, a 1% gas volume contains ~ 25% of all oxygen, and thus considerable oxygen storage potential. We propose that high oxygen
release rates by the seabed at the end of the day, as well as low oxygen consumption at night, as we observed in our AEC datasets during periods with high metabolic activity, are consistent with the gas storage effect. Oxygen accumulation in gas spaces during the day would buffer oxygen consumption requirements at night, thereby reducing the drawdown requirements of oxygen from the water column. These observations contrast with seafloor oxygen flux dynamics observed in other habitats such as seagrass beds and bare sediments, which typically show higher seafloor oxygen uptake rates in the evening than in the morning due to photosynthesis-coupled respiration (Fenchel and Glud 2000; Rheuban et al. 2014b). Further measurements are required to constrain the effects of oxygen storage on the oxygen budget within macroalgal canopies, in order to better understand its ecological and biogeochemical significance.

**Annual canopy NEM**

The results from the predictive model indicate that daily NEM was positive for 246 d (or two-thirds) of the year (Fig. 7B). When integrated over the year, the *F. vesiculosus* canopy was highly net autotrophic, with annual NEM amounting to ~25 mol O₂ m⁻² yr⁻¹ (~0.3 kg C m⁻² yr⁻¹; Fig. 7C). The process of deriving annual NEM based on the empirical relationship between daily integrated PAR and NEM for the five measurement campaigns (Figs. 6–7) is a first-order approach that invokes several assumptions about the functioning of the canopy. In particular, this approach assumes that the daily balance between primary productivity and respiration over the course of the year is driven only by changes in the amount of daily irradiance reaching the seafloor, thus not directly accounting for other potentially important factors such as seasonal changes in standing biomass and community composition, sedimentation dynamics, light acclimation, and temperature effects. We estimated annual NEM using a second approach that is independent of the PAR-NEM relationship. In this second approach, the daily NEM values from the individual measurement campaigns were plotted over the year, and a Gaussian function was fitted to this data using a least-squares approach to interpolate between data points (OriginPro 8.5, OriginLab). The Gaussian function thus described daily NEM dynamics over 12 months, and the integrated value of this curve then represents the annual NEM. The fitting function gave a tight fit to the measured data ($R^2 = 0.99$), and the estimated annual NEM of 23.5 mol O₂ m⁻² yr⁻¹ was within 6% of our previous estimate (25 mol O₂ m⁻² yr⁻¹). We can further constrain the minimum annual NEM and C export using the seasonal *F. vesiculosus* standing biomass measurements (Fig. 2). The decrease between summer and late winter (17 mol O₂ m⁻² yr⁻¹ [0.2 ± 0.2 kg C m⁻²]), and the subsequent increase between winter and summer (17 mol O₂ m⁻² yr⁻¹ [0.2 ± 0.2 kg C m⁻²]), represent the minimum annual rates of C export and NEM.

Very few studies have considered seasonal changes in canopy productivity on a community-scale when computing the annual estimate. To our knowledge, there exist none for macroalgal canopies. Comparison to available datasets for other habitats indicates that the annual NEM for the *F. vesiculosus* canopy is high. Martin et al. (2007) and Attard et al. (2015) found that coralline algal beds were net heterotrophic on an annual basis (~14 and ~4 mol m⁻² yr⁻¹, respectively), as were the bare sediments in the studies by Attard et al. (2015) (~7 mol m⁻² yr⁻¹) and Rheuban et al. (2014a) (~8 mol m⁻² yr⁻¹). Annual NEM of seagrass beds in the Virginia Coastal Reserve, ranged from net heterotrophic (~7 mol m⁻² yr⁻¹) to net autotrophic (9 mol m⁻² yr⁻¹) (Rheuban et al. 2014a). The *F. vesiculosus* canopy investigated in this study thus has higher annual NEM than some of the most productive seagrass beds (Fig. 8B), an observation that is in line...
with the global synthesis on annual NEM of submerged vegetated habitats (Duarte 2017). The annual canopy NEM of 25 mol O$_2$ m$^{-2}$ yr$^{-1}$, or 0.3 kg C m$^{-2}$ yr$^{-1}$, is comparable to the global average estimate for macroalgal bed NEM of 0.4 kg C m$^{-2}$ yr$^{-1}$ (Krause-Jensen and Duarte 2016).

It is likely that the high daily and annual NEM rates associated with macroalgal canopies are a result of a combination of factors relating to high macroalgal primary productivity, low sediment accumulation rates within-habitat, and low herbivory rates. Macroalgal canopies typically colonize submerged rocky outcrops, having low sediment and detritus accumulation. Canopy-forming macroalgae such as $F$. vesiculosus have a high proportion of their biomass that is photosynthetically active when compared to other canopy-forming vegetation (e.g., seagrass), and thus are able to photosynthesize efficiently under low light and can assign a large proportion of their new production to growth (Markager and Sand-Jensen 1992). Production of anti-grazing compounds (Rohde et al. 2004) and high frond C : N (e.g., C : N of $F$. vesiculosus in the Baltic Sea ranges seasonally from ~ 15 to ~ 60, with an annual average of ~ 37) (Ilvessalo and Tuomi 1989) may restrict extensive canopy grazing. Furthermore, many marine invertebrates that are commonly associated with macroalgal canopies in the Baltic Sea, such as the isopod Idotea balthica and the bivalve Mytilus edulis, apparently lack the carbohydrases that are required to break down the plant tissue, and thus are not able to digest the plant material (Hylleberg Kristensen 1972). Altogether, these processes would favor low rates of community respiration within macroalgal canopies, with potential important implications for biogeochemical cycles and habitat functioning.

**C export potential**

Globally, it is estimated that a large proportion of macroalgal NEM (> 40%) is exported to surrounding environments (Duarte and Cebrian 1996; Krumhansl and Scheibling 2012; Krause-Jensen and Duarte 2016). Our measurements of $F$. vesiculosus abundance indicate that the number of canopy-forming individuals remained more or less constant throughout the year, with no significant differences observed between the five measurement campaigns (Fig. 2). However, the canopy standing biomass and the estimated C standing stock decreased significantly from summer to late winter (from 0.4 ± 0.2 kg C m$^{-2}$ in August to 0.2 ± 0.1 kg C m$^{-2}$ in March), and then amassed biomass to values comparable to the previous summer between March and June (0.4 ± 0.2 kg C m$^{-2}$). These measurements, in combination with our personal observations whilst surveying the canopy, indicate that a sizable canopy is present year-round. In addition to being a perennial habitat-forming structure, the canopy is a substantial source of organic matter to its surroundings. In our study, annual integrated values of canopy NEM indicate a surplus of organic C production amounting to 25 mol O$_2$ m$^{-2}$ yr$^{-1}$ (or 0.3 kg C m$^{-2}$ yr$^{-1}$; Fig. 7D). When considering that there was no detectable accumulation of canopy biomass on an annual basis, nor a substantial accumulation of detritus on the rocky substratum within the macroalgal bed, we can assume that most of the surplus C produced by the canopy was exported as particulate and dissolved matter. Thus, our measurements indicate that annual net C export rates are comparable to the canopy standing biomass observed in summer. Due to the rocky nature of the underlying substrate, C burial within the habitat itself will be very low, so exchange of detritus will occur beyond the perimeter of the habitat itself (Krumhansl and Scheibling 2012). On multiple occasions in summer and in autumn, we observed vast underwater fields of unattached macroalgal detritus on the seabed of an enclosed embayment nearby the study site that consisted primarily of $F$. vesiculosus (Fig. 9). Factors such as seasonal decay, epiphyte growth, and grazing accelerate blade erosion and fragmentation (Knight and Parke 1950; Russell 1985; Krumhansl and Scheibling 2012; Anderson and Martone 2014). Additionally, physical forcing such as intensified wave energy and currents during storms, as well as seasonal ice scouring, would dislodge whole fronds and thalli, and thus would facilitate C export (Kiirikki and Ruskanen 1996; Krumhansl and Scheibling 2012). Analysis of wind speed and direction data collected from a weather station nearby our sampling site (Hanko Rus sarö, Finnish Meteorological Institute) indicates a prevalence of southerly winds (wind directions > 90° and < 270°) from September to December, with high southerly winds (wind speeds > 10 m s$^{-1}$) being a common occurrence in October and November (data not shown). This period coincides with a substantial reduction in biomass in the $F$. vesiculosus canopy (Fig. 2), potentially indicating that these months are particularly important for particulate C export. Accumulations of drifting detritus constitute nourishment and a temporary habitat structure for opportunistic mobile taxa such as Ostracod crustaceans and the water snail $Hydrobia$ spp. (Norkko et al. 2000). Conversely, given high enough abundance,
drifting macroalgae can represent hotspots of anoxia and hydrogen sulfide production, generating adverse conditions to coastal biota and inducing large-scale mortality of sedentary macrobenthic populations (Norkko and Bonsdorff 1996; Glud et al. 2004; Norkko et al. 2013). The exported detritus is therefore an important spatial subsidy and a form of habitat connectivity within the coastal zone that can have positive as well as deleterious effects on aquatic ecosystems (Polis et al. 1997).

A proportion of the exported C by the canopy will be in dissolved form. Canopy-forming algae such as F. vesiculosus as well as other canopy-associated algae exude a proportion of their daily C fixation as dissolved organic carbon (DOC), which exchange with the surrounding water and could make up a significant proportion of the total DOC pool in coastal waters (~20%) (Wada and Hama 2013). Exudation rates for F. vesiculosus range from <5% to 40% of net C production and vary seasonally (Sieburth 1969; Carlson and Carlson 1984), with global average values for macroalgae DOC exudation at ~20% of NEM (Maher and Eyre 2010; Krause-Jensen and Duarte 2016). DOC exudation by benthic microalgae and ephemeral macroalgae, that would be present within the canopy as epiphytes on the fronds or as a biofilm on the underlying rocks, would also contribute to DOC export (Underwood and Paterson 2003). Based on these literature values, it is possible that the F. vesiculosus canopy investigated in this study exuded DOC at a rate of 37–73 mmol m⁻² d⁻¹ (0.5–0.9 g C m⁻² d⁻¹) during peak production periods in summer, which is at the upper range of values reported for seagrass beds and macroalgae worldwide (Maher and Eyre 2010; Barrón et al. 2014). The derived annual values of DOC export from the F. vesiculosus canopy of 5–10 mol m⁻² yr⁻¹ (60–121 g C m⁻² yr⁻¹) are in good agreement with the global average value for DOC export from algal beds of 8 mol m⁻² yr⁻¹ (101 g C m⁻² yr⁻¹) (Krause-Jensen and Duarte 2016).

**Outlook and perspectives**

Decades after the influential studies by Mann (1973) and Smith (1981), the important functions of vegetated seafloor habitats in the oceanic C cycle are increasingly being recognized (Duarte 2017). Despite this advancement, the dynamics of macroalgal canopy productivity remain poorly understood. Recent technological developments, such as the AEC technique used in this study, as well as other non-invasive techniques that integrate over large seafloor surface areas (tens to hundreds of square meter) (Falter et al. 2008; McGillis et al. 2011), constitute promising new tools to investigate habitat-scale productivity rates and their drivers in macroalgal canopies under in situ conditions. The high temporal flux resolution that AEC provides enabled us to investigate hourly dynamics in detail. In this study, we have documented the importance of perennial macroalgal canopies in the Baltic Sea for primary productivity and biogeochemical functioning, on timescales ranging from 1 h to 1 yr. The F. vesiculosus canopy effected high hourly rates of oxygen production that are comparable to the most productive seagrass beds worldwide. Through detailed analyses of the hourly flux dynamics and their drivers, we were able to identify a mechanism that explains an apparent release of oxygen by the seafloor under dark conditions. The fact that high hourly rates of oxygen production were also measured in late winter under low light availability and cold water temperatures is a substantial new finding that highlights the importance of conducting seasonal measurements. The canopy was highly net autotrophic on an annual basis, and we are able to conclude, through comparing annual NEM with seasonal measurements of canopy standing biomass, that much of this surplus C production is exported to its surroundings.

Macroalgal canopies represent significantly understudied habitats when compared to other conspicuous canopy-forming vegetation such as seagrasses. More research is required to investigate NEM and its drivers within macroalgal canopies. Additional measurements in mid-winter (January), as well as during other periods of interest (e.g., storms and production blooms), would provide considerable new insight. Incorporating macrovegetation into estimates of coastal primary productivity is an important next step. Furthermore, determining the fate of the exported C from macroalgal canopies, and its importance as a resource subsidy for secondary production in adjacent seafloor communities, requires further research.

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Conflict of Interest

None declared.

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