Paramecium tetraurelia basal body structure
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Abstract
Paramecium is a free-living unicellular organism, easy to cultivate, featuring ca. 4000 motile cilia emanating from longitudinal rows of basal bodies anchored in the plasma membrane. The basal body circumferential polarity is marked by the asymmetrical organization of its associated appendages. The complex basal body plus its associated rootlets forms the kinetid. Kinetids are precisely oriented within a row in correlation with the cell polarity. Basal bodies also display a proximo-distal polarity with microtubule triplets at their proximal ends, surrounding a permanent cartwheel, and microtubule doublets at the transition zone located between the basal body and the cilium. Basal bodies remain anchored at the cell surface during the whole cell cycle. On the opposite to metazoan, there is no centriolar stage and new basal bodies develop anteriorly and at right angle from the base of the docked ones. Ciliogenesis follows a specific temporal pattern during the cell cycle and both unciliated and ciliated docked basal bodies can be observed in the same cell. The transition zone is particularly well organized with three distinct plates and a maturation of its structure is observed during the growth of the cilium. Transcriptomic and proteomic analyses have been performed in different organisms including Paramecium to understand the ciliogenesis process. The data have incremented a multi-organism database, dedicated to proteins involved in the biogenesis, composition and function of centrosomes, basal bodies or cilia. Thanks to its thousands of basal bodies and the well-known choreography of their duplication during the cell cycle, Paramecium has allowed pioneer studies focusing on the structural and functional processes underlying basal body duplication. Proteins involved in basal body anchoring are sequentially recruited to assemble the transition zone thus indicating that the anchoring process parallels the structural differentiation of the transition zone. This feature offers an opportunity to dissect spatio-temporally the mechanisms involved in the basal body anchoring process and transition zone formation.

Keywords: Paramecium, Basal body assembly, Basal body docking, Transition zone, Ciliogenesis

The organism
Paramecium tetraurelia is a unicellular eukaryote belonging to the Chromalveolata kingdom, Ciliophora phylum. It is a free-living bacteriophagous organism that is easy to cultivate, usually found in freshwater where it can swim and capture its preys thanks to its ca. 4000 cilia (Fig. 1).

Basic basal body structure
In Paramecium, basal bodies are arranged into parallel rows, the kineties, patterned along the antero-posterior axis of the cell, and their rotational polarity is marked by the asymmetrical organization of their associated structures (Fig. 1b). These basal bodies are anchored at the cell surface and embedded in a superficial cytoskeletal layer, the epiplasm, which partitions the cell surface in cortical units [5–8]. In interphasic cells, either one or two basal bodies are anchored in the middle of each cortical unit (Fig. 1c). Basal body duplication occurs close to the mother basal body, which, contrary to mammalian cells, is permanently anchored at the cell surface. During cell division, several successive duplication waves of basal bodies and associated appendages together with the formation of new cortical units lead to the duplication of the whole cell pattern [7]. Paramecium basal bodies display a ninefold symmetry of microtubule blades with, at their proximal
end, microtubule triplets surrounding a cartwheel that remains present all along their life cycle [9, 10] (Fig. 2). Microtubule triplets (A, B and C tubules) end up at the level of the transition zone and microtubule doublets are observed above [11] (Fig. 2). All Paramecium basal bodies show a similar organization, but their length varies between 330 and 600 nm depending on their location on the cell cortex (Fig. 3). The height of the cartwheel correlates with the basal body length; the longest are present in the oral apparatus, while the shortest are scarce and found only in the cell cortex [12].

Dute and Kung [11] have studied the structure of the Paramecium transition zone in detail using both thin-section and freeze-fracture electron microscopy. This transition zone is particularly well delimited and organized in three distinct plates namely the terminal, the intermediate and the axosomal plates (Fig. 3). The terminal plate marks the boundary between the basal body and the transition zone. Hufnagel [5] showed that this plate is organized around a central rim, from which nine spokes radiate into the gap between the microtubule doublets. This plate is at least partially built up with epiplasmins, the protein components of the epiplam [8]. Nine perforations in the epiplam ring located around the microtubule doublets have been observed in both Paramecium [5] and Tetrahymena [13]. These structures correspond to the ciliary pores described by Ounjai et al. [14] in Tetrahymena basal bodies after potassium phosphotungstate treatment to remove the microtubules. Transitional fibres are difficult to observe but a pinwheel structure surrounding the microtubule doublets has been disclosed. It originates from the terminal plate and ends beneath the plasma membrane. Close to the axonemal plates, peg-like Y-shaped structures called Y-links project from the common wall of the A and B tubules [11] (Fig. 2). In Paramecium, docked basal bodies are not systematically ciliated so that both non-ciliated and ciliated docked basal bodies can be observed in the same cell. Growth of cilia is accompanied by modifications of the structure of the transition zone which appears more collapsed in anchored non-ciliated basal bodies than in the ciliated ones (Fig. 3) [15]. Two recent papers concerning the function of two transition zone proteins MKS1 and MKS3 [16, 17] report loss of cilia after their depletion suggesting that in Paramecium MKS1 and MKS3 are essential for cilium stability as in mammals [18].

In the tubulin superfamily comprising six members, δ-tubulin has been shown to be required for the
C-tubule assembly [19], while ε-tubulin was necessary for B- and C-tubule assembly or stabilization of the microtubule triplet. In addition, basal body duplication is impaired after ε-tubulin depletion [20]. A mutation found in sm19, encoding the rare eta now called ζ [21], was reported to inhibit basal body duplication and to induce the delocalization of γ-tubulin [22, 23].

**Additional basal body structures or accessory structures**

Three major appendages typical of all Ciliata are found to be associated with Paramecium basal bodies: a long striated rootlet crossing over several cortical units and two microtubular rootlets, the transverse microtubules and the post-ciliary microtubules [12, 24, 25]. They protrude asymmetrically from the proximal part of the basal bodies. The direction of these cytoskeletal appendages correlates with the antero-posterior axis of the ciliary row; the pattern of these ciliary rows correlates with the global cell polarity with its right–left asymmetry and antero-posterior axis (Fig. 1b). The striated rootlet (kinetodesmal fibre) is connected to the right side of the basal body and extends along the basal body row towards the anterior pole of the cell (Fig. 1b). The transverse microtubules originate close to the anterior basal body and run perpendicular to the basal body row towards the left side of the cortical unit. The post-ciliary microtubules originate close to the posterior basal body, and run perpendicularly to the basal body row towards the posterior pole of the cell [12, 24, 25]. The striated rootlet guides the new basal body during its positioning [10].

Lynn [26] has proposed that the basal body and its associated set of rootlets form the kinetid. Monokinetids (single basal bodies) display the three rootlets, while in dikinetids (paired basal bodies) only one post-ciliary and one striated rootlet are present. They are associated with the posterior basal body, while both anterior and posterior basal bodies have associated transverse...
microtubular rootlets (Fig. 2). In both mono- and dikeratids, a delicate system of fibres links the basal bodies to their rootlets [12]. Further details on the fate of permanent appendages during duplication are available in [10].

An additional transient appendage, called the anterior left filament (ALF), develops at the anterior left from the mother basal body (triplets n°4) before duplication and disappears once the new basal body is docked at the surface [27]. This transient filament requires Centrin3 for its formation and is assumed to be involved in the tilting up of the new basal body allowing its anchoring [27].

**Basal body origins**

All basal bodies develop from a pre-existing mother basal body and in contrast to other ciliates such as *Oxytricha* [28, 29], *Sterkiella* [30] or *Paraurostyla* [31], no de novo assembly of basal bodies has ever been observed in *Paramecium*.

**Basal body life cycle and other functions**

In *Paramecium*, basal bodies are dedicated to organize cilia required for motility, food uptake, sensory functions and cell–cell recognition during sexual reproduction. Basal bodies never act as a centrosome to organize the mitotic nuclear spindle. They duplicate close to their parents and remain anchored at the cell membrane during the whole cell cycle. During division, new basal bodies act as organizing centres for the assembly of new cortical units. Their microtubular rootlets probably act as templates for the assembly of a superficial sub-membranous spindle, the cytoplasmic transitory detected during division [32, 33].

**Identification of basal body components**

In this species, no proteomic or genomic screens dedicated to dissecting basal body composition have been performed so far. However, a proteomic screen of isolated cilia [34] and a study of transcriptomic changes during ciliary biogenesis in response to deciliation have been carried out to understand the ciliogenesis process [35]. In addition, a ciliary membrane proteomic analysis has been reported recently [36]. Results from these analyses were all included in the Cildb database (http://cildb.cgm.cnrs-gif.fr/) [34, 37]. Together with 66 high-throughput studies from 15 eukaryotes having centrioles/basal bodies and cilium, they enabled Carvalho-Santos [38] to establish simplified phylogenetic profiles of the structure and function of these organelles.

**Notable basal body findings**

In 1965, Beisson and Sonneborn [39] demonstrated that the polarization of newly assembled kinetids is determined by the polarity of the cortical environment existing at the time of their development, leading to the concept of “cortical inheritance” or structural memory.

In 1968, Dippell [9], thanks to thousands of basal bodies arising almost synchronously in known positions at a recognizable stage, has followed the duplication steps of the basal body and dissected the formation of the microtubule wall in a pioneer electron microscopy study.

In 1994, Redecker et al. [40] have discovered a new tubulin modification using *Paramecium* axonemal tubulin. This post-translational modification, which affects kinetid microtubules a lag-time after their assembly, has
been suggested to act as a marker to discriminate parental and new structures during cell division [33].

Ruiz et al. [23], taking advantage of the fact that cells continue dividing after inhibition of basal body duplication, demonstrated for the first time the requirement of γ-tubulin for basal body duplication. They also demonstrated first that Centrins are required for basal body positioning but not for its duplication per se [41].

In 2000, a novel member of the tubulin superfamily was identified using the sm19 mutant, and called η/ζ-tubulin. The mutations in the gene encoding this protein caused an inhibition of basal body duplication [42]. Further physiological and genetical studies indicate an interaction with microtubules [22]. In a recent functional study of ζ-tubulin in *Xenopus* [21], no evident basal body defect was observed, leading to the conclusion that ζ-tubulin is involved in basal body orientation and distribution and might function by interacting with other tubulins.

Finally, studying the process of basal body anchoring through a combination of GFP-fusion protein expression, RNAi and low-resolution electron microscopy, Aubusson-Fleury et al. [15] discovered that three proteins, Centrin2, FOR20 and Centrin3, are sequentially recruited to allow basal body docking is not systematically coupled to ciliogenesis, which occurs throughout the cell cycle (Aubusson-Fleury, in preparation). Therefore, both non-ciliated and ciliated docked basal bodies can be observed in the same cell. In anchored non-ciliated basal bodies, the transition zone appears more collapsed than in the ciliated ones, suggesting that maturation of this region occurs during axoneme extension [15]. Multi-disciplinary approaches, combining biochemical, molecular and cell biology techniques with high-resolution ultrastructural approaches, will permit characterization of the molecular and structural bases of the transition zone assembly process in a 4D-space. This will open the way to further studies linking atomic structure with transition zone assembly.

**Strengths and future of basal body research in *Paramecium***

The strengths of the *Paramecium* model rest in its long established status as a genetic model [43] and its large number of basal bodies. Their regular arrangement over the cell surface, as well as the precise choreography of their duplication, facilitates functional analyses of ciliary proteins both at the cellular and ultrastructural levels. Such a precise organization has allowed, in the past, to perform pioneer and high-quality ultrastructural studies of the basal body duplication process [9] and the transition zone arrangement [11]. The coupling of rapid and non-costly functional analyses of candidate genes based on RNAi knock-down using feeding techniques [44], and overexpression of tagged-fusion proteins by nuclear microinjection of DNA, makes *Paramecium* an outstanding and efficient model to study basal body assembly and functions. It is also particularly suitable for assessing the process of basal body anchoring, since anchoring defects are detected by immunofluorescence using specific antibodies and can be easily followed from the early steps of protein depletion.

**What are the prospects for future work?**

Unravelling the structural aspects of cilia assembly is fundamental in the study of ciliopathies. Owing to the motility of its cilia, *Paramecium* is obviously an appropriate model to study the function of proteins involved in primary ciliary dyskinesia (PCD), a pulmonary disease arising from immotile respiratory cilia. The transition zone acts as a filter between the cellular and the cilium compartments and houses many proteins involved in human ciliopathies. However, the structural bases of the ciliary gate function are currently unknown. The large number of basal bodies in *Paramecium*, and the easiness to prepare *Paramecium* cell cortices, will allow studying the structural organization of the *Paramecium* transition zone using high-resolution microscopy such as cryo-electron tomography [45] or scanning transmission electron tomography. In addition, in *Paramecium*, the basal body docking is not systematically coupled to ciliogenesis, which occurs throughout the cell cycle (Aubusson-Fleury, in preparation). Therefore, both non-ciliated and ciliated docked basal bodies can be observed in the same cell. In anchored non-ciliated basal bodies, the transition zone appears more collapsed than in the ciliated ones, suggesting that maturation of this region occurs during axoneme extension [15]. Multi-disciplinary approaches, combining biochemical, molecular and cell biology techniques with high-resolution ultrastructural approaches, will permit characterization of the molecular and structural bases of the transition zone assembly process in a 4D-space. This will open the way to further studies linking atomic structure with transition zone assembly.
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