

1 **Microscale light management and inherent optical properties of**
2 **intact corals studied with optical coherence tomography**

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14

15 **Abstract**

16 Coral reefs are highly productive photosynthetic systems and coral optics studies suggest that such high
17 efficiency is due to optimised light scattering by coral tissue and skeleton. Here, we characterise the
18 inherent optical properties, i.e., the scattering coefficient, μ_s , and the anisotropy of scattering, g , of 8
19 intact coral species using optical coherence tomography (OCT). Specifically, we describe light scattering
20 by coral skeletons, coenoarc tissues, polyp tentacles and areas covered by fluorescent pigments (FP). Our
21 results reveal that light scattering between coral species ranges from $\mu_s = 3 \text{ mm}^{-1}$ (*Stylophora pistillata*)
22 to $\mu_s = 25 \text{ mm}^{-1}$ (*Echinopora lamellosa*). For *Platygyra pini*, μ_s was 10-fold higher for tissue vs skeleton,
23 while in other corals (e.g. *Hydnophora pilosa*) no difference was found between tissue and skeletal
24 scattering. Tissue scattering was 3-fold enhanced in coenosarc tissues ($\mu_s = 24.6 \text{ mm}^{-1}$) vs polyp tentacles
25 ($\mu_s = 8.3 \text{ mm}^{-1}$) in *Turbinaria reniformis*. FP scattering was almost isotropic when FP were organized in
26 granule chromatophores ($g=0.34$) but was forward directed when FP were distributed diffusely in the
27 tissue ($g=0.96$). Our study provides detailed measurements of coral scattering and establishes a rapid
28 approach for characterising optical properties of photosynthetic soft tissues via OCT *in vivo*.

29

30 **Introduction**

31 The form and function of an organism represents a design solution to the problems posed by a multitude
32 of environmental parameters. The evolutionary design of terrestrial plants has been studied over decades,
33 providing evidence on the prime role of irradiance exposure, hydration and mechanical stability in
34 driving plant morphology on cellular to canopy scales [1, 2]. On the scale of a plant leaf, studies showed
35 that epidermal cells act similar to a lens and can focus incident radiation [3]. Light focused by epidermal
36 cells can then be channelled deep into the plant leaf via lossy scattering between air-filled vacuoles and
37 the palisade layer (i.e. a porous matrix) [4]. Such light propagating mechanism is more pronounced in
38 sun adapted leaves that are in need of effectively distributing excess irradiance [4] compared to shade
39 adapted leaves which did not have such well-defined palisade mesophyll and instead developed a spongy
40 mesophyll, which scatters less light [5]. It is also known that some succulents that are exposed to high
41 light (e.g. *Dudleya brittonii*) can develop wax cuticles that effectively attenuate UV light and could thus
42 act photoprotective [6]. Thus for terrestrial plants, the plant leaf is highly specialised in optimising
43 irradiance exposure for chloroplast photosynthesis [3].

44 On tropical coral reefs, corals have evolved as highly efficient photosynthetic organisms [7].
45 Similar to plants, corals can be subject to variable irradiance regimes. Shallow water corals are exposed
46 to excess irradiance, reaching about $2000 \mu\text{mol photons m}^{-2} \text{s}^{-1}$ during mid-day sun and low tide [8]. In
47 contrast, deep water corals can live photosynthetically under mesophotic conditions leaving only about
48 3% of surface irradiance at a depth of over $>100 \text{ m}$ [9]. While it is well-known that the coral animal's
49 symbiotic microalgae (*Symbiodinium* sp.) adjusts pigment density (specifically chlorophyll *a*) to
50 optimise light absorption for photosynthesis [10] there is now mounting evidence that the light scattering

51 properties of coral skeletons and tissues also play a central role in light propagation and coral
52 photosynthesis [11-16].

53 Studies with fiber-optic light microsensors showed that the coral tissue surface scalar irradiance
54 (i.e. the total quantum flux in a point incident from all directions) can be about twice of the incident
55 downwelling irradiance [11]. Similar to plant tissues, coral tissues are suggested to be strongly light
56 scattering matrices [15]. Light scattering affects the photon pathlength and residence time in a given
57 tissue layer and thus the chance for light absorption and subsequent O₂ evolution or heat dissipation.
58 Monte Carlo simulations (i.e. probabilistic light propagation models [17]) were developed to quantify
59 the scattering probability of coral tissue and skeletons [16]. It was found that the scattering coefficient,
60 μ_s [mm⁻¹], i.e., the ability to scatter photons over a certain distance, of the massive coral *Favites abdita*,
61 was about 3 times higher in the tissue than in the underlying skeleton, indicating that light is effectively
62 homogenized by the tissue and then propagated by the skeleton [16]. Other studies investigated light
63 scattering from a museum collection of coral skeletons and showed that scattering is highly variable
64 between coral skeletons [14, 18]. Differences in light scattering were related to coral bleaching
65 susceptibility [14] and corals with low light scattering ability appeared more susceptible to coral
66 bleaching [19]. Despite a few studies that have extracted the scattering coefficient of coral skeletons [14,
67 16, 19, 20] the current data on intact corals and coral tissues is very limited [16].

68 Characterisation of optical properties of soft tissues in their intact state is difficult given that many
69 techniques rely on tissue preparation that can result in changes in tissue structure and hydration with the
70 potential to alter tissue optical properties [21]. In the field of biomedical tissue optics, several approaches
71 have been developed for extracting optical properties of intact tissues. A common technique is diffuse
72 optical spectroscopy, where the lateral attenuation of diffuse reflectance is measured with fibers placed

73 at several radial distances [21]. However, a key requirement for using diffusion theory is that the
74 collected light is entirely diffuse, i.e., has lost directionality. Under the assumption of diffuse light, the
75 diffuse scattering coefficient can be extracted as $\mu_s' = \mu_s(1-g)$, where g is the so-called anisotropy factor.
76 The g value [dimensionless] is defined as the mean cosine of the scattering angle θ [22, 23], which
77 describes the amount of directionality retained after a single scattering event such that $g = 1$ for entirely
78 forward scattering, $g = 0$ for isotropic scattering, and $g = -1$ for entirely backward scattering. The
79 diffusion approximation thus lumps together μ_s and g [24].

80 More recently, optical coherence tomography (OCT) has been used to extract the optical
81 properties of human tissues and optical phantoms [25, 26]. OCT is a non-invasive imaging technique
82 that generates high resolution tomographic images using low coherent near infrared radiation (NIR). It
83 measures characteristic patterns of directly elastically backscattered (low coherent ballistic and near-
84 ballistic) photons from different reflective layers in a sample, e.g. at refractive index mismatches between
85 tissue compartments with different microstructural properties [27]. If the OCT light source and imaging
86 optics are calibrated with respect to absolute reflectivity, OCT can be used to obtain quantitative
87 information on the scattering properties of the sample [25, 26, 28]. Levitz et al. [25] developed a
88 theoretical model, based on inverse Monte Carlo modelling, to fit the depth dependent attenuation of the
89 OCT signal and the local OCT signal intensity to extract μ_s and g , respectively (see methods for details).
90 If the investigated tissue/structure does not absorb strongly in the wavelength range of the light source,
91 light attenuation in OCT is primarily a function of light scattering, where the local reflectivity quantifies
92 the amount of directly backscattered photons, which can thus be used to quantify the g value [25] as a
93 decrease in g causes an increase in the local reflectivity and *vice versa*. Another advantage of OCT is

94 that it allows for very localised extraction of optical properties, while e.g. diffusion theory averages the
95 optical properties over the measurement area (usually encompassing several cm² of surface area) [21].

96 In the present study, we use OCT to characterise the optical scattering properties of intact coral
97 tissues and skeletons *in vivo*. The specific aims are to study the variability in the scattering coefficient
98 μ_s and g value of 8 coral species with a specific focus on differences between tissue and skeletal scattering
99 as well as differences among tissue types.

100 **Methods**

101 ***Coral specimens.*** The corals used in this experiment originated from the coral culture at the Centre
102 Scientifique de Monaco. Corals were kept in aquaria supplied with Mediterranean seawater (exchange
103 rate 70%/h) at a salinity of 38, temperature of 25°C, and photon irradiance (PAR, 400-700 nm) of 200
104 $\mu\text{mol photons m}^{-2} \text{ s}^{-1}$ on a 12-h/12-h photoperiod as provided by HQI-10000K metal halide lamps (BLV
105 Nepturion). Corals were fed twice a week with *Artemia salina nauplii*. We selected several coral
106 fragments from the following coral species: *Echinopora lamellosa*, *Platygyra pini*, *Hydnophora pilosa*,
107 *Pavona cactus*, *Turbinaria reniformis*, *Acropora sp.*, *Montipora capricornis* and *Stylophora pistillata*. The
108 corals were chosen to represent a diversity of coral species with different tissue thicknesses, host
109 pigmentation as well as skeletal optical properties [14]. At least three coral fragments of each species
110 were used for optical extraction.

111 ***Theory of optical parameter extraction based on OCT.*** The basic OCT operating principle is explained
112 elsewhere [27]. OCT can be used to extract the scattering coefficient μ_s and anisotropy of scattering, g ,
113 from biological tissues using a theoretical model of OCT light propagation derived from the inverse

114 Monte Carlo method [25, 26, 28]. For a homogenous biological tissue, the depth dependent OCT
115 reflectance signal, $R(z)$, is thereafter described as a simple exponential decay:

116
$$R(z) = \rho \cdot e^{-\mu \cdot z} \quad (1)$$

117 where ρ is the fraction of light sampled from the focal volume of tissue, and μ is the signal attenuation
118 to and from the focal volume (see Levitz et al.[25] for details). Further,

119
$$\rho = \mu_s \cdot b(g) \cdot \Delta z \quad (2)$$

120 and

121
$$\mu = 2 \cdot G \cdot [\mu_s \cdot a(g) + \mu_a], \quad (3)$$

122 Here, Δz is the axial resolution of the imaging system given by the coherence length l_c :

123
$$l_c = 0.44 \cdot \lambda / \Delta \lambda \quad (4)$$

124 where λ is the center wavelength of the light source (930nm) and $\Delta \lambda$ is the spectral bandwidth (63.4 nm).
125 The parameter $b(g)$ is the scatter collection efficiency factor that describes the fraction of light scattered
126 within the coherence gate (i.e. the spatial distance over which the reflected light and reference beam can
127 cause interference signals) that is backscattered within the solid angle of collection by the objective lens.
128 G is a geometry factor that accounts for the enhanced pathlength due to off-axis light propagation during
129 delivery and collection by the objective lens. For an objective with low numerical aperture (NA), G is
130 approximated by $1/\cos[\sin^{-1}(\text{NA})]$ [25]. The theoretical model further assumes a Henyey-Greenstein
131 scattering phase function $p(\theta)$ to describe the scattering of coral tissues, given that this phase function is
132 most commonly used to describe the scattering of biological tissues [22, 29]. The phase function $p(\theta)$

133 affects the parameter $b(g)$ by:

134
$$b(g) = \int_{\pi - \sin^{-1}(NA)}^{\pi} p(\theta) \cdot 2\pi \cdot \sin(\theta) d\theta \quad (5)$$

135 The factor a in Eq. 3 is the scattering efficiency factor that depends on g . It determines the ability of
136 photons to reach the focus of the OCT system despite scattering; when $a = 1$ then $g = 0$ and *vice versa*
137 [30]. The function $a(g)$ can be approximated by Monte Carlo simulations [25] yielding:

138
$$a = 1 - \exp\left(-\left((1 - g)^{0.6651}\right)/0.1555\right) \quad (6)$$

139 ***OCT imaging and signal calibration.*** OCT imaging was performed on corals as described previously
140 [31]. Briefly, imaging was done with a commercially available spectral-domain (SD) OCT system
141 (Ganymede II, Thorlabs GmbH, Dachau, Germany) equipped with an objective lens with an effective
142 focal length of 36 mm, and a working distance of 25.1 mm (LSM03; Thorlabs GmbH, Dachau, Germany;
143 Fig. 1a). The system used a 930 nm light source, yielding a maximal axial and lateral resolution in water
144 of 5.8 μm and 8 μm , respectively. OCT b-scans were acquired at a fixed pixel size of 581 x 1023, while
145 the actual field of view was variable in y but fixed in z ($z=2.2$ mm). OCT imaging was performed for
146 different coral species in a black acrylic flow chamber supplied with aerated seawater. For all
147 measurements the OCT system was optimised to yield highest OCT signal measurements at a fixed
148 distance, chosen to be in the upper 1/3 of the OCT image (at 0.4 mm from the top). To calibrate the OCT
149 reflectivity, calibration measurements were performed prior to measurements for each coral species using
150 a reflectance standard composing of (1) oil-glass interface, (2) water-glass interface, and (3) air-glass
151 interface. The reflectivity from the reflectance standard was calibrated via Fresnel's equations for the
152 case of normal incidence of light using the refractive index of air ($n=1$), water ($n=1.33$), oil ($n=1.46$) and

153 glass ($n=1.52$). The reflectance of the oil-glass interface was thus $((1.46-1.52)/(1.46+1.52))^2 = 4.05 \times 10^{-4}$,
154 while the reflectance from a mirror = 1 (Fig. S1). The OCT signal from measurements on corals,
155 OCT(dB), was converted to R via a linear fitting procedure, where the relation of OCT(dB) to $\log_{10}(R)$
156 was described by linear function, as both parameters are logarithmic. To calibrate for the focus function
157 of the objective lens, calibration measurements were performed for 8 different z positions (0.0 – 0.8 mm
158 from the image top in steps of 0.1 mm). Such measurements showed that the OCT signal fall-off, from
159 the z position of peak reflectivity ($z= 0.4$ mm) to $z=0.8$ mm, was exponential, facilitating a straight
160 forward correction of acquired scans for the focus function of the OCT system.

161 **Optical extraction.** After OCT scans were corrected for the focus function (OCT dB) and converted to
162 absolute values of reflectivity ($\log_{10}(R)$) dedicated coral tissue areas were selected for optical extraction.
163 Areas were selected to cover different tissue types, including polyp and coenosarc tissues as well as
164 tissues covered with host pigment granules. Values of ρ (local reflectivity, dimensionless) and μ (linear
165 signal attenuation [cm^{-1}]) were matched to g and μ_s using the theory described above [25, 30]. The
166 absorption coefficient, μ_a , by coral and algal pigments at 930 nm is negligible [14, 16] and the fitting
167 assumed that absorption was dominated by water, where $\mu_a= 0.43 \text{ cm}^{-1}$ for water at 930 nm [22]. The
168 water content of the tissue was assumed to be 50%, based on the hydration of human tissues [22]. The
169 effective numerical aperture was 0.15 and the OCT coherence length was 6 μm . Based on these settings
170 a grid method was applied [25] to extract the optical parameters g and μ_s . Briefly, the grid method is a
171 lookup table of experimental values of μ versus ρ generated by choosing values of μ_s and g for use in
172 Monte Carlo simulations to create $R(z)$ curves that are fit using Eq. 1 to yield $\mu(\mu_s, g)$ and $\rho(\mu_s, g)$ (Fig.
173 1). When the $\mu(\mu_s, g)$ and $\rho(\mu_s, g)$ values are plotted, and the iso- μ_s and iso- g lines are drawn, a grid is
174 formed. A pair of observed μ, ρ values now uniquely maps to a pair of μ_s, g values [25]. A 2D interpolation

175 algorithm implements the mapping, specifying $\mu_s(\mu, \rho)$ and $g(\mu, \rho)$. More details on the grid can be found
176 in [25, 30].

177 Optical extraction involved curve fitting for randomly selected tissue and skeletal spots. Curve
178 fitting was considered satisfactory if the model matched the data with a R^2 value >0.5 . The first few
179 pixels at the tissue surface were excluded from the curve fitting procedure [32], given the strong
180 reflectivity spike due to the refractive index mismatch between tissue and water. Preliminary analyses
181 showed that the OCT signal often did not attenuate within the first 20-50 μm of the coral tissue (Fig. 2).
182 Maximal OCT signal was found at about 50 μm depth and curve fitting used maximal reflectivity values
183 that showed a smooth signal fall off. We found that optical extraction for regions of interest positioned
184 deeper within the tissue, e.g. underlying coral skeletons, would not yield reasonable results due to OCT
185 signal attenuation by upper tissue layers. More robust estimates of coral skeleton optical properties were
186 obtained from OCT scans of bare skeletons, where the tissue was removed with an air gun [15].

187 **Results and discussion**

188 *Light scattering in tissue vs skeleton.* The inherent optical properties of coral tissues are largely
189 unknown, although they are fundamental for a better mechanistic understanding of coral light absorption
190 and photosynthesis, and thus coral ecophysiology [16, 33, 34]. Previous studies have primarily
191 characterised the apparent optical properties of coral tissues, i.e., light field parameters such as the scalar
192 irradiance and reflectance, suggesting that coral tissues are highly scattering [15, 35]. However, only one
193 study has presented data on the scattering coefficient of intact coral tissues, providing evidence for a high
194 reduced scattering coefficient, μ_s' , of a massive faviid coral tissue [16]. In the present study,
195 determination of μ_s in intact coral tissue for 8 corals showed high variability of μ_s from 4 to 25 mm^{-1} at

196 930 nm (Table 1). In comparison, μ_s of plant tissues ranges from about 5 to 10 mm^{-1} [36, 37], while
197 human skin can reach values of about 20 mm^{-1} [22]. Thus for the case of a highly scattering coral tissue,
198 such as in *Platygyra pini* (Fig. 2), the average distance between scattering events, i.e, the scattering mean
199 free path, $\text{MFP}=1/\mu_s$, is 46 μm (Table 1), which is about 2-fold lower than for most biological tissues
200 ($\text{MFP}\sim 100\ \mu\text{m}$) [38]. In contrast, the scattering strength of the coral *Stylophora pistillata* would rank
201 among the lower end of biological tissues, with a MFP of about 250 μm (Table 1) [22].

202 The relative role of tissue vs skeleton light scattering in modulating light propagation for coral
203 photosynthesis has been debated [16, 18, 19]. We thus compared tissue and skeletal scattering on the
204 same coral colonies using OCT *in vivo* (Fig. 2, Table 1). A comparison of tissue and skeletal scattering
205 showed that tissue scattering was 8-fold higher compared to skeleton scattering for *P. pini* (Table 1),
206 supporting earlier findings of high tissue vs low skeleton scattering in a comparable faviid coral [16]. In
207 contrast, for the corals *S. pistillata*, *P. cactus* and *H. pilosa*, μ_s was similar between tissue and skeletons
208 (Table 1). These results highlight species-specific variations in the relative role of tissue and skeletal
209 light scattering for coral light management.

210 ***Light scattering characteristics of different tissue types.*** The structural organisation of coral tissues
211 varies within a coral species [31], and given that light scattering in biological tissues is a function of
212 refractive index fluctuations due to different cell types and constituents [22], it is thus to be expected that
213 scattering variability exists within different spatial compartments of the coral tissue. We found that the
214 scattering coefficient of coenosarc tissues ($\mu_s=24.6 \pm 2.8$ SE) was about 3-fold higher compared to polyp
215 tentacle tissue ($\mu_s=8.3 \pm 0.4$ SE) in the coral *Turbinaria reniformis* (one-way ANOVA $F_{1,7}= 26.3$,
216 $p<0.01$, Fig. 3a-e). The lower scattering for the polyp tentacles could be related to a simpler tissue
217 structure as the tentacles do not have a mesoglea and an aboral gastrodermal layer as compared to the

218 coenosarc tissue [39]. The mesoglea is collagen-rich and has been suggested to have an important role
219 in tissue light scattering [15, 16, 40].

220 Close up images of the tissue surface of the coral *Pavona cactus* revealed an interesting tissue
221 surface pattern, alternating between ‘brown crevices’ and ‘white elevations’ (Fig. 2c, 3f). Although we
222 did not characterise *Symbiodinium* distribution via spectral measurements, the brown colour is clearly
223 indicative of a dense aggregation of *Symbiodinium* cells [41]. Cross-sectional OCT scanning showed
224 that ‘white elevations’ are skeletal extrusions, fully covered by living tissue but apparently lacking
225 *Symbiodinium* (Fig. 3g). These extrusions are likely coenosteal spines, which are found in branching
226 corals, including e.g. *Stylophora pistillata* and *Pocillopora damicornis* [42, 43]. Optical analyses showed
227 that the scattering coefficient was about 1.8-fold higher for the brown crevices ($\mu_s=22.42 \pm 1.67$ SE)
228 compared to the white elevations ($\mu_s=12.35 \pm 0.76$ SE) (one-way ANOVA $F_{1,12}= 29.9$, $p<0.01$, Figure
229 3). The enhanced scattering found in the brown crevices compared to the skeletal elevations of the
230 coenosarc tissue of *Pavona cactus* could indicate a link to the aggregation of *Symbiodinium* cells. The
231 scattering coefficient of *Symbiodinium* cells was estimated to $\mu_s=16 \text{ mm}^{-1}$ [20], which is 1.3 fold higher
232 than the average scattering over the white areas (Fig. 3h). The scattering of *Symbiodinium* likely depends
233 on the clade specific cell ultrastructure [44], including e.g. the density of membrane lipids, which are
234 important light scatterers [45, 46]. Although a detailed study on the scattering coefficient of *Symbiodinium*
235 was beyond the scope of the present study, these results hint towards a potential role of light scattering
236 by *Symbiodinium* for coral light management, which warrants future investigations.

237

238 We also studied the light scattering properties of fluorescent host pigments (Fig. 4a-g). The role
239 of fluorescent host pigments (FP) for coral photosynthesis and ecophysiology is disputed, as studies have

240 shown that FP can be either photoprotective (e.g. by absorbing damaging UV radiation and
241 backscattering of incident radiation [47]) or stimulate photosynthesis [48-50] depending on their spectral
242 properties and distribution in the coral tissue. It has also been suggested that such photoprotective and/or
243 photosynthesis stimulating functions depend on the type of FP and its structural aggregation within the
244 tissue [31, 51]. In our study, we found FP aggregations in *Platygyra pini*, where they formed cluster-like
245 ~50-100 μm wide granules also known as chromatophores [31, 49] (Fig. 2a, Fig. 4e,f). These
246 chromatophores are composed of smaller scattering granules, typically about 1 μm in diameter [49]. Our
247 optical analyses showed that tissue areas with such FP aggregates have a low g value of about 0.34 (\pm
248 0.09 SE) indicating a nearly isotropic scattering behaviour of FP granules (Fig. 4g). In contrast, coral
249 tissue with a more diffuse distribution of FP was strongly forward scattering with a g value of about 0.96
250 (\pm 0.008 SE, pooled for *Hydnopora pilosa* and *Echnipora lamellosa*) (Fig. 4a-d,g). Likewise, brown-
251 pigmented tissue in *P. pini* was forward scattering ($g=0.97 \pm 0.004$ SE (Fig. 4g).

252 Previous studies showed that high densities of light scattering FP granules lead to enhanced
253 tissue reflectivity and surface scalar irradiance [11, 51]. Many faviid corals have a dense network of GFP
254 granules, covering >30% of the polyp surface area [31]. Our results suggest that the almost isotropic
255 scattering behaviour of FP granules leads to an effective lateral homogenization of the vertically incident
256 irradiance, which is characterised by enhanced tissue surface scalar irradiance and a more rapid vertical
257 attenuation of irradiance compared to tissue areas with diffuse FP [11]. The g value is diagnostic for the
258 size of the scattering elements and assuming that the light scattering particles are spherical, a lower g
259 value hints towards a smaller size of the scattering elements [52]. Certainly, links between ultrastructure,
260 scattering properties of FP granules, and the photobiology of corals warrant further investigation.

261 ***Implications of coral scattering properties for coral light management.*** The present study provides
262 detailed *in vivo* estimates of the inherent optical properties of live corals which is a key requirement for
263 a better mechanistic understanding of coral optics and thus coral photosynthesis [13]. The quantification
264 of light scattering (μ_s and g) allows for improved light propagation models [16, 20]. Although a detailed
265 simulation was beyond the scope of the present study, one can exemplify basic principles of coral light
266 harvesting in a simplified simulation (see Supplementary Information). We developed a Monte Carlo
267 simulation for two corals (*Stylophora pistillata* and *Hydnophora pilosa*) with different light scattering
268 properties (Table, Fig. S2). Previous studies suggested that the optical properties of the skeleton of *S.*
269 *pistillata* lead to a strong lateral redistribution of incident irradiance which enhances photosynthetic
270 efficiency but also makes this species highly susceptible to coral bleaching [14, 18].

271 Using the *in vivo* optical properties of coral tissue and skeleton (Table 1) we show that incident
272 light is indeed strongly redistributed by light scattering in *Stylophora pistillata*. The high lateral spread
273 of light leads to an 84% absorption of light by the tissue layer (assuming $\mu_a = 3 \text{ mm}^{-1}$, Fig. S2a). In
274 contrast, the light scattering properties of *H. pilosa* lead to a much lower lateral spread of light and an
275 approximate 1.3-fold reduced light absorption by the tissue layer (Fig. S2b). Although the scattering
276 coefficient in *H. pilosa* is higher than in *S. pistillata* (Table 1), the g value in *S. pistillata* is very low
277 (tissue, $g=0.51$ skeleton, $g=0.28$) leading to a more isotropic scattering behaviour and enhanced lateral
278 spread of light (Fig. S2a). Most biological tissues are strongly forward scattering [22] and g values of
279 0.9 have been assumed for corals [16]. However, our study shows that the assumption of strongly
280 forward scattering tissues is not always correct for coral tissues and skeletons (Table 1).

281 It is important to point out, that coral morphology (mm to cm scale) also affects light distribution
282 [18, 53] and first attempts have been made to model light propagation with a 3D architecture [16]. Mesh-

283 based 3D monte Carlo simulations have been developed for brain tissues in optogenetics [54] and can
284 now in theory also be developed using 3D segmented tissue and skeletal architectures from OCT scans
285 (Fig.1) [31] combined with data on scattering properties of intact corals (this study) and skeletons [14,
286 20].

287 In conclusion, the current OCT-based approach allows for a rapid determination of *in vivo* optical
288 properties of corals *in vivo* under defined laboratory conditions. The experimental measurements are fast
289 and application of the theoretical model is straight forward, showcasing the suitability of OCT as a rapid
290 method for optical characterisation. Our approach is currently limited to the extraction of optical
291 properties in the NIR range (930 nm) and ideally this should be improved for characterising light
292 scattering in the visible range (400-700nm) via e.g. visible light OCT [55] or diffuse optical spectroscopy
293 [22].

294 **Conflict of Interest**

295 The authors declare that the research was conducted in the absence of any commercial or financial
296 relationships that could be construed as a potential conflict of interest.

297 **Author Contributions**

298 DW, CFP and MK designed study. DW, JBH and NL performed measurements. SLJ provided analytical
299 tools. CFP provided coral holding and lab facilities, maintained and prepared coral specimens. DW
300 analyzed data with input from SLJ and MK. DW wrote the article with editorial input from all co-authors.

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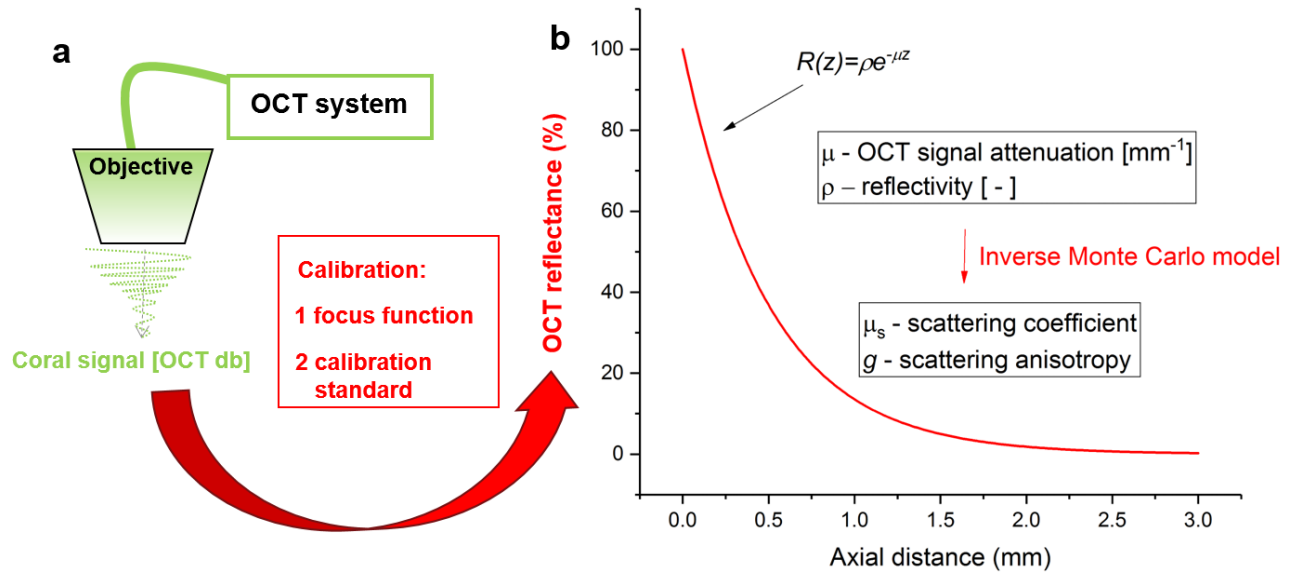
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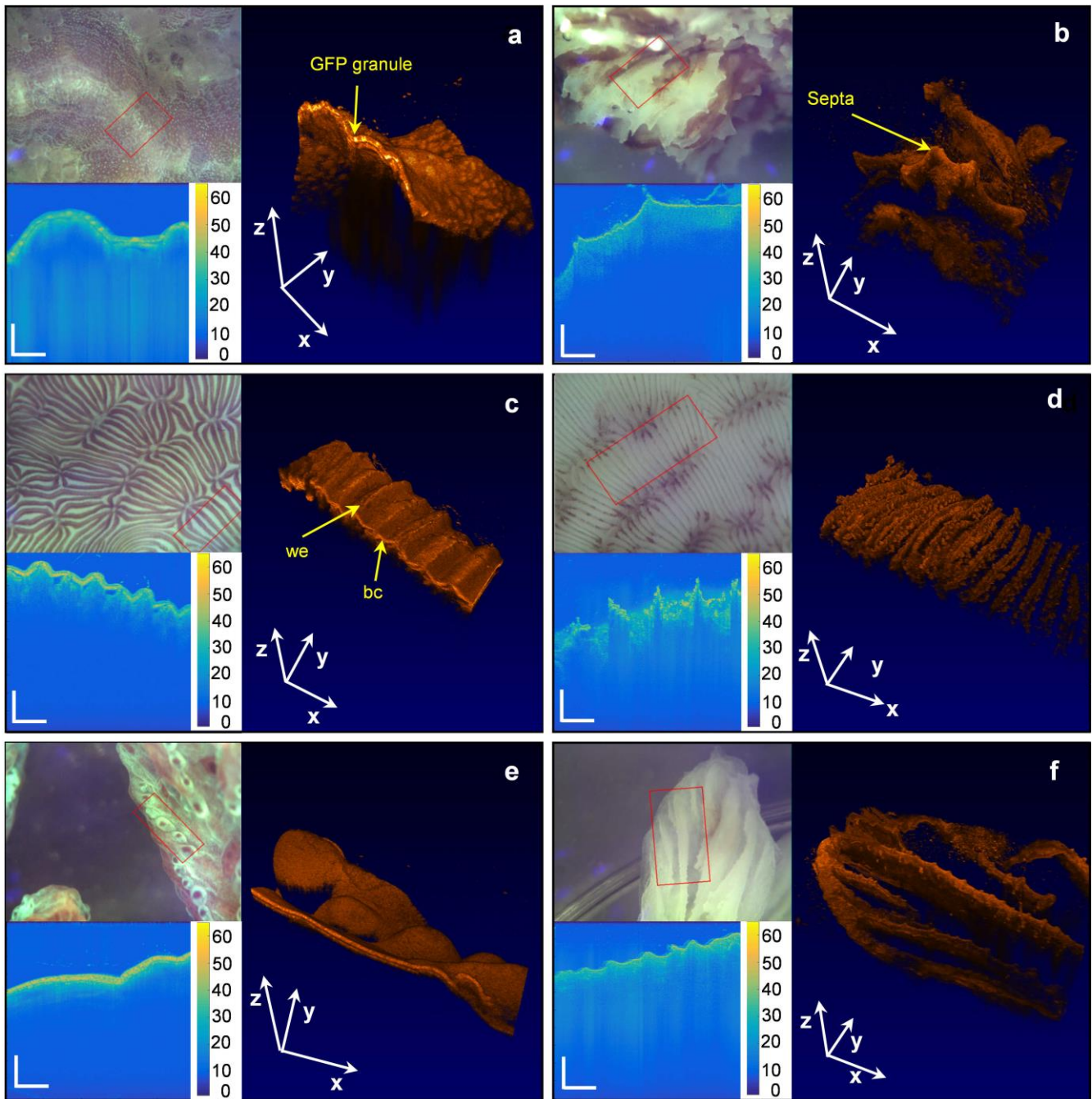
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460 **Figures**



461

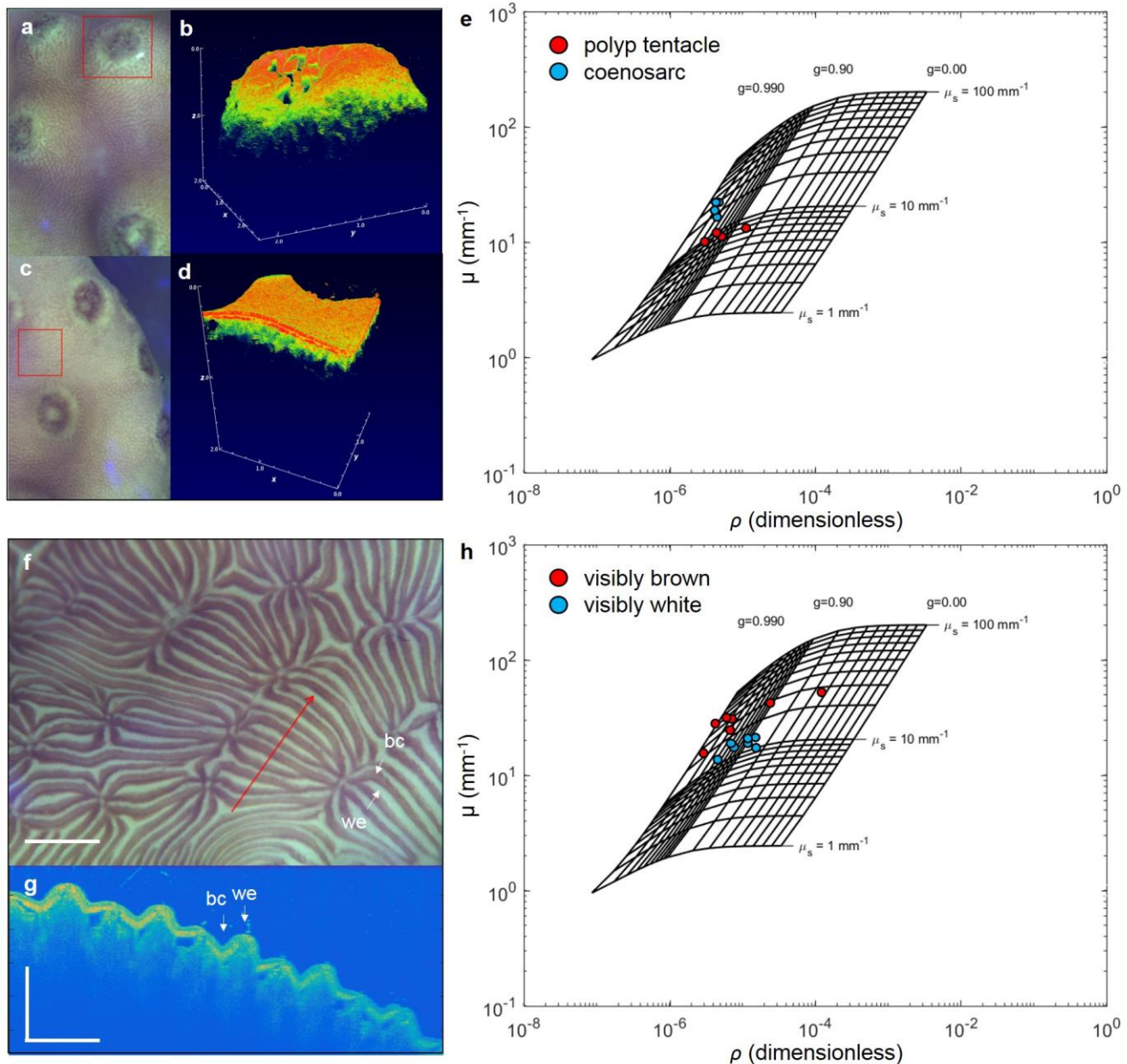
462 **Fig. 1. Experimental and theoretical approach to extract the scattering coefficient μ_s [mm⁻¹] and**
463 **anisotropy of scattering g [dimensionless] for intact living corals.** a) A spectral domain OCT system
464 provided non-invasive low coherent light (930 nm) that imaged corals in vivo without any actinic effect.
465 For each coral, the raw OCT dB signal was calibrated via correcting for the depth-dependent signal
466 attenuation (focus function calibration) and scanning against known reflectance standards to yield the
467 absolute OCT reflectance. b) The vertical coral OCT reflectance attenuation at a given area of interest
468 was then matched to a simple linear exponential function that yielded (1) the reflectivity and (2) the slope
469 of the signal attenuation, which was then matched to gain μ_s and g via an inverse Monte Carlo approach
470 (see methods for details).



471

472 **Fig. 2. Overview of OCT scanning on intact corals (left panel) and bare coral skeletons (right**
473 **panel).** OCT scans are shown for *Platygyra pini* (a, b), *Pavona cactus* (c, d) and *Hydnophora pilosa* (e,
474 f). Close-up photographs were taken with the USB camera of the OCT system. The area within the red
475 square corresponds to the three-dimensional OCT scans (displayed in black-orange false-color coding,
476 which was adjusted for optimised visualisation). The *x-y-z* scale bars represent a distance of 2 mm in

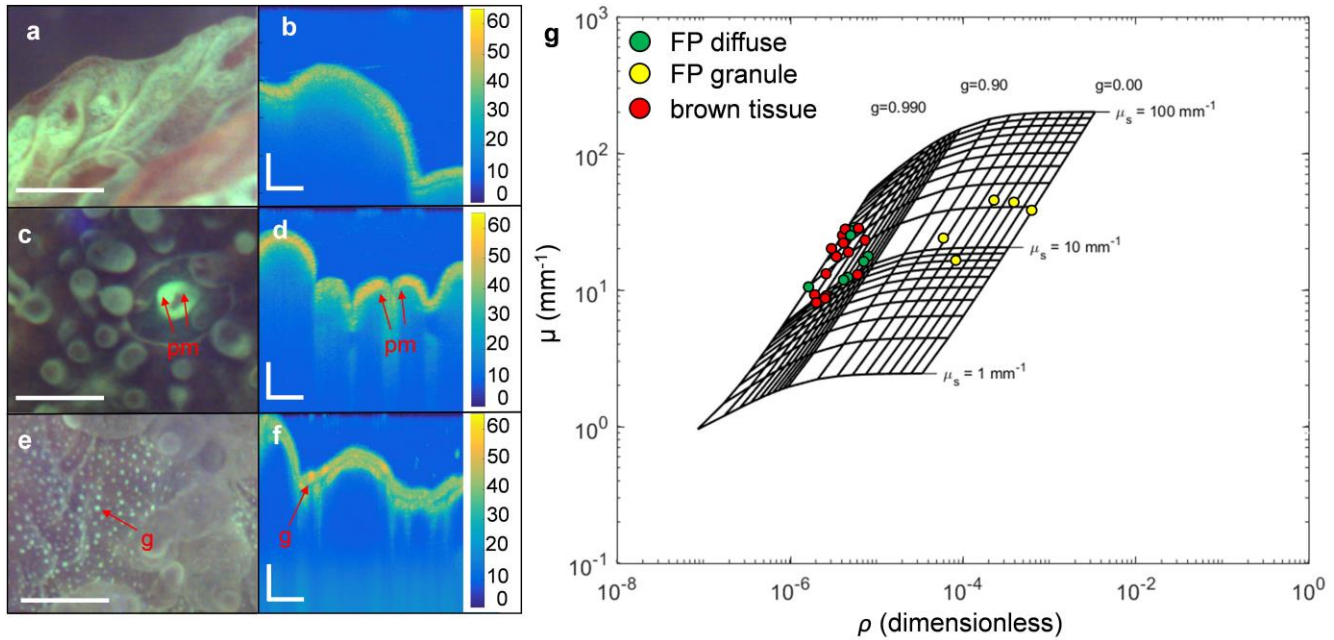
477 each dimension. Exemplary cross-sectional OCT B-scans are shown before calibration with the false-
 478 color legend representing the OCT signal from 0-60 OCT dB. Scale bars in z and x dimensions represent
 479 a distance of 400 μm . we= white elevation, bc= brown crevices.



480

481 **Fig. 3. Characterisation of scattering properties of different coral tissue types.** Close up images (a,c)
 482 and cross sectional OCT B-scans (b,d) of *Turbinaria reniformis* coenosarc and polyp tissues,
 483 respectively (as shown in red square). The optical grid (e) matches the local reflectivity (ρ) to the

484 anisotropy of scattering, g , while the vertical OCT signal attenuation, μ , yields the optical scattering
485 coefficient, μ_s . (see methods for details). Close-up image (f) of the tissue surface of *Pavona cactus*
486 covering brown cervices (bc) and white elevations (we). Scale bar = 2 mm. The red arrow shows the area
487 covered by OCT imaging in panel b. Exemplary OCT cross-sectional B-scan (g). Scale bar in x and z =
488 500 μm . Optical grid comparing white elevations and brown crevices (h).



489

490 **Fig. 4. Light scattering by fluorescent host pigments (FP) in *Hydnophora pilosa* (a,b), *Echinopora***
491 ***lamellosa* (c,d) and *Platygyra pini* (e,f). pm= polyp mouth, g= granule. The optical analyses (g) included**
492 **all three species and show extracted g values for brown tissue areas without visible FP content, tissue**
493 **areas with a diffuse FP distribution, and tissue areas with a clear aggregation of FP into granules. Scale**
494 **bars are 1mm (a, c, e) and 200 μm (b, d, f).**

495

496 **Tables**

497 **Table 1 Optical properties of 8 coral species extracted from calibrated OCT scans at 930 nm.** The
 498 extracted ρ (local reflectivity) and μ (signal attenuation) were used to fit g (anisotropy of scattering)
 499 and μ_s (scattering coefficient). The mean free path ($MFP=1/\mu_s$) and the transport mean free path
 500 ($TMFP=1/\mu_s'$) were calculated. Data are means ($\pm SE$).

Species	Type	ρ []	μ (mm^{-1})	g []	μ_s (mm^{-1})	MFP (mm)	$TMFP$ (mm)	R^2	n
<i>Acropora sp.</i>	tissue	2.10E-06 ($\pm 4.04\text{E-}07$)	10.5 (± 1.1)	0.98 (± 0.003)	14.5 (± 1.44)	0.08 (± 0.01)	5.34 (± 0.82)	0.56 (± 0.03)	10
<i>Echinopora lamelosa</i>	tissue	5.45E-06 ($\pm 7.85\text{E-}07$)	20.9 (± 1.5)	0.97 (± 0.005)	25.2 (± 1.8)	0.04 (± 0.001)	2.1 (± 0.29)	0.79 (± 0.01)	22
<i>Hydnophora pilosa</i>	tissue	4.64E-06 ($\pm 7.93\text{E-}07$)	13 (± 1.2)	0.96 (± 0.007)	12.2 (± 1.09)	0.09 (± 0.01)	2.7 (± 0.44)	0.71 (± 0.04)	12
	skeleton	8.19E-07 ($\pm 1.60\text{E-}07$)	7.7 (± 0.85)	0.99 (± 0.003)	12.1 (± 1.38)	0.09 (± 0.01)	7.34 (± 0.76)	0.55 (± 0.03)	6
<i>Montipora capricornis</i>	tissue	6.72E-05 ($\pm 2.04\text{E-}05$)	16.2 (± 1.6)	0.59 (± 0.09)	8.63 (± 0.83)	0.12 (± 0.01)	0.62 (± 0.31)	0.75 (± 0.03)	8
<i>Pavona cactus</i>	tissue	1.75E-05 ($\pm 9.13\text{E-}06$)	20.5 (± 2.3)	0.91 (± 0.03)	17.4 (± 1.65)	0.06 (± 0.01)	1.53 (± 0.31)	0.78 (± 0.02)	14
	skeleton	2.12E-06 ($\pm 4.96\text{E-}07$)	10.8 (± 1.2)	0.98 (± 0.004)	15.6 (± 1.8)	0.07 (± 0.009)	0.52 (± 0.13)	0.56 (± 0.03)	6
<i>Platygyra pini</i>	tissue	1.42E-4 ($\pm 7.51\text{E-}05$)	26 (± 3.6)	0.69 (± 0.11)	21.3 (± 2.02)	0.05 (± 0.01)	1.37 (± 0.45)	0.79 (± 0.03)	11
	skeleton	1.04E-04 ($\pm 1.96\text{E-}05$)	6.0 (± 0.74)	0.15 (± 0.05)	2.7 (± 0.35)	0.44 (± 0.05)	0.55 (± 0.08)	0.54 (± 0.01)	12
<i>Stylophora pistillata</i>	tissue	3.55E-05 ($\pm 1.74\text{E-}05$)	6.3 (± 0.5)	0.51 (± 0.16)	4 (± 1)	0.33 (± 0.06)	2.13 (± 1.01)	0.81 (± 0.03)	6
	skeleton	7.59E-05 ($\pm 2.59\text{E-}05$)	8.4 (± 1.4)	0.28 (± 0.05)	4.03 (± 0.74)	0.36 (± 0.09)	0.52 (± 0.13)	0.53 (± 0.04)	9
<i>Turbinaria reniformis</i>	tissue	4.69E-06 ($\pm 1.01\text{E-}06$)	15 (± 2)	0.96 (± 0.02)	17.3 (± 3.26)	0.08 (± 0.01)	2.32 (± 0.29)	0.77 (± 0.03)	9