One-way calcium spill-over during signal transduction in *Paramecium* cells: from the cell cortex into cilia, but not in the reverse direction

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Abstract

We asked to what extent Ca²⁺ signals in two different domains of *Paramecium* cells remain separated during different stimulations. Wild-type (7S) and pawn cells (strain d4-500r, without ciliary voltage-dependent Ca²⁺-channels) were stimulated for trichocyst exocytosis within 80 ms by quenched-flow preparation and analysed by energy-dispersive X-ray microanalysis (EDX), paralleled by fast confocal fluorochrome analysis. We also analysed depolarisation-dependent calcium signalling during ciliary beat reversal, also by EDX, after 80-ms stimulation in the quenched-flow mode. EDX and fluorochrome analysis enable to register total and free intracellular calcium concentrations, [Ca], and [Ca²⁺], respectively. After exocytosis stimulation we find by both methods that the calcium signal sweeps into the basis of cilia, not only in 7S but also in pawn cells which then also perform ciliary reversal. After depolarisation we see an increase of [Ca] along cilia selectively in 7S, but not in pawn cells. Opposite to exocytosis stimulation, during depolarisation no calcium spill-over into the nearby cytosol and no exocytosis occurs. In sum, we conclude that cilia must contain a very potent Ca²⁺ buffering system and that ciliary reversal induction, much more than exocytosis stimulation, involves strict microdomain regulation of Ca²⁺ signals.

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Keywords: Ca²⁺; Calcium; Cilia; EDX; *Paramecium*; X-ray microanalysis

1. Introduction

Ca²⁺ is a pivotal second messenger in numerous cell functions [1–5], including stimulated secretory activity [2] and ciliary beat regulation [6–9]. This also holds true for ciliated protozoa [10–14]. In ciliates dense-core exocytotic vesicles and cilia exist side-by-side in regular intervals [15]. In *Paramecium*, the distance of these organelles is ~1 μm, thus aggravating differential signal transduction [16]. In *Paramecium*, depolarisation-induced activation of ciliary Ca²⁺-channels causes ciliary beat reversal and backward swimming, respectively [10,11,13,17–20], with no exocytotic response [21]. These channels are irrelevant for exocytosis [22]. With permeabilised *Paramecium* cells, ciliary reversal is induced by a Ca²⁺ concentration of 10⁻⁶ M [23]. In contrast, trichocyst exocytosis requires ~5 × 10⁻⁶ M [24] and relies on two different Ca²⁺ sources, (i) Ca²⁺ from cortical stores (“alveolar sacs”) and (ii) from superimposed store-operated Ca²⁺ influx (SOC) via the non-ciliary cell membrane [21,25–27]. A *Paramecium* cell is able to selectively activate either one of the two phenomena, evidently by microdomain Ca²⁺ regulation.

The differential activation of Ca²⁺-mediated processes is difficult to analyse with fluorochromes. Particularly, with cilia this is problematic [8,28,29]. Difficulties in determining precise free Ca²⁺ concentrations, [Ca²⁺], within cilia and flagella have several reasons, not only the small size, but also unknown diffusion effects [30] after passing Ca²⁺-channels whose distribution along the cilium is unknown (see “Section 4”), as is that of a still increasing number of Ca²⁺-activated and Ca²⁺-binding proteins (CaBPs) [31–35], not to speak of practical problems with fluorochrome analyses in these slender and highly motile...
structures. Alternative methods, like intracellular application of Ca<sup>2+</sup> chelators of different K<sub>0</sub> or local photoactivation of caged Ca<sup>2+</sup>, as it is widely used in exocytosis research [36], proved useful merely on a rather coarse scale, at best to compare proximal and distal regions of cilia in Paramecium [37].

Recently, we have established with Paramecium cells energy-dispersive X-ray microanalysis (EDX) combined with scanning transmission electron microscopy (STEM) to determine local values of total calcium concentrations, [Ca] i.e. bound calcium and free Ca<sup>2+</sup> [38,39]. The preparative method we apply is quenched-flow/freeze freezing [40] combined with freeze-substitution at 193 K in presence of fluoride as a calcium precipitating ion of very low solubility [38,41], based on previous work by Poenie and Epel [42]. The quenched-flow/cryofixation method we developed allows stimulation times in the sub-second range. For comparison, synchronous trichocyst exocytosis occurs within 80 ms [40], whereas one ciliary beat requires ∼50 ms (see “Section 3”). While we register considerable changes in Ca K<sub>/H9251</sub> EDX signals upon exocytosis stimulation [38,39], we find no evidence of any artificial redistribution during mock stimulation. Exocytosis stimulation involves depletion of cortical Ca stores (alveolar sacs) superimposed by a Ca<sup>2+</sup> influx during exocytosis stimulation, i.e. a store-operated Ca<sup>2+</sup> influx, SOC [39]—a mechanism basically different from that occurring in cilia. We use 500 nm sections and operate the STEM at a spot size of 63 nm (bottom 72 nm [38]) for spot measurements of Ca K<sub>/H9251</sub> signals. Cilia were analysed along their shaft and signals were recorded also from other sites of the cell as indicated in figures and tables.

In the present paper, we analyse calcium in cilia by measuring both, [Ca<sup>2+</sup>]i by fluorochromes at the light microscope (LM) and [Ca] i.e. by EDX at the electron microscope (EM) level. While at LM resolution [Ca<sup>2+</sup>]i is difficult to determine in cilia, [Ca] values determined at the EM level after stimulation allow one to clearly trace calcium fluxes within fine subcellular details and eventual spill-over into nearby domains. This appears particularly promising for cilia because the arrangement of voltage-dependent Ca<sup>2+</sup>-channels is not known.

Our EDX analysis with wild-type (7S) cells reveals massive calcium fluxes into cilia, not only during depolarisation, but also during exocytosis stimulation. We also observe Ca<sup>2+</sup> spill-over in the “tightest” pawn cells (devoid of the voltage-dependent Ca<sup>2+</sup>-channels in cilia), strain d4-500r [43], into the basis in cilia, when exocytosis is stimulated. This causes ciliary reversal which these cells do not show under depolarisation conditions. We see no spill-over in the opposite direction, i.e. from the cilia into the cell soma when 7S cells are depolarised. Possible explanations of this difference may be, for instance, not only different Ca<sup>2+</sup>-sensitivity, but also the magnitude of the respective calcium fluxes and signals, as well as different Ca<sup>2+</sup>-buffering capacity. Therefore, it was useful to monitor real calcium distributions by EDX.

As to possible biological implications in Paramecium, ciliary reversal rapidly ensues exocytosis during defence against a predator attack. Both these mechanisms acting in concert may explain their enormous efficiency of this defence mechanism [44].

2. Materials and methods

2.1. Cell cultures and stimulation for EDX

Cells, wild-type strain 7S and pawn strain d4-500r [43], were cultivated as described [24] and prepared by quenched-flow/freeze-substitution. Briefly, the methodology developed in our lab was used for quenched-flow [40] and for freeze-substitution under conditions allowing to retain Ca<sup>2+</sup> [38,41]. Depolarisation was performed in media with [Ca<sup>2+</sup>]c = 1.0 mM by rapid mixing in the quenched-flow apparatus with KCl resulting in [K<sup>+</sup>]c = 20 mM. To analyse the effects of synchronous trichocyst exocytosis, cells were exposed to 2 μM aminoethylidextran (AED), routinely at [Ca<sup>2+</sup>]c = 0.5 mM [39].

2.2. EDX

This method was applied as described by Hardt and Plattner [38,39]. Briefly, semithin sections of 0.5 μm were prepared and analysed in a LEO 912 Omega EM operated in the scanning transmission (STEM) mode at 80 kV, −10 μA beam current, using a spot size of 63 nm (bottom 72 nm [38]) for spot measurements of Ca K<sub>/H9251</sub> signals. Cilia were analysed along their shaft and signals were recorded also from other sites of the cell as indicated in figures and tables.

2.3. Fluorochrome measurements

Cells were loaded with Fluo-3 and the f<sub>0</sub> signal ratio during stimulation was analysed with a confocal laser scanning microscope (CLSM) type Tracor/Noran Odyssey, equipped with a fast opto-acoustic deflection system, as described previously [24–26]. During stimulation, signals were recorded close to the site of stimulus application, not only in the cell cortex, but also in an extracellular area adjacent to the cell surface, i.e. in the layer containing beating cilia. The fields to be analysed were framed (∼5 μm wide parallel to the cell surface) and the signals recorded were processed as described [24–26].

3. Results

Table 1 summarises the effects of depolarisation in 7S and d4-500r cells. The ability and inability of 7S and d4-500r cells, respectively, was routinely tested immediately before quenched-flow preparation for subsequent EDX analysis. Under the depolarisation conditions used throughout this
Table 1

<table>
<thead>
<tr>
<th>Reaction of 7S and d4-500r cells to depolarisation conditions and to exocytosis stimulation, respectively a</th>
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<tbody>
<tr>
<td>Ciliary reversal (depolarisation)</td>
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<tr>
<td>----------------------------------</td>
</tr>
<tr>
<td>7S</td>
</tr>
<tr>
<td>20 mM [K⁺] → reversal</td>
</tr>
<tr>
<td>AED → exocytosis</td>
</tr>
<tr>
<td></td>
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a |Ca²⁺| = 1 and 10 mM in 7S and d4-500r cells, respectively. Scaling: “+” to “+++” indicate reactions from clearly positive to maximal intensity (and with shorter duration, as indicated); “−” indicates no reaction.

b Different scaling reflects different intensity and duration of ciliary reversal, i.e. about <8 s and about <5 s in 7S and d4-500r cells, respectively.

paper, only 7S, but not d4-500r cells, responded by ciliary reversal. However, both strains exhibited ciliary reversal, though with different intensity, during synchronous exocytosis induction by AED. In a series of microkinematographic images, the duration of a ciliary beat cycle was determined as 49 ms (~20 Hz, data not shown).

3.1. Fluorochrome analysis of [Ca²⁺] signals in cilia during exocytosis stimulation

The increase of [Ca²⁺] in cilia during depolarisation has been amply documented by electrophysiology (see “Section 1”). No more details are available for calcium dynamics in cilia of *Paramecium*. We now analysed different types of stimulation and we also envisaged spill-over phenomena. We loaded 7S and d4-500r cells with Fluo-3 for rapid recording of fluo signals in a CLSM equipped with an opto-acoustic beam deflection system (Fig. 1). In parallel, we performed EDX analyses.

According to previous work with 7S cells, AED-stimulated synchronous trichocyst exocytosis causes a swift rise of the cortical Ca²⁺ signal in the stimulation area to over five times basal levels within ~1 s [14,24]. We now show that this is paralleled by a similar [Ca²⁺] increase in cilia. In the typical example presented in Fig. 1A, AED stimulation is tightly followed, or superimposed by a slightly lower signal recorded in the adjacent layer of cilia in a 5 μm broad frame adjacent to the cell boundary. Although the volume contributed by cilia in such a layer is much smaller than the adjacent cortical cytoplasmic layer, the fluo evaluation applied allows one to register true changes of [Ca²⁺]. In sum, we find a considerable [Ca²⁺] increase in cilia during exocytosis stimulation. Since ciliary Ca²⁺-channels are irrelevant for exocytosis [22], this increase must be due to spill-over from the cytosol.

A similar analysis with d4-500r cells gives quite similar results. A typical example is presented in Fig. 1B. The insert shows that, after AED application, [Ca²⁺] in cilia again rises very rapidly from 20 ms on, i.e. within one ciliary stroke. As to be expected [22], this was paralleled by local trichocyst exocytosis at the stimulation site (not shown), just like in 7S cells.

Table 2 summarises cortical fluo signals thus achieved. Clearly, Ca²⁺ sweeps into the ciliary basis very rapidly. As reported, during synchronous exocytosis induction Ca²⁺ signals can be recorded by a fast CLSM analysis within such short time periods [24–26], whereby exocytosis is completed within 80 ms [40].

In sum, the new information contained in Fig. 1A and B says that Ca²⁺ spill-over into cilia occurs within one ciliary stroke and that this [Ca²⁺] increase in the basal part of cilia can induce ciliary reversal.

Fig. 1. AED-triggered exocytosis is accompanied by Ca²⁺ spill-over into cilia visible by fluorochrome analysis in 7S (A) and in d4-500r cells (B). Signals were recorded in the CLSM close to the site of AED application, not only in the outermost cell cortex, but also in a ~5 μm broad zone (containing beating cilia) adjacent to the cell surface. The time of AED application is marked by an arrow.
3.2. [Ca] recording by EDX in cilia during exocytosis stimulation

When massive synchronous trichocyst exocytosis is stimulated by AED in 7S cells spill-over of calcium into basal regions of cilia can also be seen in EDX analysis (Fig. 2). The positions of Ca K signals are marked in the STEM image, the block diagrams below show statistical evaluations from numerous such measurements. Position 3 indicates spot signals have been recorded, (B) in mock-stimulated cells and (C) in aliquots stimulated with AED (30 ms). Note that position -"5" is located on the basal body with surrounding alveolar sacs which are contained and whose Ca K signal is registered in the 0.5 μm thick section analyzed. S.E.M. = standard error of the mean. Number of measurements (from positions 1 to 7), n = 12, N = 12, each in controls and stimulation experiments.

Table 2

<table>
<thead>
<tr>
<th>Strain</th>
<th>N</th>
<th>Maximum f(3)/fx increase ± S.E.M.</th>
<th>f(3)/fx signal rise (s) ± S.E.M.</th>
<th>f(3)/fx signal decay (s) ± S.E.M.</th>
</tr>
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<tbody>
<tr>
<td>7S</td>
<td>6</td>
<td>5.4 ± 0.5</td>
<td>0.26 ± 0.03</td>
<td>1.93 ± 0.46</td>
</tr>
<tr>
<td>7S-50m</td>
<td>6</td>
<td>3.9 ± 0.9</td>
<td>0.75 ± 0.18</td>
<td>0.51 ± 0.01</td>
</tr>
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*|CaKf(3)| = 0.5 mM. N = number of cells analysed.

3.3. Effects of depolarisation on [Ca] in cilia of 7S cells

We used EDX to analyse [Ca] along the shaft of longitudinally cut cilia of 7S cells (Table 3). To facilitate comparison of data which unexpectedly showed considerable fluctuations in recordings at different positions along the ciliary shaft, we posted the Ca K values recorded within a distance of 0.5–1.5 μm from the ciliary basis and compared with those from more remote domains. This way we tried to address the question whether Ca[++] influx channels may be evenly distributed or enriched in the ciliary base. (A distance of 0.5 μm is sufficiently remote from alveolar sacs with their relatively high [Ca] [39].) It should be noted that it is very difficult to find cilia with a clear origin on a
cell body and an extension over several micrometres where, therefore, it is difficult to obtain recordings.

[Ca] values recorded in cilia from unstimulated 7S cells decrease along the cilium, from 1.5 μm on (Table 3). These cells were mock stimulated for 80 ms, i.e. they passed through the quenched-flow apparatus in the medium in which they were contained (without increased [K+]o).

In Table 3, aliquots were subjected to depolarisation for 80 ms. In this case, [Ca] rose several times, particularly in more distal regions. A similar observation was made when cross-sectioned cilia around a cell, within still larger distances from the cell body, have been analysed (data not shown), though the precise origin or distance from a cell body could not be determined in that case.

In sum, from these data one may derive the conclusion that Ca2+-channels are not restricted to the ciliary base.

3.4. No spill-over of calcium into the cytosolic compartment after depolarisation in 7S cells

Table 4 presents Ca2+ readings recorded from different regions of 7S cells, with or without 80-ms depolarisation. Table 4 also includes values recorded from alveolar sacs and from different cytosolic domains at a distance of 0.5 μm from (longitudinally cut) basal bodies. Would depolarisation cause a calcium spill-over from cilia into the nearby cytosol? Consider that, at a distance of 0.5 μm from the contact zone between docked trichocysts and alveolar sacs, we had previously recorded strong Ca2+ signals during exocytosis stimulation [39].

In alveolar sacs, Ca2+ values do not change after depolarisation (Table 4)—in agreement with the general assumption that under these conditions Ca2+ comes from the outside medium (see “Section 1”). The values have to be compared with Ca2+ readings in alveolar sacs in previous

### Table 3

<table>
<thead>
<tr>
<th>Experimental conditions</th>
<th>Distance from ciliary basis</th>
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<tbody>
<tr>
<td>Ca Kα counts recorded along longitudinally cut cilia of 7S cells, with or without 80-ms depolarisation</td>
<td></td>
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</table>

<table>
<thead>
<tr>
<th>Without depolarisation</th>
<th>0–1.5 μm ± S.E.M.</th>
<th>&gt;1.5–3.5 μm ± S.E.M.</th>
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<tbody>
<tr>
<td>Ca Kα counts</td>
<td>335 ± 5</td>
<td>59 ± 11</td>
</tr>
<tr>
<td>n = 74, N = 47</td>
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With depolarisation

| Ca Kα counts         | 436 ± 6          | 854 ± 63             |
| n = 19, N = 13        |                  |

P < 0.001

### Table 4

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<thead>
<tr>
<th>Ca Kα counts recorded in longitudinally cut cilia, alveolar sacs and cortical cytosol of 7S cells</th>
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<tr>
<td>Mock stimulation ± S.E.M.</td>
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<tr>
<td>Ca Kα counts in cilia* (mean)</td>
</tr>
<tr>
<td>n = 33, N = 8</td>
</tr>
<tr>
<td>P = 0.004</td>
</tr>
<tr>
<td>Ca Kα counts in alveolar sacs (mean)</td>
</tr>
<tr>
<td>n = 13, N = 5</td>
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<tr>
<td>P = 0.44</td>
</tr>
<tr>
<td>Ca Kα counts in cytosol** (mean)</td>
</tr>
<tr>
<td>n = 13, N = 5</td>
</tr>
<tr>
<td>P = 0.844</td>
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*All measurements from cilia pooled.

**No statistical change.

*Measured within 0.5 μm below ciliary bases.
analyses [38,39] in which we have obtained a calibrated value of \([\text{Ca}^+] \sim 43 \, \text{nM}\), Ca K0 signals in cilia 80 ms after de-polarisation must be a significant fraction of those recorded in alveolar sacs or even more (considering their small diameter in comparison to the 0.5 \, \mu m section thickness). Table 4 also shows that no spill-over beyond the ciliary basis occurs. This agrees with the fact that depolarisation-induced ciliary reversal is not paralleled by trichocyst exocytosis (Table 1).

5.5. Ciliary \([\text{Ca}^+]) in pawn cells

According to pilot experiments, the situation is rather different in d4-500r cells (data not shown). In the mock-stimulated cells, Ca K0 readings are remarkably lower than in 75 cells. Furthermore, 80-ms exposure to depolarisation conditions causes only little change. The latter had to be expected due to the absence of any Ca2+- influx under such conditions in the pawn strain (see "Section 1").

4. Discussion

Ca2+ regulates ciliary beat [8]. It is relevant for normal beat activity, not only in ciliated epithelia [9,28,29] but also in Paramecium [10,11,45]. Beyond that, a \([\text{Ca}^+] \) increase can induce a ciliary or flagellar beat reversal in lower eukaryotes, like the green alga *Chlamydomonas* [46,47], ciliates (see reviews by Preston and Saihi [19], Plattner and Kline [14] and ctenophores [48,49]). In Paramecium, several molecular targets are envisaged [53,33,50], as discussed below, with unknown distribution along the organelle.

4.1. \([\text{Ca}^+] \) increase originating from depolarisation-dependent influx is the signal for ciliary reversal

A multiplicity of signals can occur in cilia [19]. In *Paramecium*, the increase of intraciliary \([\text{Ca}^+] \) causing ciliary reversal [17,20,23] is normally initiated by activation of mechano-sensitive Ca2+-channels in the non-ciliary cell membrane which entails depolarisation of the cell membrane potential [10–13,18–20]. This in turn opens voltage-dependent Ca2+-channels in the ciliary membrane [51,52]. The very same Ca2+ rapidly inactivates Ca2+-channels from the inside [53].

A rapid intraciliary \([\text{Ca}^+] \) increase is the relevant signal for induction of ciliary reversal, but how does this compare with other signals? All evidences favour Ca2+ as the relevant signal, whereas formation of cGMP during ciliary reversal in *Paramecium* [50,54,55] may exert a modulatory effect only [45,56–60]. In *Paramecium*, guanylate cyclase is distributed over the ciliary membrane [61], but cGMP rises during ciliary reversal induction only with considerable delay [56,62], at least \( \sim 10 \) times slower than \([\text{Ca}^+] \) [55]. Formation of cGMP is even considered the result of the \([\text{Ca}^+] \) increase [63,64] occurring during ciliary reversal. Concomitantly, Ca2+, which suffices as a signal in the permeabilised *Paramecium* model [23], is the relevant parameter to be analysed in the present context. In mammalian cells, cGMP serves only to modulate ciliary beat frequency [28,65]. Our present work adds further support to this concept since \([\text{Ca}^+] \) in EDX increases in cilia within the duration of about one ciliary stroke, i.e. \( \sim 10 \) times faster than cGMP formation.

4.2. Are voltage-dependent Ca2+-channels located exclusively in the ciliary cell membrane?

Voltage-dependent Ca2+-influx channels are generally assumed to be restricted to the ciliary membrane since they disappear after deciliation and reappear during reformation of cilia [51,52]. However, Schultz et al. [66], based on their measurements of Ca2+-dependent cGMP increase after depolarisation of deciliated cells, came to the conclusion that they would be located, at least in part, also in the non-ciliary cell membrane. They assumed that such channels may still be present after deciliation, but in deactivated form, due to the different chemicals used to deciliate *Paramecium* cells.

In our EDX analyses with 75 cells, we find, upon depolarisation, an increase of \([\text{Ca}^+] \) within cilia, but not in cortical regions near basal bodies or anywhere below the non-ciliary cell membrane. This clearly favours a Ca2+ influx selectively through the ciliary membrane, particularly since, in principle, our methodology enables us to demonstrate also a Ca2+ influx via the non-ciliary cell membrane, e.g. during exocytosis stimulation [39].

4.3. Where in the ciliary membrane reside voltage-dependent Ca2+-channels?

Since antibodies for immuno-localisation are not available this question could not be resolved so far and answers have remained speculative. In fact, conflicting evidence of an equal distribution along the cilium as well as of a basal enrichment has been presented. Ca2+ spill-over from the cell soma into cilia can induce ciliary beat reversal, as we show. Similarly, in ctenophores, ciliary bending can be initiated at the basis [49]. How does that compare with other systems and which relation exists to the distribution of Ca2+-channels and Ca2+-sensitive target molecules?

A variety of Ca2+-sensitive molecular targets occur in cilia of a *Paramecium* cell where they may participate in ciliary activity. This includes calmodulin [13,67], centrin-like proteins [33], possibly a protein kinase with calmodulin-like domains [68] and perhaps, as suggested from work with *Chlamydomonas* [47], a calmodulin-dependent protein kinase (which, however, is unknown so far from *Paramecium*). Their distribution along the ciliary shaft has not been analysed in detail as yet. Moreover, in *Chlamydomonas*, immuno-localisation of p28, a protein relevant for normal beat, sometimes appears distributed equally and sometimes enriched in basal regions of flagella [69].
In an attempt to learn more about the distribution of Ca\(^{2+}\)-channels and Ca\(^{2+}\)-sensitivity along flagella and cilia, more direct approaches have been made—all resulting in suggestions which we may be able to re-evaluate. Several authors analysed ciliary aggregates in ctenophores and some flagella in Chlamydomonas. In ctenophores, initiation of ciliary bending at the basis causes mechanical self-propagation of ciliary beat [48]. In mammals, [Ca\(^{2+}\)] in basal ciliary regions accounts for accelerated ciliary beat [9]. In Chlamydomonas, voltage-dependent Ca\(^{2+}\)-conductance is lost after deflagellation, but increases to the same extent as flagella re-grow [46]. This was interpreted as localisation of Ca\(^{2+}\)-channels throughout the flagellum. A similar observation was made after de-ciliation and subsequent re-ciliation in Paramecium [51]. However, channels could also be increasingly inserted at specific sites, like at the flagellar base, as the organelle reforms. Another approach executed with Paramecium was subtraction of ciliary membranes, loading with the Ca\(^{2+}\)-fluorochrome arsenazo III and subsequent exposure to a depolarisation medium [30]. With the major vesicle subpopulations similar Ca\(^{2+}\)- signals were recorded. Although this was again interpreted as indicating equal distribution of channels over the entire cilium, the problem of sidedness after resealing, of variable vesicle subpopulations and of “hybridisation” by fusion has not been addressed.

More direct approaches were as follows. Hamasaki and Naitoh [70] have applied Ca\(^{2+}\) by micropipettes very locally along cilia. They found that basal regions are more sensitive than more distal ones. More recently, Iwadate [37] found by laser activation of caged Ca\(^{2+}\) that the distal part of the cilium also contains the Ca\(^{2+}\)-sensitive machinery relevant for ciliary beat reversal.

Our present data provide independent support of the idea that voltage-dependent Ca\(^{2+}\)-influx channels may be positioned not only in basal regions of cilia, although a [Ca\(^{2+}\)] increase in the cilium basis can suffice to induce ciliary reversal. The first argument is supported by the observation that, upon depolarisation, [Ca] does not selectively rise in the basal regions. The second aspect is supported by the occurrence of ciliary reversal in power mutants, when exocytosis stimulation provokes a Ca\(^{2+}\) spill-over from the cell body into the basis of the cilium, while normally these cells are not capable of ciliary reversal.

4.4. The ciliary Ca\(^{2+}\) signal does not sweep into exocytotic microdomains

Depolarisation does not cause exocytosis of trichocysts [21] and ciliary voltage-dependent Ca\(^{2+}\)-channels are irrelevant for this process [22]. Accordingly, in our current EDX analyses calcium is not seen to sweep into exocytotic sites and not even to the base of a cilium. This may be due to several effects, (i) the rapid deactivation of voltage-dependent Ca\(^{2+}\)-channels by the very same Ca\(^{2+}\) which enters the cilium [53], (ii) the rapid binding of Ca\(^{2+}\) by the numerous Ca\(^{2+}\)-binding target molecules (see above). These must provide a high Ca\(^{2+}\) buffer capacity, considering the magnitude of the Ca Ks signal.

4.5. Spill-over of Ca\(^{2+}\) from activated exocytosis sites into cilium

During stimulated synchronous trichocyst exocytosis Ca\(^{2+}\) mobilisation from alveolar sacs is superimposed by a “store-operated Ca\(^{2+}\) influx”, SOC, through the non-ciliary cell membrane [26,27]. SOC is triggered by the emptying of the cortical stores with which it has to act in concert. In sum, a local [Ca\(^{2+}\)] signal of ~5 μM is required for full exocytotic response [14,24,71]. Evidently this enormous Ca\(^{2+}\) flux arising from influx over the extensive non-ciliary cell membrane and from Ca\(^{2+}\) release from the extensive cortical stores is strong enough to sweep into the basis of cilia and to cause ciliary reversal, as we show. In contrast, the opposite phenomenon, i.e. spill-over from cilia into the cytosol, does not work, probably because of the rapid inactivation of ciliary Ca\(^{2+}\)-channels and of a high Ca\(^{2+}\) buffer capacity.

4.6. Comparison with electrophysiological data and balance calculations

According to Table 4, Ca Ks counts in cilia rise approximately five times after depolarisation. Since longitudinally cut cilium occupy only ~50% of the section thickness used, comparison of the Ca Ks values registered in cilia after depolarisation with those in alveolar sacs (where it remains unchanged during stimulation) indicate a comparable [Ca] in these organelles. Recall that in alveolar sacs [Ca] is ~43 mM [38,39]. Are our data compatible with estimations by other methods?

Based on voltage clamp analyses, an estimation of Ca\(^{2+}\) molecules which cross the ciliary membrane under optimal conditions can be attempted. A pilot calculation shows that this could yield a maximal [Ca] within cilia of the order of ~50 mM (R.R. Preston, personal communication) which is easily detectable by EDX [38]. Assuming that in cilia ~99.9% of Ca\(^{2+}\) are rapidly bound [72] both our data sets, those of [Ca] and [Ca\(^{2+}\)] combined, would result a residual free Ca\(^{2+}\) concentration of 50 μM, i.e. ~700 residual Ca\(^{2+}\) atoms in a cilium after activation. A broader range of estimation, from 10 μM to 1 mM, has also been reported [73,74]. All these values exceed the reported activation by 1 μM in vitro [23] when permeabilisation may cause CaBP extraction or other phenomena which may entail a lower activation threshold.

Would our estimation, [Ca\(^{2+}\)] ~ 50 μM in activated cilia, be sufficient for detection by fluorochromes? Our fNa ratio evaluation does not allow to directly compare readings in cilia and in the nearby cortical domains. Though the fluorochrome signal in cilia during reversal is smaller than that in the nearby cell cortex, relative values recorded in cilia
must considerably exceed the signal recorded in the nearby cell cortex—otherwise it would be undetectable according to our experience. A fraction of only 1.4% would be expected if [Ca²⁺] are equal in cilia and cell cortex since cilia (more precisely, their basal regions) contained in the extracellular layer occupy only ~1.4% of the volume available (one cilium per kinetid containing 3.45 μm²) [356]. Since ciliary signals are well detectable the high magnitude of [Ca²⁺] estimated above for cilia after depolarisation appears realistic. If so, calcium homeostasis must be re-established after depolarisation by a high buffer capacity, considering the absence of diffusion into the cell cortex (this paper) and the reported absence of a Ca²⁺-pump (ATPase) from ciliary membranes in *Paramecium* [75].

4.7. Functional aspects of ciliary reversal

Ciliary reversal serves the avoidance reaction when a *Paramecium* cell hits an obstacle. In a more complicated situation, when a *Paramecium* cell meets a predatory organism, the first reaction is expulsion of elongate trichocysts to keep the predator at a distance, as first shown by Harumoto and Miyake [76] and in more detail by Knoll et al. [44]. The concept of a defensive function of trichocyst expulsion [76] can be supplemented by the observation that this is followed with little delay by ciliary reversal [44]. The effect can be seen, e.g. in Fig. 3 of Ref. [44], where a 75 cell, after releasing trichocysts upon contact with a *Dileptus* cell, withdraws by ~40 μm (cell width) within 12 ms—too short a time to be captured. The present work shows the possible basis of a tandem signalling involved in these two successive reactions.

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References


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