Calcein labelling and electrophysiology: insights on coral tissue permeability and calcification

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The mechanisms behind the transfer of molecules from the surrounding sea water to the site of coral calcification are not well understood, but are critical for understanding how coral reefs are formed. We conducted experiments with the fluorescent dye calcein, which binds to calcium and is incorporated into growing calcium carbonate crystals, to determine the permeability properties of coral cells and tissues to this molecule, and to determine how it is incorporated into the coral skeleton. We also compared rates of calcein incorporation with rates of calcification measured by the alkalinity anomaly technique. Finally, by an electrophysiological approach, we investigated the electrical resistance of coral tissues in order to better understand the role of tissues in ionic permeability. Our results show that (i) calcein passes through coral tissues by a paracellular pathway, (ii) intercellular junctions control and restrict the diffusion of molecules, (iii) intercellular junctions should have pores of a size higher than 13 Å and lower than 20 nm, and (iv) the resistance of the tissues owing to paracellular junctions has a value of $477 \pm 21$ Ohm cm$^2$. We discuss the implication of our results for the transport of calcium involved in the calcification process.

Keywords: coral; biomineralization; membrane permeability; calcium transport; cell junctions; para/transcellular transport

1. INTRODUCTION

The calcification of corals underpins the largest biomineralized structures on the planet: coral reefs. Coral skeletons themselves are widely used to infer past environmental conditions in the Earth’s history. However, despite an abundant literature on coral calcification (for reviews see [1–8]), mechanistic aspects of this process still remain enigmatic, especially the mechanisms behind the transport of calcium at the level of the calcifying cells (i.e. the calicoblastic cells; for reviews see [1,9]).

Indeed, three potential pathways have been proposed: (i) an active transcellular transport of calcium through calicoblastic cells, (ii) a passive paracellular diffusion of calcium or sea water between calicoblastic cells, and (iii) a combination of transcellular and paracellular pathways.

The data in favour of an active transcellular pathway rely on physiological experiments [10–12] that have produced kinetic and pharmacological data, which show that inhibitors of transcellular calcium carriers also inhibit calcification. Moreover, data obtained from molecular approaches show the presence of a calcium channel [13] and a CaATPase [14] in calicoblastic cells that could be responsible for a transcellular transport of calcium. However, these data are not conclusive since many epithelial cells (even in non-calciifying organisms) contain similar carriers and to date no functional analysis has been conducted to establish their role in calcification. Furthermore, pharmacological inhibitors may not be specific to calcium transport for calcification but also to calcium carriers involved in other processes of cell physiology, such as regulation of endocytotic and exocytotic pathways, and can therefore indirectly affect coral calcification.

Arguments in favour of a passive paracellular pathway in corals—in which bulk sea water (external unmodified) arrives at the site of calcification, where its composition can be modified by enzymes—have been presented by several authors [14,15–18]. One important source of data supporting the existence of a paracellular pathway arises from experiments that show cell impermeant molecules such as calcein or alizarine (which are used to stain coral skeletons during calcification) are incorporated into the skeletons [3,15,17–20], suggesting that these molecules follow a paracellular pathway. These observations are puzzling when considered against histological data, where paracellular pathways are usually distinguished from transcellular pathways based on the morpho-functional properties of intercellular epithelial junctions. In corals, based on ultrastructural studies, septate junctions have been described between calicoblastic cells [7,21,22]. These junctions are usually considered in invertebrates as the counterparts of tight junctions in...
vertebrates, which constitute barriers to the paracellular transport of ions [23]. However, in a freshwater mollusc, it has been shown that these junctions are permeable to calcium (with at least 50 per cent of transport being paracellular [24]) and thus one can question their role in coral tissue permeability.

The aim of the current study was to investigate the variable lines of evidence obtained on the pathway of ions/molecules with respect to their implication for coral calcification. We thus conducted experiments with calcein to determine the permeability properties of coral cells and tissues to the molecule and to determine how it is incorporated into coral skeleton. We also compared rates of calcein incorporation with rates of calcification measured by the alkalinity anomaly technique [30] and half of the samples were used for calcification measurements by the technique. Finally, by an electrophysiological approach, we looked at the electrical resistance of coral tissues in order to better understand the role of tissues in ionic permeability.

2. MATERIAL AND METHODS

(a) Biological material

The majority of the experiments were performed on the scleractinian coral *Stylophora pistillata*, and some experiments were performed on the Mediterranean sea anemone *Anemonia viridis*; both were maintained in the laboratory in controlled conditions of culture (see the electronic supplementary material). Coral apaxes were cut into small fragments to obtain microcolonies cultivated on nylon nets [25] or grown on glass coverslips or glass slides [22,25–28]. For experiments, all incubations of living specimen were performed under the same environmental conditions of light intensity, pH and temperature as in culture conditions.

(b) Calcein experiments

Calcein is a fluorescent label that binds to calcium and is incorporated into growing calcium carbonate crystals [29]. Calcein was purchased from Sigma-Aldrich. A concentrated solution containing 2 g l⁻¹ calcein was prepared in distilled water and buffered to pH 6 with sodium bicarbonate to enhance solubility [29]. This concentrate was then diluted in filtered sea water (fSW; pore size filter 0.2 μm) buffered on glass coverslips or glass slides [22,25–28]. For experiments, all incubations of living specimen were performed under the same environmental conditions of light intensity, pH and temperature as in culture conditions.

(c) Preparation of samples for calcein experiments

(i) Qualitative measurements of calcein incorporated into calcium carbonate

To investigate the incorporation of calcein into skeletons or inorganic calcium carbonate crystals, samples were incubated for 20 min with fSW-calcein 20 μM. Observations were performed with a fluorescent microscope and spectra were obtained with a spectrophotometer (see below).

(ii) Quantitative measurements of calcein incorporated into calcium carbonate

To measure both calcein incorporation into the skeleton and calcification, microcolonies prepared as described above were incubated for 2, 4, 6 and 24 h in fSW-calcein 20 μM. Half of the samples were used for calcification measurements by the alkalinity anomaly technique [30] and half of the samples were used to determine calcein incorporated into the skeleton. For calcein incorporation, tissues were removed by the treatment with NaOCl 10 per cent in distilled water for 30 min and then skeletons were rinsed three times in distilled water and dried at room temperature. Skeletons were then dissolved with HCl 2N and the solution was neutralized with NaOH to pH 8.2.

A standard curve of calcein at concentrations varying from 0.05 to 20 μM in fSW was prepared in the same conditions as the samples (HCl treatment and NaOH neutralization, pH 8.2) and was used as a calibration curve to determine the quantity of calcein incorporated into the coral skeletons. Measurements of calcein were obtained with the confocal system (see below), by spectral analysis with excitation at 488 nm. The correlation coefficient of the calibration curve of calcein was 0.9914.

(iii) Permeability of tissues to calcein

To determine if calcein passes through a paracellular or a transcellular pathway, live microcolonies grown on glass coverslips were incubated for 20 min in 5 ml of fSW-containing calcein in fSW (concentration specified for each experiment). Observations were performed with the confocal microscope (see below).

More details on sample preparation are given in the electronic supplementary material.

(d) Fluorescent beads experiments

Red fluorescent Flospheres (carboxylate-modified microspherical beads of 20 nm, 200 nm and 2 μm were purchased from Molecular Probes (ref. F8887).

To determine if fluorescent beads were incorporated into the skeleton, microcolonies were incubated for 24 h (12 L:12 D cycle) in 50 ml fSW-containing 2 per cent fluorescent beads (each size independently) under mixing with a magnetic stirrer. Samples were then rinsed for 10 min in 25 ml H₂O and tissues were removed with a Waterpik Ultra Water Flosser. Skeletons were rinsed in ultrapure water and dried at room temperature.

To determine if fluorescent beads could pass through the cell layers, microcolonies grown on glass coverslips were incubated for 20 min in 5 ml fSW-containing 2 per cent fluorescent beads (each size independently) under mixing with a magnetic stirrer. Observations were performed with the confocal microscope (see below).

(e) Observations of samples (calcein and fluorescent beads)

As specified for each experiment, observations were made either with a fluorescent microscope (Z16APO, Leica Microsystems) or with an inverted confocal laser-scanning microscope (TCS SP5, DMI 6000 CS, Leica Microsystems).

Quantitative measurements of calcein were obtained with the confocal system by spectral analysis with excitation at 488 nm.

For calcein and fluorescent beads observed with the Leica Z16APO microscope or the Leica DMI 6000 CS microscope, samples were excited, respectively, with blue and green light, and emission fluorescence was detected using a JVC 3CCD digital camera.

For fluorescence spectra of calcein in confocal microscopy, measurements were made with a spectrophotometer coupled to the imaging system with 488 nm laser excitation.

For confocal images of calcein, excitation wavelength was 488 nm and emission was 515 ± 15 nm. For confocal images

of red fluorescent beads, excitation wavelength was 543 nm and emission was 605 ± 15 nm.

(f) Electrophysiological experiments
(i) Preparation of samples
Microcolonies of the coral S. pistillata were cultured over perforated glass slides with a hole of 2.5 mm radius and 0.2 cm² surface area, on which the apex of the coral was placed (for details of the experimental set-up, see electronic supplementary material, figure S1a–d and material and methods). The sea anemone, A. viridis, was also used since this anthozoan possesses long tentacles from which oral tissues are easy to obtain. Briefly, one piece of tentacle was cut longitudinally and placed between the two Ussing half-chambers as described previously by Bénazet-Tambutte et al. [31]. Resistance was then calculated from the values of intensity and voltage (for details see electronic supplementary material, material and methods).

3. RESULTS
(a) Calcein, calcium carbonate and fluorescence
To determine the potential autofluorescence of the samples, we observed a bare coral skeleton under excitation with blue light. As can be seen in figure 1a, the control bare coral skeleton (without calcein) shows a faint yellow–green autofluorescence. This signal is weaker than the bright green fluorescence owing to the incorporation of calcein (i) in calcium carbonate crystals precipitated in vitro (figure 1b) and (ii) in the skeleton of live microcolonies during in vivo experiments (figure 1c). The spectral analysis showed that calcein incorporated into coral skeleton of live microcolonies gives an emission spectrum (after excitation at 488 nm) similar to calcein in sea water, whereas the control bare coral (without calcein) gives negligible emission under excitation at 488 nm (figure 1d). Spectra obtained after dissolution of skeletons from live microcolonies incubated with calcein had identical spectra to those obtained for calcein in sea water treated with HCl and NaOH (final pH 8.2; data not shown).

(b) Calcein incorporation into coral skeleton
The following observations were obtained with a macroscope but results were always checked by spectral analysis in order to discriminate between autofluorescence and fluorescence owing to calcein (as described above). Figure 2a,b,c respectively, show the fluorescence of (i) the skeleton from a live microcolony (microcolony = skeleton + tissue), (ii) a bare skeleton (= skeleton without tissues) and (iii) the skeleton from a dead microcolony (dead microcolony = microcolony killed with NaCN, 1 mM, 2 h, = skeleton with dead tissues), all after incubation in calcein. Except for the bare skeleton, the incubations were performed in the presence of tissues. Further treatment of the skeletons with NaOCl revealed that calcein labelling is removed (labile) in bare skeletons and skeletons from dead microcolonies, indicating that it is adhering superficially to the CaCO₃ (results not shown). By contrast, in live microcolonies, calcein is permanently incorporated into the skeletons as a result of calcification. In live corals, calcein is incorporated into the newly forming crystals in areas such as septa and columns of corallites (figure 2d), individual granules of calcium carbonate growing on glass slides (figure 2e) and forming spines (figure 2f). Complementary experiments to show that the green granules in figure 2e are due to the incorporation of calcein in calcium carbonate granules
and not to green fluorescent proteins (GFP) are presented as the electronic supplementary material, figure S2a–f. All these results show that, while calcein may superficially bind to a bare skeleton or a skeleton from a dead microcolony, it is permanently incorporated (resistant to NaOCl treatment) into calcifying zones of live growing corals.

(c) Stoichiometry of calcein incorporation into the skeleton
The amount of calcein incorporated into the skeletons of live microcolonies was estimated using the calibration curve of calcein as described in §2. The results were obtained as a function of time (2, 4, 6 and 24 h). The rate of calcein incorporation has a value of 0.08 nmol calcein g⁻¹ dry skeleton h⁻¹. In parallel, the rate of calcification was measured by the alkalinity anomaly technique and has a value of 3302 ± 173 nmol CaCO₃ g⁻¹ dry skeleton h⁻¹. The equation of the slopes fitted with linear regression was $y = 3344.8x$ for alkalinity and $y = 0.0771x$ for calcein. Equality of the slopes was tested on normalized values (normalization by the mean, n = 5 samples) using GraphPad Prism (v. 5.0), which confirmed that they were not statistically different (ANCOVA, $p = 0.40, F = 0.82$). The results are presented in figure 3 as the rate of calcein incorporation versus the rate of calcification. In our experimental conditions, the rate of calcein incorporation is far lower than the rate of calcium incorporation (ratio of $2 \times 10^{-5}$).

(d) Paracellular versus transcellular pathway of calcein
We investigated the permeability of coral tissues to calcein (which is known to be impermeant to cells owing to its hydrophilic properties) by observing whole tissues of microcolonies grown on glass coverslips (for histology, see [28]) and incubated with calcein (figure 4). Using an inverted confocal microscope, we could focus on specific optical sections obtained by z-stack analysis through the preparation from tissues and growing crystals in contact with the coverslip to the upper tissues. As can be seen in figure 4a, calcein is present in the outside medium but also between oral ectodermal cells. Control experiments without calcein (figure 4a) allow us to determine that intracellular small green fluorescent dots are due to autofluorescent proteins. The same paracellular distribution of calcein was also observed in the endodermal tissues (figure 4d). We could also observe the tissue directly in contact with the coverslip and surrounding the growing crystals (i.e. the calcicoblastic ectoderm). In control experiments without calcein, there was no fluorescence owing to autofluorescent proteins in the calcicoblastic ectoderm. We had to use a higher concentration of calcein (160 μM) because the calcicoblastic cells are very thin and labelling was difficult to observe. As can
be seen in figure 4e, calcein is present between the calcicoblastic cells, and the two green high fluorescent zones (asterisks) are due to calcein incorporated into forming crystals (crystals arranged as fasciculi can be seen with bright field transmitted light; figure 4f).

**Figure 4.** Characterization of the paracellular pathway. (a–f) Experiments with microcolonies grown on glass coverslips. Except for (c), observations were made with the inverted microscope of a confocal and the z-stack tool was used to position the observation. (c) Representative microcolony grown on a glass coverslip and observed with bright field light, observation with a macroscope. (a) Control microcolony without any incubation, magnification of the upper zone of (c). Small green autofluorescent dots are due to autofluorescent proteins. (b) Microcolony incubated for 20 min in the presence of calcein 20 μM, magnification of the upper zone of (c). Small green fluorescent dots are due to autofluorescent proteins whereas paracellular green fluorescence in oral ectoderm is due to calcein. (d) Microcolony incubated for 20 min in the presence of calcein 20 μM, magnification of the right zone of (c). Paracellular green fluorescence of endodermal cells is due to calcein; the image is merged with a bright field transmitted light image in which zooxanthellae appear as black spheres (the optical section is 17 μm above the coverslip, taken from a z-stack with a 0.3 μm step). (e) Microcolony incubated for 20 min in the presence of calcein 160 μM, magnification of the lower zone of (c). Paracellular green fluorescence in calcicoblastic ectoderm surrounding the two fluorescent crystals (asterisks) is due to calcein (the optical section is 1 μm above the coverslip). (f) Bright field transmitted light allows confirming that the two large fluorescent zones in (e) are due to calcein incorporated into calcium carbonate crystals (arranged as fasciculi). (g) Microcolony incubated for 20 min with 200 nm red fluorescent beads, magnification of the left zone of (c). The image is a merging of bright field transmitted light and red fluorescence. Crystals can be distinguished in the growing front and are surrounded by tissues whereas red fluorescent beads remain outside the tissues.

**Permeability of the tissues to fluorescent beads**

To estimate the size of the paracellular pathway, we incubated microcolonies with fluorescent beads of different sizes (20 nm, 200 nm or 2 μm), removed the tissues and observed the skeleton. We did not observe any labelling of the skeleton whatever the size of the fluorescent beads (observations performed with a fluorescent macroscope and microscope; results not shown since no fluorescence was observed). We confirmed this result by incubating the microcolonies grown on coverslips with the fluorescent beads of different sizes. In this case, we looked at the tissues with the confocal microscope (inverted microscope) using both bright field transmitted light and fluorescence, and the z-stack tool. As can be seen in figure 4g for the 200 nm diameter beads, regardless of their size, the red fluorescent beads remained at the periphery of the tissues and were never found either in the tissues or in the new growing calcium carbonate crystals, thus confirming that the tissues are not permeable to molecules of 20 nm diameter or more.
Determination of tissues and skeleton resistance

Voltage-clamp experiments in Ussing chambers were performed to determine the electrical resistance of the tissues of *S. pistillata*. We found a value of $1483 \pm 189 \text{ Ohm cm}^2$ for the resistance of whole microcolonies (tissues covering the skeleton). We then removed the tissues and found a value of $1006 \pm 175 \text{ Ohm cm}^2$ for the resistance of the bare skeleton. By subtracting the resistance of the bare skeleton from the resistance of the whole microcolony, we obtained a value of $477 \pm 21 \text{ Ohm cm}^2$ for the resistance of the tissues (oral and aboral tissues). Since it was impossible to separate the oral tissue from the aboral tissue in *S. pistillata*, we measured the resistance of the oral tissue of the sea anemone *A. viridis* and obtained a value of $15 \pm 3 \text{ Ohm cm}^2$.

We then performed a hyperosmotic shock, which is usually used to determine if junctions are involved in the paracellular resistance of tissues [32]. When the hyperosmotic shock was performed in the hemi-chamber facing the tissues (apex) of a microcolony, the resistance decreased as a function of time (electronic supplementary material, figure S3). Since there was no decrease in the resistance of a bare skeleton under hyperosmotic shock (results not shown), we can confirm that we really measured the resistance of the tissues and we can conclude that, under hyperosmotic shock, the tissues became less resistant to electric current.

4. DISCUSSION

(a) Calcine incorporation and coral calcification rate

In corals and in other calcifying organisms, calcine is widely used to indicate skeletal growth. Indeed, organisms exposed to this fluorochrome incorporate it into growing calcified structures in the form of an internal growth mark (observable under excitation with fluorescent blue light) that can subsequently be used to estimate growth from time of exposure. We first compared the spectrum of the fluorescent signal obtained from corals labelled with calcine with the autofluorescence signal of the calcium carbonate of coral skeletons. Clearly, the spectra were different (figure 1). Then, we showed that calcine is incorporated (i) into calcium carbonate crystals precipitated inorganically and (ii) in newly formed crystals present in calcifying zones of coral skeleton, such as spines, septa or columnella. We observed that the labelling of bare skeletons or dead microcolonies was labile (after treatment with NaOCl), whereas it was stable in similarly treated skeletons in live microcolonies. In parallel to calcine incorporation, we measured the calcification rate by the alkalinity method of mineralization of calcium phosphate [34].

(b) Insight on transepithelial transport from experiments with fluorescent molecules

Evidence supporting the transport of bulk sea water to the site of calcification includes observations that molecules such as calcine or alizarin are rapidly incorporated from the surrounding sea water into coral skeleton. However, the pathway of these molecules through the tissues has never been characterized in corals nor in other calcifying organisms.

Our data allow us to conclude that the hydrophilic molecule calcine is transported via a paracellular pathway at the level of the coral tissue layers, including the calicoblastic ectoderm. Indeed, we have shown (i) that calcine is incorporated into the skeleton, indicating that it crosses the four cell layers (unless it is coming from the sea water present in the coelenteron, and then it crosses only two layers); and (ii) that calcine cannot enter into cells, suggesting that it has to pass through a paracellular pathway through the four (or two) cell layers. Using confocal microscopy, we visualized the paracellular pathway with images of calcine between cells of the oral ectoderm and calicoblastic ectoderm (and observed that this was also the case for endoderm). We have shown that fluorescent beads of different sizes were not incorporated into the skeleton and that they were not present between cells. This finding supports the idea that (i) calcine passes through intercellular spaces, such as described in different coral species [7,21,22], and (ii) there are no holes in the tissue through which molecules can pass (holes in the tissues have only been evidenced by microscopy observations in the oral tissues [35,36], never in the calicoblastic ectoderm).

In vertebrates, the gatekeeper of the paracellular pathway is the tight junction, which is an apically located cell–cell interaction of epithelial cells. This tight junction allows the selective pathway of ions while restricting the movement of large molecules. Permeability studies using membrane impermeant tracers suggest that the tight junction has pores of approximately 6–7 Å diameter [37]. In the case of corals, based on ultrastructural studies, the septate junction has been considered the counterpart of the mammalian tight junction and has been localized in the calicoblastic cell layer [7,21,22] (electronic supplementary material, figure S4). Since calcine passes through a paracellular pathway, it has to diffuse through these junctions. We can thus conclude that the septate junctions should possess pores of a larger size than the tight junction (at least twice the size of the molecular radius of calcine, i.e. 13 Å [38]). However, since fluorescent beads of 20 nm do not pass through the epithelial layers, we can conclude that the size of the septate junctions is less than 20 nm. Therefore, combined with the results on calcine, we can conclude that, in corals, the septate junctions present in the paracellular pathway should possess pores of more than 13 Å and less than 20 nm.

(c) Insight on epithelial permeability from electrophysiology experiments

Transepithelial transport of ions/molecules by a paracellular pathway is dependent not only on the size but also on the charge of ions/molecules [39]. The technique of electrophysiology with Ussing chambers has been used for many years to estimate the ‘tightness’ versus the ‘leakiness’ of the epithelial layers. Indeed, voltage clamp experiments indicate whether an epithelial layer forms a low- or high-resistance barrier to ion diffusion and give insight into the importance of the paracellular versus the...
transcellular pathway. In leaky epithelia, resistance values are classically between 6 and 100 Ohm cm$^2$, whereas in tight epithelia, these values range from 500 to 70 000 Ohm cm$^2$ [40,41]. In the present study, electrophysiology experiments show that the tissues have a resistance of 477 ± 42 Ohm cm$^2$. This value could be considered ‘intermediate’, but it is located closer to the lower range of values for mammalian tight epithelia (for comparative values, see electronic supplementary material, table S1). Moreover, this resistance is very high when compared with the resistance of the oral tissue of sea anemones (present study and [31]). Electrophysiology experiments cannot be done on the coral oral tissue owing to technical limitations, but structural similarities between coral and anemone oral tissues [42] suggest that results should be comparable, and if this was the case the higher resistance of the tissues of S. pistillata would be due to the paracellular junctions of the aboral tissue. In epithelia, permeability due to the paracellular pathway is known to increase under hyperosmotic conditions with non-electrolyte solutions by opening junctions [32,43]. In this study, we have observed that a hyperosmotic shock leads to a decrease in microcolony resistance, and thus an increase in tissue permeability. In mammals, electronic microscopic observations show changes in the geometry of junctional complexes compatible with an increase in permeability during hyperosmotic shock [43] causing the opening of septate junctions [44]. The effect of hyperosmotic shock by decreasing coral tissue resistance supports the interpretation that in normal conditions (without osmotic shock), paracellular junctions control and restrict the diffusion of molecules, and thus play a key role in determining the permeability properties of the epithelial layers.

(d) Epithelial permeability and ion transport in corals and other organisms

From the results obtained in the present study on the permeability of tissues to calcein, it is tempting to widen the discussion to ions involved in the calcification process, such as calcium. A direct comparison between calcium incorporation and calcein incorporation in the coral skeleton is not possible for the moment since there is no data in the literature for how calcein is incorporated into calcium carbonate. It is, however, possible to consider our results in context with the existing literature on calcium transport, both in corals and other systems. Studies aimed at determining the mechanisms of calcium transport across coral tissues have shown that, at the level of the oral tissues, calcium is transported either by an active transcellular pathway [21,45] or by a passive paracellular pathway [12,31]. Moreover, at the level of the calicoblastic ectoderm, data suggest that the transport of calcium for calcification is active and transcellular [10–14], and it has been shown that there is an increasing intracellular gradient of calcium from the oral cells facing external sea water to the calicoblastic layer of the aboral tissue [46]. Since the size of calcium ions (atomic radius of calcium is 1.8 Å) is far lower than the size of calcein (molecular radius of calcein is 6.5 Å), it is not possible to rule out the possibility that in addition to the transcellular pathway mentioned above, there is also a paracellular pathway of calcium. However, since the paracellular pathway also depends on the charge of ions/molecules (calcium is positively charged and calcein is negatively charged at pH 8.2), the relative contribution of the paracellular and transcellular transport mechanisms to the overall calcium transport remains unknown.

5. CONCLUSION

The results of the present study show that, in corals, calcein enters the calcification environment via a paracellular pathway with paracellular junctions controlling and restricting the diffusion of molecules. From these data, we propose a schematic of the transfer of calcein and the possible transfer of ions at the level of calicoblastic cells (figure 5). From our
results, we draw attention to some crucial points that need to be taken into account: (i) the role of paracellular junctions in ions/molecules transport; (ii) the significance of the value of the epithelial resistance owing to paracellular junctions; and (iii) the usefulness of studying molecules such as calcein for characterizing epithelial properties (but also the necessity of precaution when extrapolating results for other ions such as calcium). Whereas our results suggest at least semi-open ion transport for characterizing epithelial properties (but also the necessity of studying molecules such as calcein for improving the manuscript. This study was conducted as part of the Centre Scientifique de Monaco Research Programme, funded by the Government of the Principality of Monaco.

REFERENCES


