Distinct Mutants of Retrograde Intraflagellar Transport (IFT) Share Similar Morphological and Molecular Defects

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Abstract. A microtubule-based transport of protein complexes, which is bidirectional and occurs between the space surrounding the basal bodies and the distal part of Chlamydomonas flagella, is referred to as intraflagellar transport (IFT). The IFT involves molecular motors and particles that consist of 17S protein complexes. To identify the function of different components of the IFT machinery, we isolated and characterized four temperature-sensitive (ts) mutants of flagellar assembly that represent the loci FLA15, FLA16, and FLA17. These mutants were selected among other ts mutants of flagellar assembly because they displayed a characteristic bulge of the flagellar membrane as a non-conditional phenotype. Each of these mutants was significantly defective for the retrograde velocity of particles and the frequency of bidirectional transport but not for the anterograde velocity of particles, as revealed by a novel method of analysis of IFT that allows tracking of single particles in a sequence of video images. Furthermore, each mutant was defective for the same four subunits of a 17S complex that was identified earlier as the IFT complex A. The occurrence of the same set of phenotypes, as the result of a mutation in any one of three loci, suggests the hypothesis that complex A is a portion of the IFT particles specifically involved in retrograde intraflagellar movement.

Key words: Chlamydomonas • temperature-sensitive mutants • intraflagellar particles • video microscopy • retrograde transport

Transport of proteins from the site of synthesis to the site of function in eukaryotic cells often occurs along frameworks of microtubules with the participation of molecular motors and auxiliary molecules that link the motors to the cargo. These molecular machineries also operate in specific compartments of polarized cells, such as cilia, flagella, and axons, where they serve several functions: among others, they release a cargo at the distal part of the compartment and move back to the cell body to be recycled. Retrograde transport often requires molecular motors that differ from those working in anterograde transport.

To dissect the complexity of a microtubule-based system that transports proteins within a specific cellular compartment, we turned to the study of temperature-sensitive (ts) mutants of Chlamydomonas reinhardtii that are defective in the assembly of flagella (Huang et al., 1977). Chlamydomonas flagella contain a machinery that transports protein complexes in both directions and operates between outer doublet microtubules and the flagellar membrane (Kozminski et al., 1995). This intraflagellar transport (IFT) of protein particles was discovered by video-enhanced differential interference contrast microscopy (Kozminski et al., 1993) and currently is being analyzed at the molecular level (Piperno and Mead, 1997; Cole et al., 1998; Pazour et al., 1998).

The IFT requires KHP1FLA10 (Walther et al., 1994), the heavy chain of an heterotrimeric kinesin II (Sholey, 1996), for anterograde transport (Kozminski et al., 1995; Cole et al., 1998) and LC8FLA14, a light chain of axonemal and cytoplasmic dyneins (King et al., 1996), for retrograde transport of a cargo (Pazour et al., 1998). The cargo consists of large protein particles that are not membrane-bound (Kozminski et al., 1995) and are composed of at least 15 polypeptides comprising two 16S complexes, referred to as IFT complexes A and B (Cole et al., 1998).

Following our studies on the structure of inner dynein complexes and the frequency of bidirectional transport but not for the anterograde velocity of particles, as revealed by a novel method of analysis of IFT that allows tracking of single particles in a sequence of video images. Furthermore, each mutant was defective for the same four subunits of a 17S complex that was identified earlier as the IFT complex A. The occurrence of the same set of phenotypes, as the result of a mutation in any one of three loci, suggests the hypothesis that complex A is a portion of the IFT particles specifically involved in retrograde intraflagellar movement.

Key words: Chlamydomonas • temperature-sensitive mutants • intraflagellar particles • video microscopy • retrograde transport

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1. Abbreviations used in this paper: IFT, intraflagellar transport; MNNG, N-methyl-N′-nitro-N′-nitrosoguanidine; ts, temperature-sensitive.
arms, we analyzed the transport of p28, an inner dynein arm light chain, within the flagella of *ida4* (Piperno et al., 1996), a null mutant of p28 (LeDizet and Piperno, 1995a). We found that p28 required KHP1*FLA10* to reach the distal part of flagella where it bound to outer doublet microtubules in a concentration gradient from distal to proximal part of the axoneme. This evidence suggested that the transport of precursors of inner dynein arms to their binding site within the axoneme involved a molecular motor and not passive diffusion (Piperno et al., 1996).

We also found that a cytoplasmic 17S complex of polypeptides binds p28 in substoichiometric amounts and requires KHP1*FLA10* to be present in flagella (Piperno and Mead, 1997). The 17S complex represents 2–4% of the mass of flagellar proteins and is composed of at least 13 different subunits, nine of which turned over 37 times faster than axonemal dyneins (Piperno and Mead, 1997). This evidence suggested the following hypotheses: the 17S complex functions as a carrier of precursors of inner dynein arms and consists of two modules that differ in turn-over (Piperno and Mead, 1997). Furthermore, the 17S complex is probably identical to the ensemble of the IFT complexes A and B recently described (Cole et al., 1998).

To identify the function of different parts of this machinery of transport, we intended to isolate ts mutants of flagellar assembly with a defect in a specific aspect of the IFT, such as the anterograde motion, the retrograde motion, or the frequency of transport. To this purpose we analyzed ts mutants of flagellar assembly with phenotypes similar to that of *fla10* because the IFT is inhibited at the restrictive temperature in flagella of *fla10* (Kozminski et al., 1995), and KHP1*FLA10* and the 17S complex are parts of the same transport system (Kozminski et al., 1995; Piperno and Mead, 1997; Cole et al., 1998). In an initial phase of this study, we observed ts mutants isolated by others (Huang et al., 1977; Adams et al., 1982). However, some of them lost the original ts phenotype while others could have accumulated additional mutations. Therefore, instead of characterizing them again we isolated a new set of ts mutants of flagellar assembly by a procedure that is similar to that previously described (Huang et al., 1977; Adams et al., 1982). Furthermore, to detect subtle changes in a specific property of the IFT, we developed a method of quantitative analysis of the IFT that allows unambiguous identification of single particles and, therefore, accurate measurement of anterograde and retrograde velocity of the particles.

Here we describe four of the new ts mutants of flagellar assembly, which we selected from among the others because they have an additional phenotype, namely a distinct bulge of the flagellar membrane. These four mutants represent three loci referred to as *FLA15*, *FLA16*, and *FLA17*. They have flagella as long as those of a wild-type strain at permissive temperature and accumulated amorphous material in the flagellar bulge at both permissive and restrictive temperatures. The presence of the flagellar bulge in each mutant was correlated with a decrease of both velocity of retrograde IFT and frequency of bidirectional transport. In addition, each mutant was defective for the same four subunits of a 17S complex, three of which were identified earlier as subunits of the IFT complex A. The identification of the gene products of *FLA15*, *FLA16*, and *FLA17* should reveal whether IFT complex A is required for retrograde transport of protein particles within flagella.

**Materials and Methods**

**Cell Culture**

Wild-type cells (137°) used for mutagenesis were grown in liquid medium (Sager and Granick, 1953) as modified in Huang et al. (1977) for 3 d under intense light at 25°C. Wild-type cells (137°) or mutants used for biochemical or phenotypic analyses were grown on solid medium for 3 d under intense light at 25°C and 1 d in the dark at 21°C. Cells used for phenotypic analysis were cultured in liquid medium for at least 12 h under light at 21°C.

Proteins were extracted from wild-type cells (137°) or mutants that were grown at a steady state on modified solid medium supplemented with [35S]sulfuric acid (Luck et al., 1977).

The wild-type strains 137 and *fla10*-1 are from the collection of Dr. David Luck (Rockefeller University, New York).

**Screening of Temperature-sensitive Mutants for Flagellar Assembly**

Vegetative wild-type cells (137°, 10⁶ cells/ml) in 50 ml of 0.02 M citrate buffer, pH 5, were exposed to 1, 5, or 10 mg/ml of N-methyl-N-nitro-N-nitrosoguanidine (MNNG) for 30 min at 25°C in the dark. After two washes in medium, cells (10⁶ cells/ml) were cultured for 24 h in light at 21°C. Survival was 71, 57, and 45% after the exposure to 1, 5, or 10 mg/ml of MNNG, respectively. Screening of temperature-sensitive mutants for flagellar assembly was achieved by three repeats of the following two-step cycle: we discarded cells regenerating their flagella and swimming at the restrictive temperature of 32°C, and we also discarded cells not swimming at the permissive temperature of 21°C. At the first step of the cycle, cells were deflagellated by pH shock, pelleted by centrifugation at 1,000 rpm (in a model GH-3.7 rotor; Beckman Instruments, Fullerton, CA), resuspended in 30 ml of medium, and exposed 3 h at 32°C in light before being separated in four aliquots, one of which was enriched in cells that were sedimenting. Sedimenting cells were cultured overnight at 21°C in light. After this incubation, the upper 70% of the culture was removed and kept for the next screening at 32°C. The volumes of the medium added to the cells before the second and third exposure at 32°C were 5 and 3 ml, respectively. We cultured the remaining cells in solid medium, picked single colonies, and cultured them in 200 μl of liquid medium. We selected clones that lost the majority of their flagella within 4 h of exposure at 32°C. We retained 11 and 5 clones of mutants from cells exposed to 1 or 5 mg/ml of MNNG, respectively. Some of the clones could derive from the same mutant that divided during the process of screening.

Mutants in four distinct clones had the same phenotype: a bulge of the flagellar membrane. They were crossed with wild-type cells (137°) to verify that they represented a single mutation. They also were crossed to each other and to *fla10* to determine whether they represented different loci. Complementation tests were performed by tetrad analysis (Harris, 1989). Three of the four mutants recombined and generated wild-type cells in reciprocal crosses. In contrast, two mutants did not recombine in 204 crosses. Each of the three mutants that recombined also recombined with *fla10*. Therefore, the four mutants were referred to as *fla15, fla16, fla17-1,* and *fla17-2* to follow the nomenclature previously adopted for temperature-sensitive mutants of flagellar assembly (Adams et al., 1982; Harris, 1989). Whether any one of these mutants is an allele of a *fla* mutant other than *fla10* remains to be determined.

Recombinant strains between *pflA*, a paralyzed flagella mutant (Adams et al., 1981), and *fla10, fla15, fla16*, and *fla17-1* were isolated to carry out the analysis by video microscopy. They had straight, immotile flagella and were temperature-sensitive for flagellar assembly.

**Optical and Video Microscopy**

Phenotypic analysis of mutants was performed on cells fixed in 2% glutaraldehyde, 0.01 M sodium phosphate buffer, pH 6.9. Phase contrast microscopy was performed with a microscope (model Axioskop; Carl Zeiss, Inc., Thornwood, NY) connected to a CCD camera (SenSys, model KAF 1400; Photometrics, Tucson, AZ) through a ×3 intermediate lens.

Differential interference contrast microscopic observation of intraflagellar transport of particles in living cells was carried out using a microscope (model Axiosvert 35; Carl Zeiss, Inc.) with a 1.4 NA condenser, a
of the period. In the unprocessed data. Examples of composite plots of linescans from scans in which the contributions of both background and noise were sustained uncorrelated signal arising from noise in the data, whereas the first, dimensions vectors constitutes a matrix called singular values, and two sets of orthogonal eigenvectors (Golub and

tional linescans. The second set of eigenvectors forms a matrix \( U \) with dimensions \( n_s \times n_s \), where \( n_s \) is the number of pixels in each linescan and is formed by \( n_l \) component \( \text{spectra} \) or waveforms, which carry information related to the shape of the experimental linescans. The second set of eigenvectors forms a matrix \( V \) with dimensions \( n_s \times l \), where \( l \) is the number of linescans and carries information about the fractional contribution of each of the \( n_s \) components to each of the linescans. Finally, each of the \( n_s \) singular values in the diagonal matrix \( S \) measures the weight or contribution of the respective component to the ensemble of linescans. Multiplication of the three matrices, \( U^T V \), where \( V^T \) represents the transpose of \( V \), using the complete set of eigenvalues and eigenvectors yielded a perfect reproduction of the original ensemble of linescans, including background and noise. However, inspection of the \( U \) eigenvectors revealed that components beyond the first 10–20 contained uncorrelated signal arising from noise in the data, whereas the first, and sometimes up to the third, eigenvector contained the signal contribution of the structured background. Matrix multiplication using the subset of intermediate components yielded a reconstructed ensemble of linescans in which the contributions of both background and noise were suppressed, as a result of this procedure not allowing the unambiguous identification of moving particles but also eliminated the distortions in the time component of the time/intensity dimension caused by variations in the signal due to fluctuations in light intensity and other artifacts present in the unprocessed data. Examples of composite plots of linescans from the flagellum of \( p/15 \), a mutant with straight and immotile flagella that was used as a reference strain, and of \( f15k \), one of the recombinant strains characterized in this study, are shown in Fig. 4. The velocity of each particle was calculated from the slope of a line drawn manually along each of the diagonal ridges.

A lower limit for the value of the frequency of intraflagellar transport, expressed in particles/second, was estimated from the ratio of the total number of particles detected to the total observation time, equal to the number of linescans divided by thirty, the video frame rate. The frequency of retrograde transport probably was underestimated as a result of the low contrast generated by particles moving in that direction, whose features in the composite plots were therefore at the limit of detection.

**Electron Microscopy**

Cells were fixed in 3% glutaraldehyde in 10 mM Hepes buffer, pH 7.2, for 2 h at 4°C. The fixed cells were then washed three times for 10 min each in 10 mM Hepes buffer, pH 7.2, and postfixed in 1% OsO\(_4\) plus 0.8% K\(\text{Fe(CN)}_3\), in 4 mM phosphate buffer, pH 7.2, for 30 min at 4°C (McDonald, 1984). The cells were then washed with water for 10 min, stained with 0.15% tannic acid for 1 min, again washed with water for 10 min, and stained en bloc in 2% uranyl acetate for 1 h at room temperature in the dark. After staining, the cells were washed with water for 10 min and then dehydrated through a graded ethanol series (70, 80, 90, 100, and 100%, 3 min in each stage). The cells were then taken through a transition of propylene oxide for 15 min and 1:1 propylene oxide/epon/araldite resin for 1 h at room temperature on a rotator. They were then infiltrated with epoxi/ araldite resin overnight at room temperature on a rotator and, finally, embedded in a fresh batch of epoxi/araldite resin. The resin was polymerized at 68°C for 2 d.

Silver sections were cut on an Ultracut E ultramicrotome (Reichert-Jung, Vienna, Austria), collected on 200 mesh grids, and stained with uranyl acetate for 20 min and lead citrate for 2 min. Examination and photography of sections was carried out using a transmission electron microscope (model H-7000; Hitachi, Tokyo, Japan).

**Isolation of the 17S Complex**

Flagella were separated from the cell bodies by a pH shock method (Huang et al., 1979). After ultrasonic disruption of flagella, soluble proteins were separated from the insoluble residue by centrifugation at 14,000 rpm in a table top minicentrifuge (model 5402; Eppendorf, Madison, WI). Protein fractions containing the 17S complexes were isolated by sedimentation in a 5–20% sucrose gradient (Piperno and Mead, 1997). Sedimentation standards were 21S and 18S Chlamydomonas dynesins (Piperno et al., 1990) that were sedimented either with the flagellar extracts or in a parallel gradient.

Electrophoresis of Polypeptides

One- and two-dimensional electrophoresis of \( ^{35}\text{S}\)-labeled polypeptides were carried out as described (Piperno, 1995) with the following two modifications. SDS gels were made of a 4–11% polyacrylamide gradient. Isoelectric focusing was carried out either at 1.4 mA for 14 h or 1.6 mA for 16 h. Quantitative analysis of the isoelectric point was carried out under the second set of conditions, which allowed the subunits of the 17S complex to reach their isoelectric point. Determination of pH of the gel containing polypeptides at their isoelectric point was carried out directly in suspensions of \( 1 \times 0.1 \) cm gel slices in 2 ml of water.

Evaluation of apparent molecular weight of polypeptides was performed by the use of molecular weight standards (Pharmacia Biotech lot 7007615011 and GIBCO-BRL lot KB9418). The values reported here for 13 subunits of the 17S complex were: (1) 189,000; (2) 148,000; (3) 127,000; (4) 86,000; (5) 81,000; (6) 71,000; (7) 69,000; (8) 65,000; (9) 56,000; (10) 47,000; (11) 26,000; (12) 21,000; and (13) 19,000. These values are probably more accurate than those reported before (Piperno and Mead, 1997). The previous set of data was obtained by prestained molecular weight standards (SDS-7B lot 125H9408; Sigma Chemical Co., St. Louis, MO) that yielded nonreproducible results.

**Western Blots**

Western blot analyses of axonemal proteins and subunits of the 17S were
performed as described (LeDizet and Piperno, 1995b). To detect the presence of inner dynein arms, outer dynein arms, and central pair complexes in flagella from fla15, fla16, and fla17-1, we used antibodies specific for p28 (LeDizet and Piperno, 1995b), γ-chain of outer dynein arms (Piperno et al., 1996), and PF16 gene product (Smith and Lefebvre, 1996), respectively. Equal amounts of axonemal proteins from each mutant and a wild-type strain bound each antibody in similar amounts. To identify subunit 2 of the 17S complex (Piperno and Mead, 1997) with p144–p139 subunits of complex A (Cole et al., 1998), we used monoclonal antibody 139.1 (Cole et al., 1998).

Results

Temperature-sensitive Mutants of Flagellar Assembly with a Nonconditional Phenotype: A Bulge of the Flagellar Membrane

To identify ts mutants of flagellar assembly that are defective in the IFT, we generated mutants similar to fla10, a ts mutant of a motor involved in IFT (Walther et al., 1994; Kozminski et al., 1995; Cole et al., 1998). After mutagenesis and screening, we isolated 16 strains that lost or did not regenerate their flagella after 4 h of exposure at the restrictive temperature of 32°C. Four strains also had a nonconditional phenotype: a distinct bulge of the flagellar membrane. We selected these four strains for further analysis because such a deformation of the flagellar membrane could derive from disruptions in the transport of protein particles that occurs between outer doublet microtubules and the flagellar membrane (Kozminski et al., 1995).

We performed a complementation test on the four strains and found that three of them consisted of mutants representing different loci. Therefore, we will refer to the four mutants as fla15, fla16, fla17-1, and fla17-2, following the designation used for mutants of flagellar assembly (Adams et al., 1982). The strain fla17-2 was not used as extensively as fla17-1 in the following studies because it may be a clone of the same mutation in FLA17, as explained in Materials and Methods.

Anomalous assembly of flagella in fla15, fla16, and fla17-1 was evident at both permissive and restrictive temperatures. At the permissive temperature, we detected a defect in flagellar assembly by measuring the rate of regeneration of flagella. Increasing delay in the regeneration of flagella was observed in the following order: fla15 > fla16 > fla17-1 > wild-type (Fig. 1). However, the final length of the regenerated flagella in all mutants was similar to that of the wild-type strain (Fig. 1). Therefore, mutations in FLA15, FLA16, and FLA17 affected the rate of assembly of flagella, not the control of flagellar length.

A 4-h exposure of each mutant to the restrictive temperature led to extensive loss of flagella similar to that observed in fla10 (Table I). The flagella remaining in mutants following a 4-h exposure to the restrictive temperature were ~50% shorter than in a wild-type strain (Table I). Prolonged exposures (6–8 h) to the restrictive temperature caused complete retraction of flagella in the mutant cells.

In contrast to fla10, which has wild-type–like flagella at the permissive temperature, fla15, fla16, fla17-1, and fla17-2 have a bulge positioned randomly along the length of one flagellum at both permissive and restrictive temperatures but not in isolated flagella. Occasionally, the bulge was on both flagella or was absent. Its short diameter was ~0.4 μm, near the limit of the optical microscope resolution. The bulge was immotile at least over the time course of microscopic observation, 5–10 min. Examples of bulges on the flagellar membrane of fla15 are shown in Fig. 2, as detected by differential interference and phase contrast microscopy after fixation in glutaraldehyde. The other mutants were indistinguishable from fla15 by this criterion.

Electron microscopy of fla15, fla16, and fla17-1 revealed that the bulge of the flagellar membrane contained amor-
To determine whether IFT is inhibited in fla15, fla16, and fla17-1, as it is in fla10 at the restrictive temperature (Kozminski et al., 1995), we observed flagella in vivo by video-enhanced differential interference contrast microscopy. For this purpose, we isolated recombinants between the mutants and p/f15, a paralyzed flagella mutant, to perform the microscopic analysis on immotile and straight flagella. In each recombinant, fla10p/f15, fla15p/f15, fla16p/f15, and fla17-1p/f15 (Table II). In contrast, the velocity of retrograde transport in fla15p/f15, fla16p/f15, and fla17-1p/f15 was significantly lower than that in p/f15 and fla10p/f15 (Table II). Results of these comparisons between each recombinant and p/f15 are significant at a confidence level of $P < 0.001$, as determined by standard Student’s $t$ test.

At the restrictive temperature, the velocity of retrograde transport of particles did not change significantly in fla16p/f15 and fla17-1p/f15, whereas retrograde transport was undetectable in the majority of fla10p/f15 flagella and in all fla15p/f15 flagella.

The representative plots of singular value decomposition (SVD) processed linescans measured from p/f15 and fla15p/f15 at the permissive temperature (Fig. 4) show velocity and the frequency of IFT of protein particles in both anterograde and retrograde directions was carried out by a new method described in Materials and Methods. The analysis required both the scanning of light intensity along a flagellum in each video frame and the composition of linescans. In these composite plots, moving particles appear as diagonal ridges or streaks, whose slope is proportional to their velocity. Examples of these streaks originating from particles undergoing anterograde and retrograde transport were colored in red and green, respectively, in the composite plots generated with p/f15 and fla15p/f15 (Fig. 4). Particles undergoing anterograde transport generated a stronger light contrast than particles undergoing retrograde transport. Therefore, these particles differed in optical properties from those moving in the opposite direction.

At the permissive temperature, the velocity of anterograde movement of particles in p/f15 was similar to that in fla10p/f15, fla15p/f15, fla16p/f15, and fla17-1p/f15 (Table II). In contrast, the velocity of retrograde transport in fla15p/f15, fla16p/f15, and fla17-1p/f15 was significantly lower than that in p/f15 and fla10p/f15 (Table II). Results of these comparisons between each recombinant and p/f15 are significant at a confidence level of $P < 0.001$, as determined by standard Student’s $t$ test.

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frequency of bidirectional transport is decreased in fla15 pf15. The frequencies of bidirectional transport in fla15 pf15, fla16pf15, and fla17-1pf15 were 24, 66, and 51% of that of pf15, respectively. In contrast, the frequency of bidirectional IFT of fla10pf15 was similar to that of pf15 under the same condition.

In summary, fla15, fla16, and fla17-1 differ from fla10 for the presence of a bulge on their flagella, a lower veloc-

Figure 3. The deformation of the flagellar shape caused by fla15 was correlated with the accumulation of cytoplasmic matrix. (a and b) Electron micrographs. (a) Longitudinal section cut through the central pair microtubules. The cytoplasm accumulated between the outer doublet microtubules and the flagellar membrane in a section of the axoneme. (b) Longitudinal section cut through the cytoplasm accumulated in the bulge of the flagellar membrane. Bar, 0.1 μm.

Figure 4. Representative illustration of intraflagellar transport of particles occurring at permissive temperature. Composite plots of longitudinal linescans of light intensity along flagella of (a) pf15 and (b) fla15pf15. One linescan from the proximal to the distal part of the flagellum was measured for each successive image in a video sequence obtained at a rate of 30 frames/s. The ensemble of linescans was then subjected to singular value decomposition and reconstructed as described in the text. The processed linescans were stacked and displayed so that the origin of the x axis corresponds to both the first linescan of a sequence and the proximal part of the flagellum. The distance on the y axis was measured relative to the proximal part of the flagellum. Particles undergoing anterograde or retrograde transport are identifiable as ridges with rightward and leftward slopes, respectively. (a and b) Examples of red and green ridges represent particles undergoing anterograde and retrograde transport, respectively.
radiograms of 35S-labeled polypeptides contained in sucrose gra-

mimenting fractions from the cytoplasmic matrix of flagella. Auto-

within flagella (Kozminski et al., 1995; Piperno and Mead,

the 17S complex are parts of the same transport system

indicate the rest of the subunits of 17S complexes.

ity of retrograde IFT and a lower frequency of bidirec-

tional transport. These phenotypes are nonconditional, in

contrast to the lack of flagellar assembly or flagellar regen-

eration that is expressed at the restrictive temperature in

every mutant including fla10.

fla15, fla16, and fla17-1 Are Deficient for the Same

Subunits of the IFT Complex A

The FLA10 product is KHP1^{FLA10} (Walther et al., 1994), a

subunit of a kinesin II (Scholey, 1996), and KHP1^{FLA10} and

the 17S complex are parts of the same transport system

within flagella (Kozminski et al., 1995; Piperno and Mead,

1997; Cole et al., 1998). To determine whether the 17S

complex was defective in fla15, fla16, and fla17-1, we ex-

tracted it from flagella of each mutant grown at the per-

missive temperature. Under this condition, the concen-

tration of the 17S complex in flagella of the mutants was

similar to that of a wild-type strain. In contrast, the con-

centration of the 17S complex decreased several folds in

flagella of mutants exposed to the restrictive temperature.

The sedimentation profiles of 35S-labeled flagellar pro-

teins from mutant strains were indistinguishable from that

of a wild-type strain. However, the electrophoreto-

grams of protein fractions from the sucrose gradients revealed

that the 17S complexes from each mutant were deficient to

a different extent for the same two polypeptides with ap-

parent molecular weights of 148,000 and 127,000. The elec-

trophoretic bands of defective polypeptides from fla15 are

indicated by asterisks located between lanes 9 and 10 of

the electrophoretograms (Fig. 5 b). The other subunits of

17S complexes from fla15 and a wild-type strain are indi-

cated by lines (Fig. 5, b and a, respectively).

The polypeptides deficient from flagella of fla15, fla16,

and fla17-1 had apparent molecular weights similar to

those of two subunits of complex A (Cole et al., 1998) (Ta-

ble III). Furthermore, in the absence of these polypepti-

des the sedimentation properties of the remaining subunits of

the 17S complexes were not altered. Therefore, polypep-

tides that are defective from each mutant may form a dis-

tinct 17S complex in a wild-type strain, and this complex

may be the IFT complex A (Cole et al., 1998).

To test these hypotheses and determine whether other

deficiencies occur in each mutant, we separated the pol-

ypeptides of 17S sedimenting fractions from wild-type and

mutant strains by high-resolution one- and two dimen-

sional gel electrophoresis. In these experiments, we iden-

tified the 13 subunits of the two 17S complexes by their ap-

parent molecular weights. These subunits are indicated by

lines and progressive numbers in Fig. 6.

Flagella of fla15, fla16, and fla17-1 contained reduced

amounts of subunits 2 and 3 relative to flagella of a wild-
type strain (Fig. 6). In addition, flagella of fla15 lacked a

polypeptide of apparent molecular weight 43,000, indicated

d by a line between a and b in Fig. 6. This polypeptide

was not previously identified as a subunit of the 17S com-
p lex (Piperno and Mead, 1997) or complexes A and B
(Cole et al., 1998). In contrast, flagella of fla16 and fla17-1
had reduced amounts of the 43,000 polypeptide and con-
tained an additional polypeptide, apparent molecular

weight 135,000, indicated by a dot between the second and

third lanes in Fig. 6 b. This last polypeptide is absent in fla-
gella of fla15 and of the wild-type (Fig. 6 a).

Comparison of two-dimensional electrophoreto-
grams of

| Strain | Anterograde | Retrograde |
|--------|-------------|------------|
| pf15   | 0.2 (SD: 0.3; n: 43) | 3.9 (SD: 0.6; n: 32) |
| pf15fla10 | 1.8 (SD: 0.2; n: 64) | 4.2 (SD: 0.6; n: 81) |
| pf15fla15 | 2.1 (SD: 0.3; n: 47) | 2.9 (SD: 0.5; n: 17) |
| pf15fla16 | 2.1 (SD: 0.2; n: 104) | 2.7 (SD: 0.6; n: 40) |
| pf15fla17-1 | 1.9 (SD: 0.3; n: 121) | 2.3 (SD: 0.4; n: 36) |

n, number of particles analyzed.

Figure 5. fla15 was defective for two polypeptides of the 17S sed-
imentong fractions from the cytoplasmic matrix of flagella. Auto-
radiograms of 35S-labeled polypeptides contained in sucrose gra-
dient fractions 6–16 after separation by gel electrophoresis.
Molecular weight standards are indicated on the left. (a) Proteins
from a wild-type strain. Lines between lanes 9 and 10 indicate the
presence of components 1–13 of the two 17S complexes. (b) Pro-
teins from fla15. Asterisks between lanes 9 and 10 indicate the
position of the two polypeptides that are deficient in fla15. Lines
indicate the rest of the subunits of 17S complexes.

Table II. Velocity of Intraflagellar Particles

| Strain | Anterograde | Retrograde |
|--------|-------------|------------|
| pf15   | 0.2 (SD: 0.3; n: 43) | 3.9 (SD: 0.6; n: 32) |
| pf15fla10 | 1.8 (SD: 0.2; n: 64) | 4.2 (SD: 0.6; n: 81) |
| pf15fla15 | 2.1 (SD: 0.3; n: 47) | 2.9 (SD: 0.5; n: 17) |
| pf15fla16 | 2.1 (SD: 0.2; n: 104) | 2.7 (SD: 0.6; n: 40) |
| pf15fla17-1 | 1.9 (SD: 0.3; n: 121) | 2.3 (SD: 0.4; n: 36) |

n, number of particles analyzed.

Table III. Polypeptide Composition of IFT Complex A from a

Wild-Type Strain

| Cole et al., 1998 | This study |
|------------------|-----------|
| Apparent molecular weight | pI | Apparent molecular weight | pI |
| 144,000 | 5.7–5.8 | 148,000 | 6.5 |
| 140,000 | 6.0 | 148,000 | 6.0 |
| 139,000 | 5.9 | 127,000 | 6.3 |
| 122,000 | 5.8–6.0 | 43,000 | 4.5 |

Piperno et al. Mutants of Retrograde Intraflagellar Transport

1597
polypeptides from \textit{fla15} and a wild-type strain revealed that the mutant lacked the majority of two polypeptides of apparent molecular weight 148,000, as well as polypeptides of molecular weights 127,000 and 43,000, respectively (Table III). These four polypeptides are indicated by oblique lines in the maps of polypeptides from wild-type and \textit{fla15} (Fig. 7, a and b). Two-dimensional maps of polypeptides from \textit{fla17-1} had reduced amounts of the same four polypeptides defective in \textit{fla15} (Fig. 7, c and b). The new 135,000 apparent molecular weight polypeptide present in \textit{fla17-1} is indicated by an arrowhead in Fig. 7 c. The two-dimensional map of polypeptides from \textit{fla16} was indistinguishable from that of \textit{fla17-1}.

Three of the polypeptides deficient in \textit{fla15}, \textit{fla16}, and \textit{fla17-1} were identified as subunits of IFT complex A on the basis of their isoelectric points and apparent molecular weight (Table III). The fourth polypeptide of 43,000 molecular weight that is defective in each mutant could be a previously undetected subunit of IFT complex A.

To determine whether these four polypeptides remain associated in a complex after chromatography and exposure to a high concentration of salt, we performed one- and two-dimensional electrophoresis of wild-type proteins that were eluted by a gradient of NaCl at pH 6.8 from a DEAE-Sepharose column. At least three protein fractions eluted at 0.29–0.30 M NaCl had identical compositions, as assessed by one-dimensional electrophoresis. Polypeptides eluted in that range of NaCl molarity formed a map (Fig. 8) that was simpler than that shown in Fig. 7 a but still included the four polypeptides that were deficient from flagella of \textit{fla15}, \textit{fla16}, and \textit{fla17-1} (Fig. 8), indicated by oblique lines. Therefore, these polypeptides behaved as subunits of a complex that was stable after two subsequent procedures of protein purification.

In addition to these four putative subunits of complex A, 30 other polypeptides were eluted from the column at 0.3 M NaCl and were detectable in the map (Fig. 8). After column chromatography, they remained in proportions similar to those recorded after sedimentation in sucrose gradient (Fig. 7 a). Only a subset of these polypeptides were tentatively identified by their apparent molecular weights and isoelectric points as subunits of the IFT complex B, previously described (Cole et al., 1998). Therefore, the molecular composition of complex B may include a larger number of subunits than that reported previously (Cole et al., 1998).

**Discussion**

To identify the function of components of the machinery that carries out the IFT of particles in \textit{Chlamydomonas}, we have isolated and characterized four ts mutants of flagellar assembly that represent three loci and are defective in the same four characteristics: the flagellar shape, the retrograde velocity of IFT, the frequency of bidirectional IFT, and the concentration of IFT complex A in flagella. This evidence suggests the hypothesis that the complex A is the component of the IFT particles involved in the retrograde transport of proteins within flagella. We also identified a new subunit of the IFT complex A and indicated that the molecular composition of both complex A and complex B includes a number of subunits larger than that reported previously. Finally, to perform this study, we devised a new method for the quantitative analysis of the IFT that measures the velocity as well as the frequency of bidirectional transport and the optical properties of IFT particles. This method will be valuable to determine whether other ts mutants of flagellar assembly are defective in IFT and to identify the structural differences existing between particles moving in opposite directions.

The four mutants of retrograde IFT, \textit{fla15}, \textit{fla16}, \textit{fla17-1}, and \textit{fla17-2}, were selected among 16 strains defective in flagellar assembly for the presence of a characteristic bulge on their flagella. These mutants, like \textit{fla10}, normally have long flagella at the permissive temperature and disassemble or do not regenerate flagella at the restrictive temperature. They are ts in the concentration of complex A in flagella and probably for this reason they are ts in the assembly of flagella. In addition, they are nonconditional for both the occurrence of a bulge of the flagellar membrane and a decrease in the velocity of the retrograde IFT. Four other ts mutants of flagellar assembly, which were isolated in parallel to \textit{fla15}, \textit{fla16}, \textit{fla17-1}, and \textit{fla17-2}, did not
present morphological defects of the flagellar membrane nor were they defective in the intraflagellar concentration of complex A. Therefore, the correlation between the phenotype of the bulge and the deficiency in complex A defines a specific subset of ts mutants of flagellar assembly.

The bulge of the flagellar membrane in fla15, fla16, fla17-1, and fla17-2 has both a distinct morphology and rare occurrence. In contrast, deformations of the flagellar shape were various and extensive in fla14, a short flagella mutant of LC8FLA14 (Pazour et al., 1998), and in sea urchin embryos, where ciliary assembly was inhibited (Morris and Scholey, 1997). Furthermore, the bulge of fla15, fla16, fla17-1, and fla17-2 occurred once per flagellum and not in every cell and flagellum. This evidence suggested that this phenotype required both a mutation in FLA15 or FLA16, or FLA17 and an unknown defect that occurs once per flagellum and only in a fraction of them.

The deformation of fla15, fla16, fla17-1, and fla17-2 flagella also is distinct from that found in fla14 for its content. The first was filled with amorphous material that is indistinguishable from the cytoplasm. In contrast, the second contained stacked structures connected to both outer doublet microtubules and the membrane. This material was tentatively identified as the ensemble of complexes A and B (Pazour et al., 1998).

In addition to this morphological defect, we observed a decrease in the velocity of the retrograde IFT in fla15, fla16, and fla17-1 by a new method of quantitative analysis of the IFT that we have developed. Previous analyses of quantitative aspects of IFT were performed by measuring the relative position of individual particles in a series of still-frame video images. This approach was hampered by both the difficulty of identifying the same particle in each frame and the difficulty of detecting particles undergoing retrograde transport. The approach was also laborious because it involved manual processing of individual images. In contrast, the procedure that we have described here involved automatic collection of data from video image sequences lasting over 10-s intervals. The processed linescan data allowed the unambiguous identification of single particles undergoing both anterograde and retrograde transport and the determination of the frequency of bidirectional transport. The numerical data also confirmed the visual perception that the particles undergoing anterograde transport are different in structure and/or size from the particles undergoing retrograde transport. Finally, the use of this procedure allowed us to detect significant changes in the velocity of retrograde IFT in fla15, fla16, and fla17-1.

The occurrence of both the bulge of the flagellar membrane and the substantially slower velocity of retrograde IFT in each mutant was correlated also with a defect in the concentration of the IFT complex A within flagella. Our identification of the flagellar polypeptides deficient in each mutant as subunits of complex A was based on the values of apparent molecular weight and isoelectric point of these polypeptides and on the sedimentation coefficient of the complex that they form. Although the values of these parameters were not identical to those published by others, they were sufficiently close to allow the identification of IFT complex A. We recently confirmed that the electrophoretic band including the two 148,000 molecular
weight subunits of the 17S complex corresponds to the band including the three p144–p139 subunits of complex A because both bind the monoclonal antibody 139.1 (Cole et al., 1998).

Two other differences between the previous and present characterization of complex A are as follows: first, the IFT complex A characterized here included two rather than three subunits of molecular weight in the 144,000–139,000 range, as reported previously (Cole et al., 1998). Second, a polypeptide of 43,000 apparent molecular weight was deficient in each mutant but was not identified previously as a subunit of either complex A (Cole et al., 1998) or the 17S complex (Piperno and Mead, 1997). These discrepancies could result from differences in the methods of purification and detection of the protein complexes used in these studies. Thus, the third subunit of complex A in the 144,000–139,000 molecular weight range was identified in flagellar extracts that were obtained by exposure to detergent (Cole et al., 1998) rather than by ultrasonic treatment as reported here. Furthermore, the polypeptide of 43,000 molecular weight was identified by autoradiography of 35S-labeled polypeptides that were resolved in two dimensions. It is interesting to note that a polypeptide of molecular weight close to 43,000 was detected by Coomassie blue staining in the immunoprecipitate that included the other subunits of complex A (Fig. 8 B by Cole et al., 1998).

Independently from a complete definition of the composition of IFT complex A, three specific characteristics of fla15, fla16, and fla17-1 (namely the distinct bulge of the flagellar membrane, the defect in the concentration of IFT complex A within flagella, and the decrease of retrograde velocity of IFT) occur together as the result of a mutation in any one of three independent loci. This evidence suggests the hypothesis that IFT complex A is involved in retrograde IFT of particles. A defect of the product of FLA15, FLA16, or FLA17 may affect assembly and concentration of complex A and, therefore, the frequency of binding of the complex to a retrograde motor. Alternatively, a structural deficiency of complex A alters the activity of the retrograde motor. In both instances, the velocity of retrograde transport would be decreased.

Phenotypic analysis suggested that fla15, fla16, and fla17-1 affect different aspects of retrograde IFT. The highest defect in the concentration of complex A in flagella of fla15 was correlated with the highest deficiency in flagellar regeneration. In contrast, the highest defect in the velocity of retrograde transport in flagella of fla17-1 was correlated with the lowest deficiency in flagellar regeneration. The causes of these differences likely will be understood when the molecular identity of the defective gene product of FLA15, FLA16, and FLA17 is determined.

The defective gene product of FLA15 could be the subunit of complex A that has the apparent molecular weight 43,000. This subunit of complex A is absent from flagella of fla15, whereas the other subunits remain present in trace amounts. If fla15 is a null mutant of the 43,000 molecular weight subunit, the assembly of the remaining subunits of complex A may be minimal at the permissive temperature and nonexistent at the restrictive temperature, as we have observed. A similar scenario was described for the radial spoke mutant p24. In this mutant, subunit 2 of the radial spoke stalk was absent, whereas the subunits of the radial spoke head were present in reduced amounts. Furthermore, subunit 2 of the radial spoke stalk was identified as the putative gene product of PF24 by both dikaryon rescue and revertant analysis (Huang et al., 1981).

Complexes A and B together likely constitute the particles moving between outer doublet microtubules and flagellar membrane. Three lines of evidence supporting this hypothesis are derived from the characterization of different fla mutants. A decrease in the concentration of complex A in flagella of fla15, fla16, and fla17-1 resulted in a decrease in the frequency of bidirectional transport of particles. Furthermore, the deficiency of complexes A and B from flagella of fla10 at the restrictive temperature was correlated with a dramatic decrease in the number of particles. Finally, the accumulation of subunits of complexes A and B in flagella of fla14 was shown to occur concurrently with the accumulation of particles (Pazour et al., 1998).

Complexes A and B together likely constitute the 17S complex previously described. This identification is supported by the analysis of fla10, where KHPlFLA10 activity independently was shown to be required for the presence of the 17S complex (Piperno and Mead, 1997) or complex A and complex B within flagella (Cole et al., 1998). The identification also is supported by the evidence described here. 17S sedimenting fractions from the cytoplasmic matrix of fla15, fla16, and fla17-1 flagella included two protein complexes, one of which was identified as complex A.

The number of subunits of complex B may be higher than that reported earlier (Cole et al., 1998). We have

Figure 8. The four polypeptides that are deficient from flagella of fla15, fla16, and fla17-1 behaved as subunits of a complex. Autoradiography of a two-dimensional map of 35S-labeled polypeptides from flagella of a wild-type strain after subsequent sedimentation in sucrose gradient and chromatography in a DEAE-Sepharose column. Oblique lines indicate the four polypeptides that are deficient in each mutant but were not identified previously as subunits of complex A because both bind the monoclonal antibody 139.1 (Cole et al., 1998).

The defective gene product of FLA15 could be the subunit of complex A that has the apparent molecular weight 43,000. This subunit of complex A is absent from flagella of fla15, whereas the other subunits remain present in trace amounts. If fla15 is a null mutant of the 43,000 molecular weight subunit, the assembly of the remaining subunits of complex A may be minimal at the permissive temperature and nonexistent at the restrictive temperature, as we have observed. A similar scenario was described for the radial spoke mutant p24. In this mutant, subunit 2 of the radial spoke stalk was absent, whereas the subunits of the radial spoke head were present in reduced amounts. Furthermore, subunit 2 of the radial spoke stalk was identified as the putative gene product of PF24 by both dikaryon rescue and revertant analysis (Huang et al., 1981).

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Complexes A and B together likely constitute the 17S complex previously described. This identification is supported by the analysis of fla10, where KHPlFLA10 activity independently was shown to be required for the presence of the 17S complex (Piperno and Mead, 1997) or complex A and complex B within flagella (Cole et al., 1998). The identification also is supported by the evidence described here. 17S sedimenting fractions from the cytoplasmic matrix of fla15, fla16, and fla17-1 flagella included two protein complexes, one of which was identified as complex A.

The number of subunits of complex B may be higher than that reported earlier (Cole et al., 1998). We have
found that chromatography of $^{35}$S-labeled 17S sedimenting fractions on a DEAE-Sepharose column followed by two-dimensional electrophoresis of the polypeptides leads to the isolation of 34 molecules, including the subunits of both complexes A and B. Therefore, all of these polypeptides may form complexes that are stable at 0.3 M NaCl, pH 6.8, the salt concentration needed for elution from the column. Some of these molecules were at the limit of autoradiographic detection and, therefore, may represent modified subunits of complexes A and B or polypeptides that remained associated to these complexes in substoichiometric amounts, such as the inner arm subunit p28 (Piperno and Mead, 1997). However, other molecules present in the maps at higher concentration may represent additional subunits of complex B. This hypothesis is also supported by evidence reported by others. The two-dimensional map of the ensemble of complexes A and B prepared by Cole et al. (1998) also included several polypeptides that were as prevalent as identified subunits of complex A or B.

Complex A and a cytoplasmic dynein alone may constitute the whole machinery for retrograde intraflagellar transport. This hypothesis can be confirmed by the analysis of additional mutants. The tight correlation between the occurrence of a specific flagellar bulge and the deficiency of retrograde transport suggests that mutations of retrograde transport in other loci may be identified by the procedure described here. Characterization of the primary defects of these mutants combined with quantitative analyses of intraflagellar transport, such as that reported here, should yield a complete description of the molecular machinery carrying out retrograde transport within flagella.

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