Abnormal Basal-Body Number, Location, and Orientation in a Striated Fiber-defective Mutant of *Chlamydomonas reinhardtii*

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**ABSTRACT** We describe a mutant of *Chlamydomonas reinhardtii* in which basal body associated striated fibers are absent or incomplete. Basal body spacing, angle, and relative rotational orientation are abnormal and extremely variable. Abnormal partitioning of cellular contents at cytokinesis is also evident. Mating, maintenance of flagellar length equality, and backward swimming response are normal. Genetic analysis indicates mutation of a new Mendelian gene—vfl-3—linked to the centromere of Chromosome VI.

Striated fibers associated with basal bodies (6, 26, 30, 44) and centrioles (21, 34, 39) have been observed in numerous eukaryotic cells, and they have been given a variety of names, including rhizoplasts (8), kinetodesmal fibers (1), and ciliary or flagellar rootlets (20). The association of striated fibers with basal bodies and centrioles is so characteristic that a relationship among all striated fibers—or at least the existence of a limited number of fiber families—seems probable (5, 26), in spite of the considerable variation in size, shape, and periodicity shown by fibers from different sources.

The very ubiquity of striated fibers makes it likely that they play important cellular roles. A number of passive roles such as anchoring or supporting basal bodies, maintaining basal body position or dissipating mechanical stress have been proposed for the fibers (30, 44). More active roles have also been considered. For instance, it has been suggested that the fibers coordinate motility, since when basal bodies are linked by striated fibers their flagella typically show coordinated beating (3, 7, 12, 15). Observations of ATPase activity in fiber cross-bands (2, 24), proximity between mitochondria and striated fibers (28, 40, 42), and variations in fiber periodicity (14, 35, 36) have all been cited as suggesting an active contractile function for the striated fibers (30, 37). Salisbury and Floyd (35) demonstrated calcium-dependent contraction of a large striated fiber—the rhizoplast of *Tetraselmis*—and they were able to produce cycles of contraction/relaxation by alternate incubations with and without CaCl₂ and ATP. A calcium-dependent decrease in the normal 90° angle between the paired basal bodies of isolated *Chlamydomonas* flagellar apparatuses was reported by Hyams and Borisy (15), who speculated that this might be due to striated fiber contraction.

We decided to approach the question of striated fiber function by isolating and examining fiber-defective mutants in the biflagellate alga *Chlamydomonas reinhardtii*. The two basal bodies of *C. reinhardtii* are oriented at right angles to one another and are connected by three striated fibers (32). The largest of these, the “distal striated fiber,” connects the basal bodies near their cell-distal ends. The distal fiber has approximate dimensions of 300 nm x 250 nm x 75 nm and a banding pattern consisting of a central pair of dark striations flanked on either side by two light bands, two dark bands, and a dark band close to the basal body (see Fig. 2b). Two parallel “proximal striated fibers” connect the basal bodies at their cell-proximal ends. Proximal fibers have approximate dimensions of 140 nm x 45 nm, and each carries three or four cross striations (see Fig. 2c).

Having reasoned that striated fibers are probably necessary for normal coordination of flagellar motility, we isolated a number of aberrant motility mutants and examined them for striated fiber defects. Here we describe the phenotype and ultrastructure of a mutant clearly defective with respect to its striated fibers.

**MATERIALS AND METHODS**

**Strains and Culture Conditions:** Wild-type strains used were the l37c derivatives NOmt* and NOmt- provided by Dr. U. Goodenough (Washington University). Strains CC530 (acl7mt), CC 624 (pfl6mt), CC1084 (nr-lthi-lma*) and CC1139 (nr-lthi-lma*) were provided by the Chlamydomonas Genetics Center, Duke University (Durham, NC). vfl-2 was vfl-2-220 (19). Cells were grown at 25°C in medium I of Sager and Granick (33), bubbled continuously with filtered air. Cultures were illuminated with white light (45 cm from two General Electric F48T12-CW-HO fluorescent tubes; General Electric Co., Wilmington, MA) and synchronized by a 14 h light/10 h dark cycle. Experiments were typically performed between hours 2 and 6 of the light segment of the cycle.

**Mutagenesis and Mutant Isolation:** Log phase NOmt* cells were spread on 1.5% agar plates and exposed to ultraviolet light (48 cm from a General Electric Co. G30T8 germicidal lamp) for 90 s (16, 23). Cell survival was...
Measurements of Flagellar, Location, and Length: Cells were fixed by addition of ½ volume of 1% aqueous glutaraldehyde, and phase-contrast microscopy at \( \times 800 \) was used to score number and location of flagella. Flagellar length was measured with an ocular micrometer, using the same optics. Distribution of flagella in a population was based on scoring no less than 200 cells per sample. Average values for flagellar length were determined from measurements of 20 or more cells of each flagella-number class, unless otherwise noted.

Motility Observations: Observation of living cells used phase-contrast microscopy at magnifications between 125 and 800. Flagella of live C. reinhardtii often become momentarily attached to the glass slide or coverslip; by counting the number of flagella on a cell when the flagella were attached and by observing motility after the cell detached, correlation between flagella number and cell motility could be made.

Electron Microscopy: Cells between the second and fourth hours of the light phase of the illumination schedule were harvested by gentle centrifugation (3 min at 2,500 rpm in a Beckman T-6 centrifuge. Beckman Instruments, Fullerton, CA) in glass culture tubes and then subjected to either of the following fixation protocols.

Collidine-Based Fixation: To the pellet of cells, ½ ml of an ice-cold solution composed of 2.5% glutaraldehyde, 0.2% tannic acid, and 0.1 M sodium cacodylate, pH 7.8, was added. The pellet was resuspended in the fixative, then replicated as described above. The initial fixation solution was removed and fresh fixative added carefully to avoid disturbing the pellet. (Centrifugation was repeated throughout the protocol, as required, to maintain the pellet.) Fixation was allowed to proceed on ice, for 1 h. Cells were then rinsed with at least five changes of ice-cold 0.1 M sodium cacodylate buffer, pH 7.8, over a period of at least 1 h, keeping cells on ice. At this point, samples were often stored at 4°C overnight.

Samples were postfixed for 1 ½ to 2 h in ice-cold 1% osmium tetroxide in 0.1 M sodium cacodylate buffer, pH 7.8, rinsed twice in water, and stained en bloc with 1% aqueous uranyl acetate for 15–20 min.

Cacodylate-Based Fixation: To the pellet of cells we added 0.5 ml of an ice-cold freshly prepared solution of 2.5% glutaraldehyde, 2% OsO\(_4\), in 0.1 M sodium cacodylate buffer, pH 7.8. The pellet was resuspended in the fixative and allowed to stand on ice for 3 min. The cells were replicated and the initial fixative was replaced with an equal volume of fresh fixative. After 30 min, we washed samples at least five times in 0.1 M sodium cacodylate buffer, pH 7.8. Samples could be stored overnight at 4°C in buffer at this point. Pellets were postfixed with 2% OsO\(_4\), in 0.1 M sodium cacodylate buffer, pH 7.8, for 1.5–2 h and rinsed twice in water. If en bloc staining was to be performed, we also rinsed the sample twice with 50% aqueous ethanol. En bloc staining was accomplished with 0.25 to 0.5% aqueous uranyl acetate for 15 min.

All samples were rapidly dehydrated in a graded alcohol series, followed by two rinses in propylene oxide, and infiltrated with resin (60 ml of DDSA, 20 ml of Araldite 502, 20 ml of Embed 812, and 3 ml of DMP-30). Infiltration was complete in two or more flagella. Distinct pairing of the flagella is commonly observed on cells with two or more flagella, so that many cells can be scored as having one pair plus one single, two pairs, etc. Average flagellar length is 9.8 \( \mu \)m, about 2 \( \mu \)m shorter than wild-type controls. With increased flagellar number, average flagellar length decreases (Table I), but the effect is not dramatic. As in wild-type biflagellates, all flagella on individual multiflagellate vfl-3 cells are equal in length.

**Mitotic Pedigree Analysis**

A pfl6,vfl-3 double mutant was constructed and used in mitotic pedigree analysis. The pfl6 mutation renders cells immotile, so that daughters of individual cells can be scored for flagellar number before and after division. The results indicate no close relationship between flagellar number of a parent cell and flagellar number of its progeny. We frequently observed abnormally small daughters among the progeny of individual vfl-3 cells, and cell debris was present in

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**RESULTS**

**Isolation of vfl-3**

In a screening of 2037 mutagenized clones, we identified 33 aberrant swimming mutants. Among the initial five that we examined by thin-section electron microscopy to determine striated fiber phenotype, one (RW207) was found to present distinct fiber defects. Genetic analysis (see below) revealed that RW207 carries a mutation in a previously unidentified mendelian gene. We have given this gene the name vfl-3 (for variable flagellar number), and the allele the name vfl-3-207. As this is the only vfl-3 allele identified to date, we will refer to it simply as vfl-3 throughout this paper.

**Distribution, Location, and Length of Flagella in RW207**

Variable number of flagella per cell is a striking feature of the vfl-3 phenotype. As indicated in Table I, roughly equal numbers of cells with zero, one, and two flagella comprise ~90% of a mutant population; cells with three or four flagella are not uncommon, and at low frequency (~0.3%) cells with five or more flagella are present. Flagellar number per cell averages 1.3.

In addition to flagellar number defects, many vfl-3 cells show abnormalities in the sites of flagellar placement on the cell surface. In ~10% of the cells, flagella are located offcenter or to the side, whereas in wild-type controls they are found exclusively at the anterior cell apex. Such flagellar displacements are more likely to occur on cells with two or more flagella. Distinct pairing of the flagella is commonly observed on cells with two or more flagella, so that many cells can be scored as having one pair plus one single, two pairs, etc. Average flagellar length is 9.8 \( \mu \)m, about 2 \( \mu \)m shorter than wild-type controls. With increased flagellar number, average flagellar length decreases (Table I), but the effect is not dramatic. As in wild-type biflagellates, all flagella on individual multiflagellate vfl-3 cells are equal in length.

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**Table I**

| Flagellar Number Class | % Total Cells Scored | Average Flagellar Length \( \mu \)m |
|-------------------------|----------------------|-------------------------------|
| 0                       | 20                   | 10.8                          |
| 1                       | 36                   | 10.8                          |
| 2                       | 34                   | 9.8                           |
| 3                       | 7.5                  | 9.5                           |
| 4                       | 1.7                  | 8.5                           |

*2,061 cells scored.

1 Average of 45 or more cells, except for quadriflagellate class, which is the average of 11 cells.
the mutant microcolonies in many cases. Furthermore, many mutant microcolonies contained odd numbers of cells, whereas most control \((pf-16)\) microcolonies contained either four or eight cells. The death and lysis of some daughters, and the abnormally small size of others, suggest a defect in the control of cell division.

**Light Microscopic Observations — Motility**

\(Vfl-3\) cells display a diverse and complex motility pattern. The majority of motile cells rotate in place. Monoflagellates show this kind of twirling motility, but, surprisingly, most biflagellates show twirling behavior as well. A "motile swirl" pattern is displayed by some biflagellate cells; these cells swirl as described above, but they also progress slowly across the field of view, looking like spinning tops. In addition to the predominant twirlers, cells with a wide variety of other abnormal patterns are present. Wildly erratic motility, zig-zagging, flagella-first tumbling, and slow swimming are all observed in mutant cultures. Very rare cells swim in a pattern approaching normal, but truly wild-type motility has not been observed.

In *Chlamydomonas* a sudden increase in illumination or collision with an object produces a backward-swimming response: flagella are thrown forward, the beat pattern changes from a ciliary to a flagellar (i.e., symmetric) form, and the cell moves slowly backward. A normal backward-swimming response has been observed in mutant cells of all motility types, regardless of number of flagella.

\(Vfl-3\) cells are not generally capable of phototaxis. Thus cells in cultures illuminated from above do not gather at the meniscus as wild-type cells do; rather, they fall to the bottom of the culture vessel. The absence of phototactic ability in mutant cells is probably a consequence of their grossly aberrant motility, and we believe that it does not reflect a primary defect in phototaxis itself. A few cells (\(<6\%) make it to the surface, and observation of these cells indicates a pronounced enrichment for the less severely affected motility types such as tumblers and slow-swimmers.

**Defective Basal-Body Orientation and Position in \(vfl-3\)**

Basal body, transition region, and axoneme structure in \(vfl-3\) are generally normal, as the thin-section electron micrographs in Fig. 2 illustrate. However, striking abnormalities in basal-body position and orientation are evident. Whereas wild-type control cells show little variation in angle and distance between their two basal bodies, we observed wide variation in the mutant. Results of measurements made on micrographs containing longitudinal sections through two or more basal bodies are tabulated in Table II. Note that the majority of mutant pairs have angles \(<60^\circ\), whereas we observed no angle \(<62^\circ\) in wild type. In wild-type cells, measured distances between the centers of transition regions exhibited only a limited range (600-770 nm), while the distance in mutant cells ranged widely (330-1,980 nm). Perfect longitudinal sections through both basal bodies are never seen in wild-type cells because the longitudinal axes of the two basal bodies lie in different planes, separated by slightly less than one basal-body.
diameter, but such sections, sometimes including three basal bodies, have been seen a number of times in vfl-3 (e.g., Fig. 2 d and f). It is our clear impression that the number of basal bodies per vfl-3 cell is considerably greater than in wild type even though the number of flagella per cell averages only 1.3; thus, mutant cells must contain many basal bodies without attached axonemes.

A variety of arrangements are seen in sections showing three basal bodies. Observed arrangements include basal bodies approximately parallel and close together in the same plane, basal bodies approximately parallel and widely separated, and two basal bodies parallel, close together, and at a right angle to a third.

Abnormal flagellar location with respect to the cell as a whole is fairly common, with flagella extending from the sides of cells or from other abnormal positions. Basal bodies have been found deep in the cell interior rather than at the cell surface, and they can be parallel to the surface or even upside down (Fig. 3).

**TABLE II**

| NO* (wild type) | vfl-3 |
|-----------------|-------|
| Angle in degrees | Distance | Angle in degrees | Distance |
| 9               | 360 nm  | 62               | 720 9 nm  |
| 67              | 600 nm  | 71               | 630 25* nm |
| 73              | 650 nm  | 76               | 660 25 nm  |
| 77              | 680 nm  | 84               | 720 38 nm  |
| 86              | 770 nm  | 87               | 710 38 nm  |
| 89              | 690 nm  | 90               | 680 55 nm  |
| 90              | 770 nm  |                  |        |

* Triflagellate section.

![Figure 3](https://example.com/figure3.png) Abnormal basal-body placement in vfl-3 cells (A) Basal body lying nearly parallel to cell surface. Collidine fixation. × 44,000. (B) Basal body upside down, with transition region facing cell interior. Note striated projection on other, transversely sectioned, basal body. Collidine fixation. Bar, 500 nm. × 44,000.

![Figure 2](https://example.com/figure2.png) Relative placement of basal bodies in wild type (A, B, and C) and vfl-3 (D, E, and F). (A and B) Wild type basal-body pairs, showing distal striated fiber. Collidine fixation. (A) × 33,000 and (B) 54,000. (C) Wild-type basal-body pair, showing proximal striated fibers. Note microtubular rootlets and probasal body. Collidine fixation. × 33,000. (D) Abnormal angle and spacing between vfl-3 basal bodies. Note Absence of distal striated fiber. (E) Longitudinal section through vfl-3 basal body, transition region, and axoneme. Basal-body structure appears fully wild type, but note abnormal projects from basal body (arrows). Collidine fixation. × 54,000. (F) Three vfl-3 basal bodies in nearly perfect longitudinal section. Collidine fixation, × 45,000. (b) Basal body. (c) Chloroplast. (df) Distal fiber. (mtr) Microtubular rootlet. (p) Probasal body. (pf) Proximal fiber. (tr) Transition region. Bar, 500 nm.
been observed. Mutant basal bodies may be completely free of fibers, or they may carry striated or non-striated projections at a variety of locations. Often, the projections have a diffuse and tenuous appearance as compared to wild-type striated fibers. With one exception, a projection has never been observed to fully span the space between the basal bodies, and even in this exceptional case fiber striations and basal body orientation were abnormal. Fig. 4 illustrates a variety of striated fiber defects observed in mutant cells.

Only 16% (19 of 116 basal bodies scored) of sections through mutant basal bodies showed evidence of attached projections or densities which resembled striated fibers, whereas 48% (71 of 147 basal bodies scored) of such sections in wild-type controls showed striated fibers of standard morphology. Because vfl-3 cells contain extra basal bodies—i.e., basal bodies without attached axonemes—the high frequency of total fiber-less basal bodies could reflect an abundant class of basal bodies with neither fibers nor axonemes.

In wild-type Chlamydomonas, four microtubular rootlets radiate into the cytoplasm from a dense plate (32) (also described as a "finely striated fiber" [10]) located between the basal bodies and beneath the distal striated fiber. Two of the rootlets typically consist of four microtubules which show a "3 over 1" arrangement as they approach the inter-basal body region, and the other two consist of a pair of microtubules each (10, 27, 32). In vfl-3 the organization and positioning of the rootlet microtubules varies from cell to cell. The "3 over 1" configuration has not been observed, and in many cases microtubules are seen to extend through, rather than terminate at, the region between basal bodies. The dense plate is usually absent as well. Fig. 5 shows representative rootlet abnormalities in vfl-3, as well as patterns seen in wild-type controls.

**Positioning of Other Organelles in vfl-3**

Although the configuration of the entire basal-body complex is abnormal and highly varied in vfl-3 cells, the overall orientation of other cellular organelles is usually normal. However, occasional displacements of the nucleus and/or pyrenoid occur, with the nucleus sometimes located far anterior near the basal bodies or with nucleus and pyrenoid side-by-side. Rare cells with two nuclei or pyrenoids have also been observed.

**Genetic Analysis**

Tetrad analysis of vfl-3 is summarized in Table III. The mutation segregates 2:2 from wild type in tetrads derived from vfl-3/wild-type meioses. It is centromere-linked based on crosses against the centromere marker ac17 (Table III, line 1). It is not linked to nr-1 (Table III, line 2); since nr-1 and the variable flagellar number mutation vfl-1 (M. Adams, personal communication) are both on the right arm of Chromosome VIII, it follows that vfl-3 and vfl-1 are not linked. It is not linked to vfl-2, since of 32 vfl-3 × vfl-2 progeny examined eight were wild type (recombination frequency = 50%). Finally, we found that vfl-3 is linked to the mating type locus (Table III, line 3), placing it on Chromosome VI. The calculated distance between vfl-3 and mating type is 32 map units. This is fully consistent with vfl-3's centromere linkage, since mt and the centromere are about 35 map units apart. We conclude that the vfl-3 mutation defines a new gene closely linked to the centromere of Chromosome VI.

**DISCUSSION**

**Vfl-3 Defect Expressed Early in Basal-Body Cycle**

In wild-type Chlamydomonas basal-body replication and segregation occur at characteristic points in the cell cycle. As early as 1 h after cell division two probasal bodies appear near the mature basal bodies (4) (Fig. 5b). Fibrous connectors join each probasal body to the mature basal bodies to form a basal body/striated fiber/probasal body complex which can be isolated intact from lysed cells (4, 11, 38). Elongation and maturation of the probasal bodies occurs during interphase, so that as cell division approaches four mature basal bodies are available for segregation to the two daughters.

Recent evidence suggests that basal-body segregation is semiconservative, with each daughter receiving one newly matured and one old basal body (13). Such a mode of segregation would clearly require that the striated fiber connections between the parental basal bodies be broken and that new connections between old and newly matured basal bodies be established. There are conflicting reports in the literature as to just when these events occur. Johnson and Porter (17) observed that cells in the midst of cytokinesis carry two pairs of basal bodies with the members of each pair connected by striated fibers—strongly suggesting that fiber dissolution and reappearance both precede cell division. In contrast, Cavalier-Smith (4) found that daughter cells shortly after cell division contained basal-body pairs which lacked striated fibers. The basal bodies were close together and nearly parallel (an arrangement reminiscent of what is frequently seen in vfl-3). Differences in strains or culture conditions could account for these discrepancies. We have not determined which pattern prevails in our cultures.

In vfl-3 isolated probasal bodies unconnected to mature basal bodies or other probasal bodies are frequently observed by mid-interphase (e.g., Fig. 6f). We conclude that the vfl-3 defect is expressed before this time—i.e., early in the cell cycle. Indeed, it is reasonable to propose that the host of ultrastructural defects observed in vfl-3 all derive from a primary failure to properly position probasal bodies in the young cell, since without appropriate basal-body placement: (a) striated fibers might be unable to form properly, (b) microtubular rootlets might be unable to arrange themselves correctly, and (a) orderly basal-body segregation at cell division might be impossible.
TABLE III
Tetrad Analysis

| Markers scored | PD | NPD | T  |
|----------------|-----|-----|----|
| vfl-3, ac-17   | 5   | 8   | 0  |
| vfl-3, nr-1    | 11  | 14  | 39 |
| vfl-3, mt      | 36  | 1   | 51 |

The absence of a simple mitotic segregation pattern with respect to flagellar number is consistent with such unordered segregation. However, because vfl-3 appears to contain many basal bodies without attached flagella, it is not possible to unequivocally infer the pattern of basal-body segregation from the flagellar number data.

**Generalized Somatic Segregation Defects in vfl-3**

Vfl-3 cells appear to possess a defect in accurately partitioning their contents at cytokinesis. Cell size is highly variable as compared to wild type (Fig. 1), and individual cells have been directly observed to yield daughters of unequal size. Consistent with a cell division defect, a small but significant fraction of cells are binucleate or bipyramidal. In light of the fact that the cleavage furrow in the dividing C. reinhardtii cell is lined with numerous microtubules which radiate from very near the basal bodies (17), we feel it likely that improper cleavage furrow positioning, and hence unequal partitioning of the cell's contents at division, represents a consequence of improper basal-body positioning.

**Inferences about Basal-Body and Striated Fiber Polarity Based on Symmetry Considerations**

*Chlamydomonas* flagella beat with the highly asymmetrical pattern typically exhibited by cilia. The two wave forms display a 180° rotational symmetry with respect to one another, resulting in a flagella-first swimming pattern. While the structural basis for flagellar beat asymmetry is not known, it is clear that in spite of its dominant ninefold rotational symmetry the flagellum has a number of elements which are not ninefold rotationally symmetric (43), and so it is safe to say that a structural polarity underlies the functional polarity expressed in the organelle's highly asymmetric beating.

Our results suggest the hypothesis that each basal body has an intrinsic polarity which determines the rotational orientation of the attached flagellum and hence the direction of the effective stroke. We reason as follows: the basal body shows ninefold rotational symmetry with respect to its major structural components, but is not ninefold symmetrical with respect to its striated fibers. If we hypothesize that the polarity associated with fiber attachment is determined by an intrinsic polarity in the basal body, and if we make two conservative assumptions—

(a) that the two basal bodies are structurally identical, and (b) that the striated fiber attachment sites on each basal body are unique—it follows from symmetry considerations alone (see Fig. 6) that the two basal bodies are 180° rotationally symmetric (and thereby show the same rotational symmetry as the two flagellar wave forms).

Symmetry considerations lead to several additional speculations about striated fiber length, orientation, and banding patterns. First, it should be noted that the sites of distal fiber attachment are equivalent on the two basal bodies. Thus if fiber assembly can initiate on one basal body it ought to be able to initiate on the other as well. This leads us to propose a model wherein complete fibers arise by the joining of two half fibers from neighboring basal bodies. If there is no neighboring basal body, as is the typical case in vfl-3, a half-length distal fiber phenotype should result. Apparent half distal fibers have in fact been observed many times in vfl-3 sections (for example Figs. 2e and 4f). The model also predicts that the banding pattern of the complete distal fiber should show apparent mirror-image symmetry about its midline, as it indeed does.

Unlike the single distal fiber, each of the two shorter proximal fibers does not connect geometrically equivalent sites on the two basal bodies. Here it can be imagined that one site serves to initiate fiber assembly and the other site serves to receive a fiber initiated from its partner basal body. This model predicts that proximal fibers will grow to full length on solitary basal bodies, and this does appear to be the case in vfl-3 (for example Fig. 4b and c). Furthermore, if proximal fiber assembly originates at only one of the two attachment sites on each basal body, an unpaired basal body should never carry two proximal fibers; this, too, has proved to be the rule.

**FIGURE 5** Abnormal rootlet microtubule arrangements in vfl-3. (A) Wild-type section showing the cruciate orientation of the rootlet microtubules which converge between the basal bodies. Both proximal striated fibers are visible as well. Collidine fixation. (B) Wild-type section similar to that in A. Note the two probasal bodies in their characteristic positions. Collidine fixation. (C) vfl-3 section showing microtubule rootlets oriented parallel to each other rather than converging between basal bodies. Collidine fixation. (D) vfl-3 section with microtubules criss-crossing near basal body. Electron-dense material of unknown identity appears to be associated with the microtubules. Collidine fixation. (E) Apparently normal microtubular rootlet associated with a solitary vfl-3 basal body. Collidine fixation. (F) vfl-3 section showing three mature basal bodies in cross section. Note microtubules criss-crossing between one basal-body pair, and an incomplete distal fiber pointing from one basal body towards the other. The third basal body appears free of rootlet associations, but possesses a striated projection. One probasal body is in a normal position relative to the basal-body pair and associated rootlet microtubules, but the other two show no association of association with rootlets, mature basal bodies or each other. Collidine fixation. (G) Basal body. (p) Probasal body. (pf) Proximal fiber. Bar, 500 nm. × 48,000.

**FIGURE 6** Cartoon view of basal body pair seen from above, showing the large distal fiber and the two smaller proximal fibers. Four radial coordinates—N, S, E, and W—are included to emphasize the rotational symmetry.

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Comparison to Other Mutants

We recently described a variable flagellar number mutation in the gene vfl-2 (19). The vfl-3 and vfl-2 phenotypes differ in several significant respects. (a) Cell size is normal in vfl-2. (b) Striated fibers of normal morphology are present in many vfl-2 cells (our unpublished micrographs). (c) Extra basal bodies are not present in vfl-2. It is thus clear that not all variable flagellar number mutants have striated fiber or cell division defects. Analysis of additional mutants should reveal whether or not such defects are rare or common features of the vfl phenotype.

The variable flagellar number mutant cyt-I which was described by Warr (41) is also quite different in phenotype from vfl-3. cyt-I cells are large, multi-lobed, and generally multinucleate. The cyt-I flagella exist in pairs, usually with one pair for each nucleus. The defect in cyt-I appears to concern the completion of cytokinesis, and there is no reason to expect it to have striated fiber defects.

There exists a mutant of the multicellular alga Volvox carteri which resembles vfl-3 in a number of respects. In this mutant, S-16 (25), cells divide abnormally to yield progeny of unequal size and of improper orientation with respect to the organism as a whole. As an apparent consequence of its defect in cytokinesis, embryogenesis in S-16 is grossly aberrant. Like vfl-3, cells of the mutant show a variable flagellar number phenotype (D. Kirk, personal communication).

Inferences about Striated Fiber Function

Because striated fibers are incomplete or absent in vfl-3, we can rule out an essential role for complete fibers in all functions which vfl-3 cells perform correctly. We therefore conclude that complete fibers are not required: (a) for the backward swimming response, (b) for mating, (c) to anchor the basal bodies in the cytoplasm so that they are not torn out by the force of flagellar beating, and (d) to maintain equality of flagellar length in multiply flagellated cells.

vfl-3 is clearly defective in regulation of flagellar number, proper placement of basal bodies in the cell, the establishment or maintenance of proper spatial orientation of flagellar wave forms, and accurate partitioning of the cytoplasm at cytokinesis. Striated fibers could be essential to any or all of these phenomena; alternatively, the vfl-3 phenotype could represent a defect in an underlying cellular organizing system about which we are not aware. However, an indirect argument, based on our failure to see complete fibers in vfl-3 sections, does suggest a primary defect in fiber assembly. In 1975 Goodenough and St. Clair (9) reported that in the basal-body defective mutant bald-2 complete distal fibers unattached to basal bodies were present in the basal-body region of the cell. Since bald-2 cells have only very rudimentary basal bodies their result suggests that complete fiber assembly can occur without basal bodies. If complete fibers can assemble independent of the basal bodies, it becomes difficult to explain the absence of complete fibers in vfl-3 on the basis of a defect in basal-body structure or placement, making the possibility of a primary defect in fiber assembly more attractive.

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