Photosynthetic capacity in seagrass seeds and early-stage seedlings of Zostera marina

Brodersen, Kasper Elgetti; Kühl, Michael

Published in:
New Phytologist

DOI:
10.1111/nph.18986

Publication date:
2023

Document version
Publisher's PDF, also known as Version of record

Document license:
CC BY

Citation for published version (APA):
Photosynthetic capacity in seagrass seeds and early-stage seedlings of *Zostera marina*

Kasper Elgetti Brodersen1,2 and Michael Kiil1

1Marine Biological Section, Department of Biology, University of Copenhagen, 3000 Helsingør, Denmark; 2Present address: Environmental Dynamics, Department of Science and Environment, Roskilde University, 4000 Roskilde, Denmark

Summary

- In many terrestrial seeds, photosynthetic activity supplies O\(_2\) to the developing plant embryo to sustain aerobic metabolism and enhance biosynthetic activity. However, whether seagrass seeds possess similar photosynthetic capacity to alleviate intra-seed hypoxic stress conditions is unknown.
- We used a novel combination of microscale variable chlorophyll fluorescence imaging, a custom-made O\(_2\) optode microrespirometry system and planar optode O\(_2\) imaging, to determine the O\(_2\) microenvironment and photosynthetic activity in developing seeds and seedlings of seagrass (*Zostera marina*).
- Developing, sheath-covered seeds exhibited high O\(_2\) concentrations in the photosynthetic active seed sheath and low O\(_2\) concentrations in the centre of the seed at the position of the embryo. In light, photosynthesis in the seed sheath increased O\(_2\) availability in central parts of the seed enabling enhanced respiratory energy generation for biosynthetic activity. Early-stage seedlings also displayed photosynthetic capacity in hypocotyl and cotyledonary tissues, which may be beneficial for seedling establishment.
- Sheath O\(_2\) production is important for alleviating intra-seed hypoxic stress, which might increase endosperm storage activity, improving the conditions for successful seed maturation and germination.

Introduction

Seagrass plants are marine angiosperms that form important coastal ecosystems playing a key role for coastal protection, biodiversity of marine organisms and sequestration of nutrients and carbon (Madsen et al., 2001; Fourqurean et al., 2012; Bertelli & Unsworth, 2014). Yet, seagrass meadows are declining worldwide due to climate change and other anthropogenic threats (Orth et al., 2006; Waycott et al., 2009), which has stimulated increasing research on seagrass habitat restoration (Katwijk et al., 2016). Seed-based restoration is of major interest due to the potential for restoring larger areas and at the same time ensuring high genetic diversity in re-established seagrass meadows (Marion & Orth, 2012; Katwijk et al., 2016; Cumming et al., 2017; Unsworth et al., 2019). However, there are still large knowledge gaps in the understanding of fundamental mechanism and processes influencing seed development, germination and survival (York et al., 2016). Seagrass seed production is initiated in flowering shoots in the warming waters during spring (Silberhorn et al., 1983). During seed development, the seeds are either: released from the plant sedimenting directly onto the sea floor; or dispersed via detached decaying reproductive shoots or in seed-containing spathes on the sediment surface (Harwell & Orth, 2002; Marion & Orth, 2012; Hosokawa et al., 2015). Most seeds settle within or close to the meadow of origin (Harwell & Orth, 2002; Hosokawa et al., 2015). For *Zostera marina* plants growing in temperate regions, seed germination occurs over the autumn/winter, whereafter the young seedlings commence rapid growth during the following springtime with the increased light availability (Marion & Orth, 2012).

Photosynthetic activity during seagrass seed germination and seedling development has been demonstrated for nondormant seagrass species like *Posidonia oceanica* and *Thalassia testudinum* (Celdrán & Marín, 2011, 2013; Celdran, 2017). Seed photosynthesis enhances *Posidonia oceanica* seedling (leaf and root) growth (Celdran & Marín, 2013), where the autotrophic production complements stored carbohydrate reserves until the seedling’s photosynthetic capacity is capable of supporting its own carbon demands (Celdran & Marín, 2013). The *Posidonia* seed photosynthesis also seems to support the seedlings O\(_2\) demand, a mechanism that could be important for avoiding hypoxia during seed embryogenesis (Celdran et al., 2015). Low internal O\(_2\) conditions can limit respiration rates and seed photosynthesis could potentially alleviate this hypoxic stress through an O\(_2\) supported supply of respiratory energy (Celdran et al., 2015). *Posidonia* seed photosynthesis could thus help alleviate the O\(_2\) demand required to metabolize the carbohydrate reserves stored in the seeds (Celdran et al., 2015). However, a similar mechanism remains to be explored.
be demonstrated in dormant seagrass seeds such as found in the key model seagrass species *Z. marina*. These seeds differ substantially from *Posidonia* seeds both in terms of their architecture and life cycle (i.e. seed sheath, hard seed coat and dormancy of up to 12 months; Orth *et al.*, 2000) and may, therefore, have higher similarity to seed development of terrestrial monocot plants.

Seeds of *Z. marina* are known to have distinct dormancy and a hard seed coat (Orth *et al.*, 2000), but its functional roles in terms of seed protection, photosynthetic activity and overall architecture remain largely unknown. In terrestrial seeds (e.g. seeds of soybean and grains of barley and maize; Rolletschek *et al.*, 2004, 2005a,b; Borisjuk *et al.*, 2005), the function of the seed coat has mainly been related to protective properties. However, the seed coat also functions as a connection to the external microenvironment by transferring environmental cues into the seed, which leads to adjustments of the metabolic activity in terrestrial seeds (Radchuk & Borisjuk, 2014). The epidermal cuticle of the seed coat in terrestrial plant seeds forms a physical barrier but does not impede the exchange of gases such as O₂ and CO₂ (Radchuk & Borisjuk, 2014).

Developing seeds of terrestrial plants have high respiratory needs, generating hypoxic to near anoxic internal conditions, which thus requires additional O₂ supplies to sustain aerobic respiration (e.g. Rolletschek *et al.*, 2004, 2005a,b). Low internal O₂ conditions may be beneficial to the seed during development as it can stimulate high carbon use efficiencies through increased bioenergetic efficiency of mitochondria (Radchuk & Borisjuk, 2014 and references therein), and hypoxic conditions could also play a major role in enabling the longevity of seeds and provide means for regulating local metabolic activity (Borisjuk & Rolletschek, 2009). However, nutrient transport and endosperm storage activity largely depend on sufficient energy supply from respiration, which is dependent on sufficient O₂ availability that can be enhanced by photosynthesis in the surrounding chlorenchymatic layer (Radchuk & Borisjuk, 2014). Photosynthesis in the seed coat has been demonstrated in many terrestrial seeds, driving seed carbon fixation and O₂ production (Rolletschek *et al.*, 2004), often at much lower light conditions than in leaves (Radchuk & Borisjuk, 2014). Seed coat photosynthesis can thus relieve hypoxic stress and enhance the biosynthetic activity of the developing seed (Rolletschek *et al.*, 2003, 2005a,b). The main input of nutrients to the developing seed is, however, supplied by the mother plant leaf, where the seeds’ ability to sense light enables metabolic adaptation to often uneven nutrient flows (Radchuk & Borisjuk, 2014). Gradients of seed photosynthesis in terrestrial monocotyledonous plant species (e.g. barley) can lead to internal O₂ concentration gradients from high concentrations of O₂ in the photosynthetic active chlorophyll-containing pericarp (chlorenchymatic regions) to low O₂ concentrations in the starchy endosperm (storage organ) and embryo (Tschiersch *et al.*, 2011). The endosperm in monocots is thus specialized for storage activity.

Internal O₂ deficiency in the developing seed can limit respiration, which can be alleviated by seed photosynthesis improving respiration and likely higher synthesis of storage compounds positively affecting the overall yield of the developing seed (Tschiersch *et al.*, 2011; Galili *et al.*, 2014). Storage products such as carbohydrates and proteins are vital during the initial stages of seed germination. High respiration rates in the seed coat along with low tissue permeability can lead to high internal CO₂ concentrations that promote internal CO₂ re-fixation and thus enable an improved seed carbon budget (Wager, 1974; Flinn, 1985; Auras *et al.*, 1993; Vigeolas *et al.*, 2003; Rolletschek *et al.*, 2004). Rubisco activity during seed photosynthesis mediates CO₂ re-fixation, but the contribution of seed photosynthesis to biomass production via net CO₂ fixation is considered low (Goffman *et al.*, 2004; Ruuska *et al.*, 2004; Radchuk & Borisjuk, 2014). In *Z. marina* seeds, the seed sheath is green during the early-stage development (e.g. Infantes & Moksnes, 2018), indicating the presence of a chlorenchymatic layer with active photosynthesis surrounding the nongreen endosperm (i.e. the assimilating storage organ) in immature *Z. marina* seeds. However, such photosynthetic capacity of the *Z. marina* seed sheath (i.e. similar to the pericarp) and its implications for internal O₂ concentration gradients and availability in the early-stages of seed development have not been investigated.

In this study, we used high-resolution variable chlorophyll fluorescence imaging, custom-made micro-chambers for photosynthesis and respiration measurements (employing O₂ sensitive sensor foils), and chemical imaging via planar O₂ optodes to investigate the photosynthetic capacity and internal O₂ concentration gradients of *Z. marina* seeds with and without an intact seed sheath as a function of increasing irradiance. We compared such detailed metabolic activity measurements on seeds to similar measurements on early-stage seedlings. We hypothesized that immature seeds with sheath and early-stage seedlings exhibit photosynthetic capacity that can support growth of the developing seed/plant via an O₂-induced supply of respiratory energy that positively affects biosynthetic activity.

**Materials and Methods**

**Seagrass seed sampling and storage**

Flowering shoots of *Zostera marina* L. with seeds were collected at Julebæk, Helsingør, Denmark (see Brodersen *et al.*, 2020 for detailed description of the sampling site). After harvesting, the flowering shoots were stored in a large continuously aerated and illuminated (14 h : 10 h, light : dark cycle) aquarium with similar light, temperature and salinity conditions as at the sampling site (temperature = 20°C; salinity = 18; incident photon irradiance = 200 μmol photons m⁻² s⁻¹). Immature sheath-covered seeds were collected directly from the spathes. Mature seeds were collected from the bottom of smaller tanks after naturally being released from the spathes. Only the denser seeds, that is rapidly sinking to the bottom after physical resuspension, were used for this study. Seed germination (i.e. split of seed coat and hypocotyl extension) was achieved via a low salinity (drop from 30 to 10) and high temperature (increase from 9°C to 13°C) pulse (e.g. Xu *et al.*, 2016; Yue *et al.*, 2019), where after the early-stage seedlings (< 5 cm; Marion & Orth, 2012) and harvested seeds were...
used for variable chlorophyll fluorescence imaging and O2 dynamics measurements (to be described later).

Experimental setup and treatments
All seagrass seeds and early-stage seedlings were kept in filter-sterilized seawater (0.2 μm; temperature of 21°C, salinity of 18) during measurements. For measurements of photosynthetic activity: photon irradiances of 0, 9, 99, 137, 232 and 323 μmol photons m−2 s−1 (where the highest light intensity was only used for the sheath-covered seeds and the early-stage seedlings) were evenly provided by a fibre-optic tungsten halogen lamp equipped with a collimating lens (KL-2500/LCD; Schott GmbH, Mainz, Germany). The incident photon irradiance (PAR, 400–700 nm) was measured for different lamp light settings with a scalar irradiance mini-sensor (US-SQS/L; Walz GmbH, Effeltrich, Germany) interfaced to a precalibrated photon irradiance meter (ULM-500). To exclude the potential contribution of photosynthetic epiphytes on the O2 production and consumption measurements, selected seeds (n=4) were surface-sterilized via submersion into a saline c. 1.05% hypochlorite solution for 30 s, followed by 3×1 min rinses in filter-sterilized (0.2 μm) seawater before measurements (Blaabjerg & Finster, 1998; Brodersen et al., 2018). To mimic the natural in situ environment of the mature seeds and early-stage seedlings, all O2 dynamic measurements were done at low O2 conditions of <30 μmol l−1 obtained by preflushing with N2 (Brodersen et al., 2017, 2019; Schrämeyer et al., 2018; Trevathan-Tackett et al., 2020).

Variable chlorophyll fluorescence imaging
Variable chlorophyll fluorescence imaging was used to determine the photosynthetic capacity and activity of Z. marina seed sheaths, seeds with and without epiphytes, and seedlings. A pulse–amplitude–modulation (PAM)-based variable chlorophyll fluorescence imaging system (MINI-IPAM; Walz GmbH) was used with a blue LED for measuring and actinic light (Ralph et al., 2005). Before measurements, the downwelling photon irradiance levels of the actinic light for specific program settings were calibrated with a quantum meter equipped with a cosine sensor. Images of the maximum photochemical quantum yield of PSII (i.e. Fv/Fm; Table 1) were calculated as (Schreiber, 2004): Fv/Fm = (Fm – F0)/Fm, where Fm is the maximal fluorescence yield measured during a strong saturation pulse fully closing all PSII reaction centres and F0 is the minimal fluorescence yield before the saturation pulse (measured after a dark-adaptation period of c. 15 min). Average Fv/Fm values were calculated by integrating over the seed’s sheath or small photosynthetic active sites on mature seeds themselves, separately (i.e. within defined regions of interest), followed by calculating an average of all the investigated seeds for the respective development stages. Rapid light curves (RLCs) of the seeds and seedlings photosynthetic activity were measured using series of images of effective PSII photochemical quantum yields (YII) and relative photosynthetic electron transport rates (rETR) at increasing irradiance with 20 s incubation at each irradiance step (Ralph & Gademann, 2005; Trampe et al., 2011).

Table 1 Photosynthetic terms, definitions and units.

<table>
<thead>
<tr>
<th>Fv/Fm</th>
<th>Maximum photochemical quantum yield of PSII</th>
</tr>
</thead>
<tbody>
<tr>
<td>rETR</td>
<td>Relative photosynthetic electron transport rate</td>
</tr>
<tr>
<td>Y(II)</td>
<td>Effective photochemical quantum yield of PSII</td>
</tr>
<tr>
<td>Pn</td>
<td>Gross photosynthesis rate (nmol O2 mg WW−1 h−1)</td>
</tr>
<tr>
<td>α</td>
<td>Light use efficiency (μmol O2 mg WW−1 h−1)</td>
</tr>
<tr>
<td>Fmax</td>
<td>Maximum photosynthesis rate (μmol O2 mg WW−1 h−1)</td>
</tr>
<tr>
<td>EC</td>
<td>Compensation photon irradiance (μmol photons m−2 s−1)</td>
</tr>
<tr>
<td>Ek</td>
<td>Photon irradiance at onset of photosynthesis saturation (μmol photons m−2 s−1)</td>
</tr>
</tbody>
</table>

*Calculated from the exponential saturation models, here representing: (nmol O2 mg WW−1 h−1) (μmol photons m−2 s−1). Data originate from: n=6, seeds with intact seed sheath; n=3, seeds with epiphytes; n=4, surface-sterilized seeds; and n=1, seedling just after germination. Values are means ± SE (for O2 measurements normalized to wet weight, WW).
fluorescence image deconvolution into four different pigmentation types (Trampe et al., 2011). The analysis of the fluorescence excitation spectra differences between the actual measurements and a reference spectral matrix subsequently produces a colour-coded deconvoluted image, which enables identification of the abundance and distribution of diatoms, green algae, red algae and cyanobacteria. This is achieved as diatoms and green algae can be characterized by strong fluorescence excitation by blue light (460 nm) that overlaps with the absorption bands of Chl b and Chl c. Diatoms can thereafter be further distinguished by excitation with green light (525 nm) owing to presence of antenna such as fucoxanthin. Most cyanobacteria express maximal fluorescence yield with red-orange excitation (c. 620 nm), owing to phycocyanin absorption, whereas red algae due to the presence of phycoerythrin display extraordinarily high fluorescence upon excitation with green light.

Gas exchange measurements

Seed O2 production and consumption rates were measured in custom-made gas exchange chambers (1.8 ml) with presterilized (0.2 μm) seawater employing calibrated O2 sensitive sensor spots (OXSP5; PyroScience GmbH, Aachen, Germany) connected to a fibre-optic O2 sensor system (FireSting O2; PyroScience GmbH, Aachen, Germany), which was interfaced to a PC running dedicated data acquisition software (O2 logger, PyroScience). The oxygen sensor spots were 2-point calibrated in air saturated (100%) and anoxic (0%; obtained by flushing with N2) seawater at experimental temperature and salinity. Water movement in the micro-photosynthesis and respiration chambers was maintained with glass coated magnets (Spinbar®; Pyrex® Glass-coated magnetic stirring bars; Sigma-Aldrich) controlled by a magnet stirrer (IKA® big-squid; VWR International, Radnor, PA, USA). Glass coated magnets were used to avoid potential O2 consumption and/or release from the magnets during measurements. Closed gas exchange chambers (i.e., control and seagrass seed/seedling samples) were submerged into a small seawater tank, wherein the O2 concentration could be manipulated via flushing with N2. This setup constellation enabled replacing seeds and/or young seedling samples for biological replication without O2 intruding into the chambers when taking off the lid to change the samples. The O2 level in the small tank was monitored with a calibrated protected tip O2 minisensor (OXF500PT-OI; PyroScience) connected to a handheld fibre-optic O2 meter (FireSting GO2; PyroScience GmbH). Light at defined photon irradiance levels was provided by a fibre-optic tungsten halogen lamp equipped with a collimating lens (KL-2500LCD; Schott GmbH). We calculated gross photosynthesis, net photosynthesis and respiration rates from the measured O2 consumption or production rates as a function of incident irradiance as nmol O2 mg WW−1 h−1 (where controls served as blanks). In brief, the photosynthesis and respiration rates were calculated as follows: GP (B) = NP (E) + R(E) |, where respiration (R) and net photosynthesis (NP) rates were calculated from the linear O2 concentration slopes over time at the given photon irradiance (E), by multiplying the linear O2 increase or depletion rate in the chamber (nmol L−1 h−1) with the volume of seawater in the measuring chamber (L) divided by the sample wet weight. The gross photosynthesis (GP) rates were calculated by adding the absolute values of the respective respiration rates (i.e., dark respiration or postillumination respiration; measured immediately after the light period at each irradiance level and used as a proxy for the respiration in the previous light period) to the respective net photosynthesis rates (further described in Hansen et al., 2022).

Planar optode based O2 imaging

Planar O2 optodes with an isolating carbon black layer (Glud et al., 1996; Brodersen et al., 2014) and a ratiometric RGB camera system (Larsen et al., 2011) were used for O2 imaging of cross tissue sections of sheath-covered seagrass seeds. The planar optode foil provides a physical diffusion barrier between the cut seed surface and the surrounding environment, and thus largely seals-off the exposed inner part of the seed during the short-term measurements (Santner et al., 2015). The O2 sensitive planar optodes were prepared via knife-coating a sensor cocktail onto a transparent polyethylene terephthalate (PET) foil. The O2 optode sensor cocktail consisted of 1.5 mg of platinum(II)-meso(2,3,4,5,6-pentafluoro)phenyl-porphyrin (PtTPPF; indicator dye), 1.5 mg of Macrolux® fluorescence yellow 10GN (MY; reference dye), 100 mg of polystyrene (PS) and 1 g of CHCL3. After dissolution of all components in the CHCL3, the sensor solution was spread onto a dust-free PET foil using a film applicator (byk.com) yielding a c. 10 μm thick sensor film after evaporation of the solvent. To exclude background fluorescence and achieve the highest possible spatial resolution of the optical sensor, an optical isolation layer consisting of 1% w/w (10 mg) carbon black dispersed in 1 g 10% w/w solution of polyurethane hydrogel (D4; EtOH; water, 9 : 1 w/w) was knife-coated on top of the optical sensor film yielding a final isolation layer thickness of c. 7.5 μm. The ratiometric RGB camera setup consisted of a SLR camera (EOS 1000D; Canon, Tokyo, Japan) equipped with a macro-objective lens (Macro 100 f2.8D; Tokina, Tokyo, Japan), and a 530 nm long-pass filter (uggoptics.com). Excitation of the planar O2 optode was achieved via a 455 nm multi-chip LED (LedEngin Inc., RS Components Ltd, Corby, UK) combined with a bandpass filter. The excitation LED was powered by a USB-controlled LED driver unit (Triggerbox; imaging.fish-n-chips.de) interfaced with a PC running the custom software look@RGB (imaging.fish-n-chips.de) for image acquisition and control of the SLR camera and LED unit. The obtained RGB images were split into the red, green and blue channels, and analysed via the software IMAGEJ. For acquiring calibrated O2 concentration images, the red channel images (O2 sensitive emission of the indicator dye) and the respective green channel images (emission of the inert reference dye) were divided using the IMAGEJ plugin Ratio Plus (ratio = red : green), whereafter the acquired ratio images were fitted to a previously obtained calibration curve (described in Supporting Information Fig. S1) using the exponential decay function of the Curve Fitting tool in IMAGEJ.
Data analysis

The software program ORIGINPro 2017 (OriginLab Corp., Northampton, MA, USA) was used for fitting and analysing the measured photosynthesis and respiration rates, and to obtain photosynthetic parameters as follows:

An exponential saturation model (Webb et al., 1974) was fitted for the gross photosynthesis (GP) rates as a function of photon scalar irradiance (E):

\[ GP(E) = GP_{MAX} \times \left(1 - \exp \left(-\alpha \times \frac{E}{GP_{MAX}} \right) \right) \]

Similar exponential saturation model with an additional term to account for respiration (Spilling et al., 2010) was used for fitting net photosynthesis (NP) rates as a function of photon scalar irradiance (E):

\[ NP(E) = NP_{MAX} \times \left(1 - \exp \left(-\alpha \times \frac{E}{NP_{MAX}} \right) \right) + R(E) \]

where \( P_{MAX} \) is the calculated maximum photosynthesis rate at saturating photon irradiance, \( \alpha \) is the light use efficiency related to the photosynthetic activity and efficiency, \( R \) is the respiration rate, and \( E \) is the respective photon scalar irradiance.

The compensation photon scalar irradiance (\( E_C \)), that is the irradiance above which net photosynthetic production of O₂ occurs, was determined from the achieved photosynthetic parameters as:

\[ E_C = NP_{MAX} \times \log_{10} \left( \frac{1 + \frac{R}{NP_{MAX}}}{-\alpha} \right) \]

And finally, the photon scalar irradiance at the onset of photosynthesis saturation (\( E_K \)) was calculated as:

\[ E_K = NP_{MAX}/\alpha \]

All oxygen production and consumption data were normalized to mg WW⁻¹ (wet weight), to allow for utilizing the same seeds (i.e. as many as possible) for all measurements at the different development stages throughout the entire experiment, and thereby minimize biological differences between replicates.

Results

Photosynthesis and O₂ microgradients in sheath-covered seeds

Variable chlorophyll fluorescence imaging revealed photosynthetic capacity in immature sheath-covered seagrass seeds (Z. marina), with an averaged maximum PSII photochemical quantum yield of 0.61 ± 0.04 (Fig. 1a; mean ± SD; \( n = 3 \), biological replicates). Further development of the immature seed during opening of the seed sheath decreased the maximum PSII photochemical quantum yield to 0.50 ± 0.03 (Fig. 1b; mean ± SD; \( n = 6 \), technical replicates), while seeds with detached sheaths displayed no photosynthetic capacity (Fig. 1c). Rapid light curves (RLC), that is measurements of the effective PSII photochemical yield (YII) and relative photosynthetic electron transport rates (rETR) at increasing irradiance, further confirmed photosynthetic activity in Z. marina seed sheaths (Fig. 1d; \( n = 3 \), biological replicates) following a typical saturation curve with increasing irradiance. Interestingly, mature Z. marina seeds without seed sheath also exhibited photosynthetic capacity, albeit with a markedly lower maximum PSII photochemical quantum yield of 0.28 ± 0.04 and a more patchy distribution of activity over the seed surface (Fig. 1c; mean ± SD; \( n = 4 \), technical replicates).

Microscopic variable chlorophyll fluorescence imaging of cross tissue sections of sheath-covered immature Z. marina seeds showed that the photosynthetic capacity was solely restricted to the seed sheath (Fig. 2a). Photosynthetic activity of the seagrass seed sheath resulted in marked O₂ production with increasing irradiance (Fig. 2b). At saturating irradiance, the maximal net and gross photosynthesis rates of seed sheaths were 6.08 ± 0.45 and 6.61 ± 0.44 nmol O₂ mg WW⁻¹ h⁻¹, respectively, with light use efficiencies (\( \alpha \)) under subsaturating irradiance of 0.10 ± 0.02 and 0.11 ± 0.03 (Table 2; mean ± SE; \( n = 6 \), biological replicates). The compensation photon irradiance (\( E_C \)) and the onset of photosynthesis saturation (\( E_K \)) of the seed sheaths were calculated to 30.1 and 62.9 µmol photons m⁻² s⁻¹, respectively (Table 2), indicating light requirements similar to seagrass leaves. The dark respiration rate of sheath-covered seeds was -4.06 ± 0.3 nmol O₂ mg WW⁻¹ h⁻¹ (Table 2; mean ± SE; \( n = 6 \), biological replicates), whereas postillumination respiration, that is the respiration rate measured immediately after a light period under a defined irradiance, displayed a slight saturating increase with increasing irradiance (Fig. 2b). The photosynthetic activity resulted in increased internal O₂ concentrations in the centre of the immature seeds (i.e. the cotyledon and hypocotyl tissue area) enabling higher respiratory activity (Fig. 2c,d). Colour-coded O₂ images of sheath-covered seed tissue cross-sections confirmed marked differences in the internal seed O₂ concentration between measurements in light and darkness (Fig. 2c), where extracted line profiles across sheath-covered seeds showed a 5.9-fold increase in the internal O₂ concentration from 4.6 to 27.3 µmol l⁻¹ under a photon irradiance (400–700 nm) of 232 µmol photons m⁻² s⁻¹ in seed replicate 1 (Fig. 2d) and a 4.2- and 8.0-fold increase in the internal O₂ concentration from 2.8 µmol l⁻¹ in darkness to 12.1 and 22.7 µmol l⁻¹ under photon irradiances of 99 and 232 µmol photons m⁻² s⁻¹, respectively, in seed replicate 2 (Fig. 2d). Hence, the photosynthetic O₂ production of the seed sheath in the light is highly beneficial for the intra-seed O₂ availability in developing immature Z. marina seeds (Fig. 2).

Functional role of epiphytes for photosynthesis and respiration of mature seeds

Epiphyte cover on mature seagrass seed coats enabled active photosynthesis of the mature seed/epiphyte community (Figs 1c, 3, S2, S3). A biofilm of seed epiphytes covered most of the surface of the mature seed coat (Fig. 3a-d), which was dominated by
by diatoms but also harboured cyanobacteria, red and green algae (Fig. S4). The seed epiphyte layer had a maximum PSII photochemical quantum yield of up to 0.57 ± 0.01 (Fig. 3e; mean ± SD; n = 6, technical replicates) and rapid light curves (RLC), that is measurements of the effective PSII photochemical quantum yield (YII) and relative photosynthetic electron transport rates (rETR) at increasing irradiance, showed photosynthesis saturation and inhibition with increasing irradiance (Fig. 3f).

The photosynthetic activity of epiphytes on mature seeds led to marked O₂ production and emission from the seed coat (Fig. 4). Initial gas exchange measurements during light/dark transitions revealed net photosynthesis rates of up to 15.8 ± 4.8 nmol O₂ mg WW⁻¹ h⁻¹ and dark respiration rates of c. −2.5 ± 0.6 nmol O₂ mg WW⁻¹ h⁻¹ (Fig. 4a; mean ± SE; n = 6, technical replicates). Further detailed measurements and calculations of the photosynthetic O₂ production of mature seed epiphytes under increasing irradiance determined maximal net and gross photosynthesis rates of 13.28 ± 0.25 and 12.28 ± 0.82 nmol O₂ mg WW⁻¹ h⁻¹ at saturating irradiance with light use efficiencies under subsaturating irradiance of 0.13 ± 0.00 and 0.17 ± 0.02, respectively (Fig. 4b; Table 2; mean ± SE; n = 3, biological replicates). The compensation photon irradiance was calculated to 10.2 μmol photons m⁻² s⁻¹, and the onset of photosynthesis saturation first commenced at 98.6 μmol photons m⁻² s⁻¹ (Table 2).

During darkness, the averaged dark respiration rate of the seeds with epiphytes was calculated to −2.82 ± 0.09 nmol O₂ mg WW⁻¹ h⁻¹ (Table 2; mean ± SE; n = 3, biological replicates), whereas the postillumination respiration increased to −4.0 nmol O₂ mg WW⁻¹ h⁻¹ at a photon irradiance of 137 μmol photons m⁻² s⁻¹ followed by a drop in postillumination respiration towards the highest measured irradiance of 232 μmol photons m⁻² s⁻¹ (Fig. 4b).
Surface-sterilized mature seeds showed no photosynthetic capacity and activity (Fig. 4c; \( n = 3 \), biological replicates), but displayed similar dark respiration rates of \(-2.8 \pm 0.5 \text{ nmol O}_2 \text{ mg WW}^{-1} \text{ h}^{-1}\) (Fig. 4d; mean ± SE; \( n = 4 \), biological replicates) as the mature seeds with epiphytes (Fig. 4a,b). The photosynthetic \( \text{O}_2 \) production of mature \( Z. \text{marina} \) seeds is thus mainly driven by epiphytic diatoms, while the mature seeds themselves have relatively high respiratory needs.
Photosynthetic capacity and O2 balance of early-stage seedlings of *Z. marina*

The photosynthetic capacity of early-stage *Z. marina* seedlings was investigated via variable chlorophyll fluorescence imaging and gas exchange measurements (Fig. 5). The basal hypocotyl (ROI 1), cotyledonary sheath (ROI 2) and cotyledonary blade (ROI 3) tissue area exhibited a maximum PSII photochemical quantum yield of 0.50, 0.44 and 0.30, respectively (Fig. 5a). The seedling tissue region with active root formation displayed no photosynthetic capacity (Fig. 5a). Rapid light curves showing effective PSII photochemical quantum yields (YII) and relative photosynthetic electron transport rates (rETR) under increasing photon irradiance also confirmed photosynthetic activity in early-stage seedlings, where earlier photosynthesis inhibition with increasing irradiance was observed in the basal hypocotyl and cotyledonary sheath, as compared to the cotyledonary blade tissue area (Fig. 5b). Such photosynthetic capacity of the cotyledonary tissue was likely supported by the innermost growing first true leaf (Fig. 5c). Microscopic variable chlorophyll fluorescence imaging of the cotyledonary blade apex showed that the photosynthetic capacity ceased at the tip of the cotyledonal blade (Fig. 5c). The main function of the cotyledonal blade apex may thus be sensing light and/or gravity. The variable chlorophyll fluorescence imaging also showed indications of minor and heterogeneous epiphyte growth on the cotyledonal tissue surface (Fig. 5c). Measurements of photosynthetic O2 production and respiration of the early-stage seedling under increasing irradiance revealed a relatively high respiratory demand and low photosynthetic activity of the seedling tissue with maximal net and gross photosynthesis rates of 3.23 ± 2.93 and 1.98 ± 0.63 nmol O2 mg WW−1 h−1 at high irradiance (Fig. 5d; Table 2; mean ± SE of the nonlinear curve fit; n = 1, biological replicate). Light utilization efficiencies were also very low, that is 0.01 ± 0.00 for both net and gross photosynthesis rates (Fig. 1d; Table 2; mean ± SE of the nonlinear curve fit; n = 1, biological replicate), leading to very high calculated compensation (EC) and onset of photosynthesis saturation (Ec) irradiances of 125.9 and 405.0 μmol photons m−2 s−1, respectively (Table 2), as compared to seeds with sheath and epiphyte cover. The dark respiration rate of the early-stage seedling was −1.65 ± 0.17 nmol O2 mg WW−1 h−1 (Table 2; mean ± SE of the nonlinear curve fit; n =

### Table 2 Multiple photosynthetic parameters derived from the light response curves in Figs 2, 4 and 5.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pn (nmol O2 mg WW−1 h−1)</td>
<td>0.10 ± 0.02</td>
</tr>
<tr>
<td>Pc (nmol O2 mg WW−1 h−1)</td>
<td>0.11 ± 0.03</td>
</tr>
<tr>
<td>α</td>
<td>6.08 ± 0.45</td>
</tr>
<tr>
<td>R (nmol O2 mg WW−1 h−1)</td>
<td>−4.06 ± 0.3</td>
</tr>
<tr>
<td>EC (μmol photons m−2 s−1)</td>
<td>30.1</td>
</tr>
<tr>
<td>EK (μmol photons m−2 s−1)</td>
<td>62.9</td>
</tr>
<tr>
<td>Seeding</td>
<td></td>
</tr>
<tr>
<td>Pn (nmol O2 mg WW−1 h−1)</td>
<td>0.13 ± 0.00</td>
</tr>
<tr>
<td>Pc (nmol O2 mg WW−1 h−1)</td>
<td>0.17 ± 0.02</td>
</tr>
<tr>
<td>α</td>
<td>13.28 ± 0.25</td>
</tr>
<tr>
<td>R (nmol O2 mg WW−1 h−1)</td>
<td>−2.82 ± 0.09</td>
</tr>
<tr>
<td>EC (μmol photons m−2 s−1)</td>
<td>10.2</td>
</tr>
<tr>
<td>EK (μmol photons m−2 s−1)</td>
<td>98.6</td>
</tr>
<tr>
<td>Surface-sterilized seeds</td>
<td></td>
</tr>
<tr>
<td>Pn (nmol O2 mg WW−1 h−1)</td>
<td>−2.8 ± 0.5</td>
</tr>
<tr>
<td>Pc (nmol O2 mg WW−1 h−1)</td>
<td>3.23 ± 2.93</td>
</tr>
<tr>
<td>α</td>
<td>−1.65 ± 0.17</td>
</tr>
<tr>
<td>R (nmol O2 mg WW−1 h−1)</td>
<td>125.9</td>
</tr>
<tr>
<td>EC (μmol photons m−2 s−1)</td>
<td>405.0</td>
</tr>
</tbody>
</table>

Data originate from seeds with intact seed sheath, seeds with epiphytes, surface-sterilized seeds and young seedlings just after germination. Abbreviations refer to: Pn, net photosynthesis; Pc, gross photosynthesis; α = initial slope of the PI-curve in the light-limiting phase; Pmax, maximum net photosynthesis rate; R, dark respiration rate; EC, compensation photon irradiance; Ec, onset of photosynthesis saturation. Values are means ± SE of: (i) six seeds with intact seed sheath, (ii) three *Z. marina* seeds with epiphytes, (iii) four surface-sterilized seeds and (iv) one seedling just after germination. Data originate from low O2 conditions in the seawater to mimic the seeds/seedlings natural sediment environment (Figs 4a, b, d); except for the surface-sterilized seeds dark respiration (Fig. 4d), as no O2 production was found for surface-sterilized seeds during light illumination (Fig. 4c), and for seeds with sheath as they are normally growing in a naturally air saturated water-column environment (Fig. 2b). All data have been normalized to: mg WW−1.
Fig. 3 Photosynthetic capacity of mature seagrass seed with epiphytes. (a, b) Image of mature Zostera marina seed showing the position of epiphytes on the seed coat. (c, d) Fluorescence microscope images of epiphyte chloroplasts, showing the seed coat surface (c) and a cross-section of the seed coat (d). (e) Microscopic variable chlorophyll fluorescence measurements of the PSII photochemical efficiency, that is the maximum quantum yield of PSII, $F_{v}/F_{m}$, of seed coat epiphytes (mainly consisting of diatoms, but also cyanobacteria, green and red algae, Supporting Information Fig. S4). The legend depicts the $F_{v}/F_{m}$ value. Further biological replication can be found in the Figs S2 and S3. (f) Rapid light curves of the effective quantum yield of PSII [Y(II)] and the relative electron transport rate (rETR) of the seed epiphytes ($n = 3$, mean ± SD).
Early-stage seedlings are thus able to produce some O$_2$ for their respiratory needs during growth, which likely is supported by photosynthesis of the innermost first true leaf, albeit with much lower production rates as compared to the seed sheath photosynthetic capacity.

**Discussion**

Our results provide first experimental evidence that seagrass (*Z. marina*) seed sheaths produce O$_2$ in the light, leading to markedly increased internal O$_2$ availability inside the cotyledon and hypocotyl tissue area of the seed (i.e. embryo), which ensures sufficient O$_2$ support for respiration and thereby likely energy for biosynthetic activity in the developing seeds.

**Photosynthetic capacity and internal O$_2$ gradients in developing seagrass seeds**

Illumination of *Z. marina* seed sheaths led to strong gradients of seed photosynthesis and internal O$_2$ concentrations in the immature seagrass seeds, from relatively high O$_2$ production and concentration areas in the chlorophyll-containing and photosynthetic active
surrounding seed sheath to hypoxic conditions in the inner embryo and endosperm tissue. After opening and detachment of the seed sheath, the photosynthetic capacity of the seed sheath ceased. Such seed sheath photosynthetic \( \text{O}_2 \) production likely acts to relieve severe internal hypoxic conditions and stress in the developing seagrass seed, which may increase the supply of respiratory energy and thereby positively affect the seeds biosynthetic activity and thus carbohydrate reserves stored in the endosperm (Borisjuk & Rolletschek, 2009). This was supported by a 19% higher respiration rate of sheath-covered seeds in light as compared to in darkness (Student’s \( t \)-test: \( P = 0.04 \); Fig. S7). Such \( \text{O}_2 \) evolution capacity in sheath-covered seagrass seeds is similar to observations in immature barley caryopsis seeds, which exhibit (i) net \( \text{O}_2 \) production at saturating light conditions (\( c. 350 \mu \text{mol photons m}^{-2} \text{s}^{-1} \)), (ii) gross photosynthesis rates of 1.8 \( \mu \text{mol O}_2 \text{ grain}^{-1} \text{ h}^{-1} \) and respiratory \( \text{O}_2 \) demand of 0.4 \( \mu \text{mol O}_2 \text{ grain}^{-1} \text{ h}^{-1} \), and (iii) cessation of photosynthetic activity during maturation (Tschiersch et al., 2011). The seagrass seed sheath gross photosynthesis rates also appeared similar to what has been measured in other \( \text{O}_2 \) producing seeds like soybeans and peas (Rolletschek et al., 2002, 2005b; c. 50–110 nmol \( 2 \text{ g}^{-1} \text{ fresh weight} \text{ min}^{-1} \); however, no direct comparison is possible due to different units used for normalization). The seagrass seed coat and sheath thus serve several physiological functions that have evolved to promote the development of the seed. Here, we speculate that seed sheath photosynthetic activity plays a vital role for endosperm storage activity, while the seagrass seed coat provides protection for the mature seed, which enables dormancy and seed dispersal and thus is of vital importance for successful establishment of seedlings in adjacent sediment areas. Comparing to terrestrial seeds, the seagrass seed sheath photosynthetic activity may actually be even more important for the developing seed biosynthetic activity as gas diffusion in water is \( c. 10,000 \) times slower than in air (Madsen & Sand-Jensen, 1991) and thus can largely restrict the passive support of \( \text{O}_2 \) from the water-column to the surface of the seed coat, especially under low water velocities within the seagrass meadow.

Fig. 5 Photosynthetic capacity and oxygen production in seagrass seedling. (a) Maximum quantum yield of PSII \( (F_v/F_m) \) of young Zostera marina seedling after germination. BH, basal hypocotyl (ROI 1); AH, axial hypocotyl; CS, cotyledonary sheath (ROI 2); CB, cotyledonary blade (ROI 3). A more detailed description of the seedling anatomy and development stages is provided in Supporting Information Fig. S6. (b) Effective quantum yield of PSII \( (Y(\text{II})) \) and relative electron transport rate \( (r\text{ETR}) \) within the three selected regions of interest (ROIs 1–3) on the seedling as shown in panel A (\( n = 3 \)), determined at increasing irradiance via rapid light curves. (c) Microscopic variable chlorophyll imaging showing close-up image of the photosynthetic capacity \( (F_v/F_m) \) of the tip of the seedling cotyledonary blade with inner growing first true leaf. (a, c) Legend depicts the \( F_v/F_m \) value. Greyscale pictures show the position of the seedling (\( Z. \text{ marina} \)) and the cotyledonary blade apex. (d) Oxygen production and respiration rates of the seedling at increasing irradiance. The measured respiration, net- and gross photosynthesis rates are expressed as \( \text{O}_2 \) evolution \( \text{mg}^{-1} \) biomass (WW). Photosynthesis values were fitted with an exponential saturation function (Webb et al., 1974), with an added respiration term \( (R) \) to account for respiration in the case of net photosynthetic rates (Spilling et al., 2010). \( R^2 \geq 0.9 \).
Impacts of epiphytic photosynthesis on the success of seed germination

Surface-sterilized, mature seagrass seeds did not exhibit photosynthetic capacity, however, showed sustained respiratory activity. Seed coat epiphytes (mainly diatoms) led to photosynthetic O₂ production on the seed coat surface, which slightly enhanced postillumination respiration rates of mature seagrass seeds, while it had no effects on the dark respiration rate. However, no significant difference was found between respiration rates in light and darkness of mature seeds with epiphytes (Student’s t-test: P = 0.76; Fig. S7). Nevertheless, whether such epiphyte-induced O₂ production can support the mature seeds respiratory demand and thereby biosynthetic activity remains largely unknown, as it may depend on the epiphyte biomass and community composition, and deserves further attention in future studies.

Epiphytic photosynthesis on the seed coat of mature seeds could also play a protective role via oxidation of phytotoxic compounds such as H₂S during seed germination in the sediment surface layers. Such plant-derived oxic microhabitats in the sediment have been shown to be very important for seagrasses, as it functions as a chemical and biological defence mechanism in the seagrass rhizosphere against toxic H₂S intrusion (Brodersen et al., 2015, 2018), and can also solubilize essential nutrients for growth (Brodersen et al., 2017). Nevertheless, the observed respiratory need of mature seagrass seeds may explain why seeds germinate earlier in anoxic sediment as compared to more oxygenated sediment conditions (e.g. Kawasaki, 1993; Moore et al., 1993; Orth et al., 2000), simply as a necessity for ensuring sufficient O₂ support and respiratory energy for biosynthetic activity during germination and establishment of the seedling. Such demand for respiration energy in mature seed is supported by previous findings of improper development and reduced survival of Z. marina seeds exposed to anaerobic conditions (Hootsmans et al., 1987; Churchill, 1992), which could be linked to increased internal reactive oxygen species (ROS) formation (e.g. Perez-Perez et al., 2012).

Comparison of metabolic activity in seeds and early-stage seagrass seedlings

Early-stage Z. marina seedlings showed photosynthesis capacity and activity of the hypocotyl and cotyledonary tissues, except at the site of adventitious root formation and at the apex of the cotyledonary blade. Generally, the cotyledonary tissue of monocotyledons (grass-like flowering plants such as seagrasses) is not considered photosynthetic, in contrast to dicot seedlings whose cotyledons display photosynthetic capacity (Ampofo et al., 1976; Harris et al., 1986; Brown & Huber, 1987; Zheng et al., 2011). The higher photosynthetic capacity at the base of the cotyledonary blade, as compared to along the cotyledonary blade, could be derived from the innermost growing first true seagrass leaf, which elongates from the inside of the cotyledonal sheath (Xu et al., 2016). In addition to this photosynthetic activity by the first true leaf/leaves, some epiphytic growth and photosynthetic activity also seemed to take place along the cotyledonal blade (see Fig. 5c). Hypocotyl photosynthesis has been shown in other plants, like in the green hypocotyl tissue of pine seedlings but is considered less important than cotyledon photosynthesis for seedling growth and development (Sasaki & Kozlowski, 1970).

The importance of cotyledon and hypocotyl photosynthesis on seedling development has largely been attributed to mobilization of stored reserves to support early seedling growth (Ampofo et al., 1976; Harris et al., 1986; Brown & Huber, 1987; Zheng et al., 2011). In soybeans, the cotyledons synthesize chlorophyll upon emergence from the soil, and cotyledon photosynthesis contributes carbon via assimilation to the developing seedling (Brown & Huber, 1987), albeit cotyledon photosynthesis rates are very low as compared with the true leaves (Harris et al., 1986). In castor seedlings, cotyledon photosynthesis provides carbohydrate and energy for the first true leaf to appear and maintains seedling growth until the first true leaf has expanded (Zheng et al., 2011). In castor seedlings, the cotyledon photosynthesis ceases during seedling development, however, cotyledon photosynthesis is sufficient to balance respiratory losses during early-stage seedling establishment (Zheng et al., 2011). Cotyledon photosynthates are essential for leaf production in Acer, where the photosynthates (such as sucrose) is first exported to the developing first leaf and subsequently to the hypocotyl and roots (Ampofo et al., 1976). Thus, similar functions of the determined cotyledon and hypocotyl photosynthesis in early-stage seagrass seedlings are likely. We found no photosynthetic capacity in the cotyledonal blade apex, and we speculate that this region of the cotyledonal tissue mainly plays a role for light and gravitropic responses during early-stage seagrass seedling growth, as observed in Arabidopsis where photoreceptors regulate the development of hypocotyls and cotyledons as a response to light conditions (Sullivan & Deng, 2003) and generally in higher plant coleoptiles and root caps containing statocytes that are involved in graviperception (Raven & Edwards, 2001).

Early-stage Z. marina seedlings exhibited lower photosynthetic efficiency (α), maximum net photosynthesis rate (P max) and respiration rate (R), as compared to Z. marina seeds with sheath, which resulted in markedly higher photosynthesis saturation (Eₚ) and compensation photon irradiance (E_c) of the young seedling. The respiration rate and compensation photon irradiance of the immature sheath-covered seeds were markedly higher (i.e. 1.4- and 3-fold, respectively) indicating high metabolic activity in the developing seagrass seeds. Hence, the functioning of the seagrass seed sheath photosynthesis seems to be like in other angiosperms by likely enhancing the endosperm storage activity, while the development and growth of the young seagrass seedlings is supported by both cotyledon and hypocotyl photosynthesis, as well as, by stored energy reserves.

In summary, we have demonstrated that the sheath of seagrass (Z. marina) seeds has photosynthetic capacity, which results in increased O₂ availability for the plant embryo in the light and can alleviate intra-seed hypoxic stress conditions that may allow for increased biosynthetic activity. The photosynthetic architecture of the immature seagrass seeds drives strong gradients of photosynthesis and O₂ concentrations across the seed tissues, from high O₂ levels in the photosynthetic active tissue of the seed sheath to low O₂ levels in the nonphotosynthetic embryotic
centre of the seed. Such photosynthetic capacity of the developing seagrass seed ceases during maturation with the detachment of the seed sheath. Early-stage seedlings also possess photosynthetic capacity, which seemed supported by the development of the first green leaf inside the cotyledonary sheath, where the lack of photosynthetic activity in the cotyledonary blade apex suggest that this part of the cotyledon may rather function as a sensing organ for, for example light and/or gravity. The resulting increased respiratory energy could enhance nutrient transport and endosperm storage capacity of the seagrass seed and seedling, and may be vital for the successful transition from seed to established seedling and thereby for the long-term performance of seagrass meadows.

Acknowledgements
The research was funded by grants from the Carlsberg Foundation (CF16-0899; KEB), the Villum Foundation (00028156; KEB) and the Independent Research Fund Denmark (DFF-8022-00301B; MK).

Competing interest
None declared.

Author contributions
KEB planned and designed the research with support from MK. KEB performed the experiments. KEB processed and analysed the data. KEB wrote the manuscript with editorial help from MK. Both authors have given approval to the final version of the manuscript.

ORCID
Kasper Elgetti Brodersen https://orcid.org/0000-0001-9010-1179
Michael Kühl https://orcid.org/0000-0002-1792-4790

Data availability
All data are included as part of the manuscript or as Supporting Information.

References


**Supporting Information**

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

**Fig. S1** Planar optode calibration plot and fitting.

**Fig. S2** Photosynthetic capacity of seeds with epiphytes #2 and 3.

**Fig. S3** Variable chlorophyll fluorescence and RLC of seeds with epiphytes #4–6.

**Fig. S4** RGB function fits as indicator of the seed epiphyte community composition (*n* = 3).

**Fig. S5** *Zostera marina* seed development stages and close-up images of *F*<sub>v</sub>/*F*<sub>m</sub>.

**Fig. S6** *Zostera marina* seedling anatomy and development.

**Fig. S7** Dark and light respiration of sheath-covered seeds and seeds with epiphytes (*n* = 3).

Please note: Wiley is not responsible for the content or functionality of any Supporting Information supplied by the authors. Any queries (other than missing material) should be directed to the *New Phytologist* Central Office.