Partially Functional Outer-Arm Dynein in a Novel *Chlamydomonas* Mutant Expressing a Truncated γ Heavy Chain

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The outer dynein arm of *Chlamydomonas* flagella contains three heavy chains (α, β, and γ), each of which exhibits motor activity. How they assemble and cooperate is of considerable interest. Here we report the isolation of a novel mutant, *oda2-t*, whose γ heavy chain is truncated at about 30% of the sequence. While the previously isolated γ chain mutant *oda2* lacks the entire outer arm, *oda2-t* retains outer arms that contain α and β heavy chains, suggesting that the N-terminal sequence (corresponding to the tail region) is necessary and sufficient for stable outer-arm assembly. Thin-section electron microscopy and image analysis localize the γ heavy chain to a basal region of the outer-arm image in the axonemal cross section. The motility of *oda2-t* is lower than that of the wild type and *oda11* (lacking the α heavy chain) but higher than that of *oda2* and *oda4-s7* (lacking the motor domain of the β heavy chain). Thus, the outer-arm dynein lacking the γ heavy-chain motor domain is partially functional. The availability of mutants lacking individual heavy chains should greatly facilitate studies on the structure and function of the outer-arm dynein.

Dyneins are molecular motors that drive various kinds of microtubule-based motility, including intracellular vesicle transport and ciliary/flagellar beating. Axonemal dyneins, the motors responsible for ciliary and flagellar beating, are present as multiple species, which are generally classified into outer-arm and inner-arm dyneins according to their positions in the axoneme. The outer-arm dynein consists of a single species of a multisubunit complex containing two or three distinct dynein heavy chains (DHCs), each of which exhibits motor activity. The inner-arm dynein, in contrast, comprises multiple species, consisting of a two-headed dynein containing two different DHCs and multiple single-headed species, each containing a single DHC. Motility analyses of *Chlamydomonas* mutants lacking particular dynein species, and in vitro motility assays using isolated dyneins, have shown that the various dyneins differ in their motile properties, suggesting that regular axonemal beating is based on the coordinated function of multiple dynein motors with distinct properties (for reviews, see references 23 and 25). However, the mechanisms by which dynein motor function is coordinated within the axoneme are not understood.

How different DHCs coordinate with each other is an important issue also for the functioning of multiheaded dyneins such as the outer-arm dynein. This is because distinct outer-arm DHCs have been shown to display strikingly different in vitro motility in sea urchins (31, 45) and *Chlamydomonas* (41). The *Chlamydomonas* outer-arm dynein is composed of three DHCs (α, β, and γ), two intermediate chains (IC1 and IC2), and 11 light chains (LCs) (6, 25). The N-terminal third of each ∼500-kDa DHC polypeptide is termed the tail or the stem, to which the two intermediate chains and most of the LCs are bound. The proximal region of the tail is the site where the three DHCs are associated with each other and attach to the A tubule of the outer doublet microtubule. The C-terminal region of each DHC consists of a AAA+ ring structure with four phosphate-binding motifs (P-loops) and a microtubule-binding stalk; the general organization of this region is conserved in all dynein species. LC1 is associated with this domain of the γ DHC (2, 43). When isolated and adsorbed to a surface, the outer arm takes on a bouquet structure, with the three AAA+ head domains projecting from the common base (11, 18, 47, 52). In the axoneme, this assembly is folded into a compact structure, which has been extensively studied using various electron microscopic techniques (10, 11, 17, 33, 34).

For studies aimed at elucidating the function and organization of individual DHCs in multiheaded dyneins, mutants missing a specific DHC are invaluable. In *Chlamydomonas*, mutants are available that lack the total outer arm (14, 21, 24, 29), only the α DHC (39), or the motor domain of the β DHC (40). The mutant lacking the α DHC, *oda11*, swims slower than the wild type but faster than mutants missing the entire outer arm, such as *oda2* and *oda4*. Thus, the outer-arm dynein containing only the β and γ DHCs can function in the axoneme, and the α DHC apparently increases the activity of the β/γ DHCs. The *oda4-s7* mutant, expressing only the N-terminal 160-kDa region of the β DHC and lacking its motor domain, assembles outer arms with the α and γ heavy chains, whereas the *oda4* mutant, with more severe defects in the β DHC gene, lacks the

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entire outer arm. In contrast to *oda11, oda4*-*s7* swims at almost the same speed as *oda4*. Thus, the motor domain of the β DHC appears to be essential for the function of the outer-arm dynein, and outer-arm dyneins containing only the α and γ DHCs are almost completely nonfunctional. However, it is not known how significantly the motor domain of the γ DHC contributes to overall outer-arm function.

The *oda11* and *oda4*-*s7* mutants were also useful for assigning the location of the α and β DHCs within the outer arm. Averaged outer-arm images in cross-section micrographs of the mutant axonemes located the α DHC at the tip of the outer arm and the motor domain of the β chain at an intermediate position between the base and tip (39, 40). From the combined images and the images of the *oda11 oda4*-*s7* double mutant, the γ DHC motor domain was predicted to localize to the inner lobe of the outer-arm image (40). These data have provided a basis for interpreting three-dimensional images obtained by cryo-electron microscopy and tomography (17, 33, 34).

Although *oda11* and *oda4*-*s7*, as well as other oda mutants lacking the entire arm, have greatly facilitated studies on the structure and function of outer-arm dyneins, another type of mutant has been awaited that lacks only the γ DHC motor domain. Such a mutant would greatly advance studies on the structure and function of individual DHCs. Here we report the isolation and characterization of such a mutant, *oda2*-4, which lacks only the motor domain of the γ DHC.

**MATERIALS AND METHODS**

**Strains.** The following Chlamydomonas reinhardtii strains were used: 137C (wild type), A54-e18 (a nit1 strain used for insertional mutagenesis; 46); the *oda2* strain lacking the entire outer dynein arm due to a mutation in the γ DHC gene (21); the *oda11* strain lacking the α DHC and the 16-kDa LC (LCS) of the outer arm due to a mutation in the α DHC gene (39); the *oda4*-*s7* strain lacking the motor domain of the outer arm β DHC (40); the *pf14* strain lacking the radial spokes (12); and the *pf18* strain lacking the central pair (12).

**Mutant isolation.** A54-e18 was mutagenized by insertional mutagenesis (48) using a linearized plasmid DNA, pMN56, composed of the pUC119 vector designed based on the cDNA sequences of the *oda11* gene (39); the *oda2* strain of *C. reinhardtii* (12); and the *oda4*-*s7* strain lacking the central pair (12).

**Electrophoresis.** The flagellar DHCs were analyzed by urea SDSPAGE with a 3 to 5% polyacrylamide gradient and a 3 to 8 M urea gradient (20). Intermediate chains and LCs were analyzed using SDS-PAGE with a 5 to 20% acrylamide gradient. All gels were stained with silver (4).

**Motility assays.** The swimming velocities of the mutant and wild-type cells were measured using a dark field microscope at a total magnification of ×100, a video recording system, and a personal computer. A red filter with a cutoff wavelength of 630 nm was used to suppress the response of the cells to light. The flagellar beat frequency was measured by analyzing the vibration of the cell bodies with a fast Fourier transform analyzer, which yields an approximate average of the beat frequencies in a large population of cells. All measurements were carried out in the Tris-acetate-phosphate medium at 25°C.

**Motility of isolated axonemes was examined** using the method of Besen et al. (3). Movements were observed using a BX50 microscope (Olympus, Tokyo) equipped with a dark field condenser, a 40× objective (numerical aperture, 0.85), and a 100-W mercury arc lamp. The images were recorded with a high-speed charge-coupled-device video camera (HAS-200IR; Ditec Co., Tokyo, Japan) operating at 200 frames/s.

**Southern blot analysis.** DNA was isolated from the *oda2*-4 mutant and the parent (A54-e18) cells using the method of Weeks et al. (50). The DNA was digested with restriction enzymes, loaded on a 0.8% agarose gel, transferred to nitro blue tetrazolium (KPL, Gaithersburg, MD).

**Immunoblotting assay.** Axonal polypeptides separated by SDS-PAGE in 5% or 10% acrylamide gels were transferred to a polyvinylidene difluoride membrane and incubated with monoclonal antibodies against γ DHC (12B and 25-b) (27), LC2 (R5391) (35), and LC4 (CT61) (44). Antibody reactivity was visualized by color development with 5-bromo-4-chloro-3-indolyl phosphate and nitro blue tetrazolium (KPL, Gaithersburg, MD).

**3° RACE.** For 3° rapid amplification of cDNA ends (3° RACE) (7), total RNA was isolated with TRIzol reagent (Invitrogen). Five micrograms of total RNA was subjected to reverse transcription using Superscript III reverse transcriptase (Invitrogen) and the specific adapter primer (5’-GGGACCGCTGAGTACGTGCTAC-3’). The first-strand cDNA amplification was carried out with an Expand High Fidelity PCR system (Roche) and two primers, a gene-specific primer (5’-GGGACCGCTGAGTACGTGAC-3’) and an abridged universal amplification primer (5’-GGGACCGCTGAGTACGTGCTAC-3’) (3° RACE system; Invitrogen). Dimethyl sulfoxide was added to the reaction mixtures at a final concentration of 10%. Subsequent double-strand cDNA amplifications were carried out with five primers, designed based on the cDNA sequences of the γ DHC (http://www.ncbi.nlm.nih.gov; U15303) and Nit1 (AF203033).

**DNA sequencing.** The PCR product, or DNA extracted from gel slices, was purified using the Wizard SV gel and PCR clean-up system (Promega). DNA fragments were sequenced using the BigDye Terminator v3.1/1.1 cycle sequencing kit (Applied Biosystems).

**Isolation of axonemes and fractionation of dynein.** The culture of cells, isolation of flagella, preparation of crude dynein extract, and fractionation of dynein by high-pressure liquid chromatography on an ion-exchange column were as described by Kagami and Kamiya (20). Instead of a Mono Q column as previously used, a Uno Q column (Bio-Rad) was used, as it provides better separation of the outer-arm dyneins.

**Electrophoresis.** The composition of the flagellar DHCs was analyzed by urea SDSPAGE with a 3 to 5% polyacrylamide gradient and a 3 to 8 M urea gradient (20). Intermediate chains and LCs were analyzed using SDS-PAGE with a 5 to 20% acrylamide gradient. All gels were stained with silver (4).

**ATPase assay.** The ATPase activities in the axonemes and extracts were assayed using EnzChek phosphate assay kit (Molecular Probes) in the presence of 1 mM ATP in HMDEK (30 mM HEPES [pH 7.4], 5 mM MgSO4, 1 mM dithiothreitol, 1 mM EGTA, and 50 mM K-acetate) at 25°C. Under these conditions, the axonemes continued beating for >10 min. When the activity in the high-salt axonemal extract was measured, the sample also contained 60 mM KCl in addition to ATP and HMDEK. The absorbance at 360 nm was monitored for 5 min, and ATPase activity was determined from the rate of phosphate release. The measurements were performed in duplicate and repeated at least three times using different preparations for each sample.

**Electron microscopy and image analysis.** The axoneme samples were fixed with 2% glutaraldehyde in the presence of 2% tannic acid, postfixed with 1% OsO4, dehydrated through a series of ethanol solutions, and embedded in Epon 812. Gray/silver sections, 60 to 80 nm thick, were cut, double-stained with uranyl acetate and lead citrate, and observed with a JEM1010 microscope (Jeol, Tokyo, Japan) at a magnification of ×40,000.

**Image analysis** was carried out basically following the protocol in reference 54 with some modifications. The cross sections of axonemes in which each doublet could be clearly seen as an assembly of protofilaments were selected. After the electron micrographs were corrected for the phase contrast transfer function following the estimated defocusing values, a set of images consisting of an outer doublet, inner and outer dynein arms, and the adjacent B tubule were boxed out as regions of interest (ROIs). All the ROIs were aligned so that they corresponded to the views seen from the proximal end of the axoneme. The ROIs were then classified by a hierarchical cluster analysis (55). The number of images was used was 126 (wild type), 234 (*oda11*), 324 (*oda4*-*s7*), and 153 (*oda2*-4).

The analysis, a threshold was set so that the largest cluster in each sample contained more than 20 images. This resulted in classification into 7 (wild type), 12 (*oda11*), 19 (*oda4*-*s7*), and 9 (*oda2*-4) clusters. Each major cluster was refined, averaged, and normalized so that the difference between each mutant and the wild type could be detected by the Student t test.

**RESULTS**

A novel type of γ DHC mutant retaining the α and β DHCs. In an attempt to isolate new types of mutants deficient in flagellar function, we generated ~5,000 clones of *nit1* transfectants from a *nit1* strain (A54-e18) and examined their motility phenotypes. Seven clones displayed jerky swimming, characteristic of oda mutants lacking outer-arm dyneins. SDS-PAGE of isolated axonemes indicated that six of them lacked...
the three outer-arm DHCs (α, β, and γ) (not shown), while one clone appeared to lack only the γ DHC, retaining the α and β DHCs (Fig. 1A). A genetic cross with oda2, a mutant lacking the entire outer-arm dynein due to a mutation in the γ DHC gene, did not yield wild-type daughter cells in 50 tetrads, suggesting that this clone is an oda2 allele. To confirm that this mutant in fact has a mutation in the γ DHC gene, we carried out genomic Southern analysis using a specific probe for the γ DHC gene (Fig. 3A). The hybridization pattern in the DNAs from the parent and this mutant, after digestion with three restriction enzymes, differed (Fig. 1B). Thus, this clone most likely carries a mutation in the γ DHC gene. To distinguish this mutant from the previous oda2 alleles lacking the entire outer arm, we named it oda2-t, because it produces a truncated γ DHC, as shown below.

Western blot analysis. Since the oda2-t mutant retains the α and β DHCs in the axoneme while the previously isolated oda2 alleles lack the entire outer arm, we speculated that oda2-t expresses the N-terminal tail portion of the heavy chain. The necessity and sufficiency of the tail portion of DHCs for the assembly of dynein complexes in the axoneme have been noted (27, 51). As shown in Fig. 2B and C, a moderately strong ~150-kDa band (asterisk) and two faint bands at ~180 kDa and ~105 kDa were detected by the antibody 12yB (B) and 25-8 (C). As shown in panel C, in contrast, the 25-8 antibody detected a DHC band in the wild type, nothing in oda2-t, and an ~180-kDa band (arrow), an ~150-kDa band (asterisk), and an ~105-kDa band in oda2-t.

because the C-terminally attached NIT1 fragment (see below) assumes a disordered structure and is easily degraded. Polypeptides of 180 kDa and 150 kDa were also detected by Western blot analysis (Fig. 4C). These results indicate that oda2-t indeed expresses the N-terminal portion of the γ DHC, whereas oda2 does not. Thus, it is likely that oda2-t axonemes contain a truncated form of the γ DHC.

The structure of the γ DHC gene in oda2-t. To examine the mutation in the γ DHC gene in detail, we first arbitrarily chose nine 500- to 1,500-bp sequences in the cDNA sequence (Fig. 3A) and examined whether the corresponding DNA fragments could be amplified by PCR from oda2-t genomic DNA. We found that the sequence corresponding to residues 3889 to 4363 of the registered cDNA sequence cannot be amplified but that the other eight fragments can be amplified. Since this observation suggested that oda2-t carries a gene insertion or deletion near the region that failed to amplify, we designed a 25-bp primer corresponding to residues 3381 to 3405 and carried out 3′ RACE with this and an abridged universal amplification primer using cDNA produced from the total oda2-r RNA. An ~2,000-bp DNA segment was obtained that contained partial sequences of the γ DHC and the NIT1 genes. Double-strand cDNA was then amplified by PCR using primers designed based on the sequences of the γ DHC and the NIT1 genes, followed by 3′ RACE. This resulted in the isolation of the cDNA corresponding to the γ DHC message expressed in this mutant. The amino acid sequence deduced from the cDNA sequence (Fig. 3B) indicated that the γ DHC in oda2-t has only 1,623 amino acids, containing a 1,270-amino-
Subunit composition of the _oda2-t_ outer-arm dynein. To examine the composition of the outer-arm dynein from _oda2-t_, we analyzed an axonemal high salt extract using ion-exchange chromatography with a Mono Q column or a Uno Q column. We separated the peaks that would normally contain the γ DHC (fraction 2 [arrow]) and the α, β, and γ DHCs (fraction 19) but retained a peak that contains the α and β DHCs (fraction 24). SDS-PAGE and Western blotting showed that _oda2-t_ axonemes lack the γ DHC but contain a novel polypeptide of an apparent molecular mass of 150 kDa (asterisk), corresponding to the N-terminal segment (for fractions 1 to 4 and 22 to 25) (Fig. 4C). In addition, an extra ~180-kDa band (arrow) is present for fractions 22 to 25. As for the Western blotting of the axonemes (Fig. 2), we interpret the ~180-kDa band to represent the full-length product and the ~150-kDa band to be a proteolytic fragment, the latter being predominant. We predict that fractions 1 to 4 correspond to a complex containing the truncated γ chain (including both the 150-kDa and 180-kDa polypeptides) and that fractions 22 to 25 correspond to a mixture of a complex containing the α, β, and truncated γ DHCs and a complex containing only the α and β DHCs.

Of the 11 LCs currently identified within the outer-arm dynein, LC1 (22 kDa) and LC4 (18 kDa) are known to associate with the γ DHC; LC1 is bound to the motor domain (2), and LC4 is permanently associated with the tail portion (26) and interacts in a Ca²⁺-dependent manner with a region (residues 875 to 1182) containing two IQ motifs (44). As shown in Fig. 4B and C, in the wild-type axoneme, LC1 is present in the peak fraction of the γ DHC (fraction 2) and the αβγ DHCs (fraction 19), but it is totally absent in the corresponding fractions from _oda2-t_ axonemes. In contrast, LC4 was found in both wild-type and _oda2-t_ samples. Western blot patterns indicate that it is present in the fractions of the γ DHC (fraction 2) and the αβγ DHCs (fraction 19) of the wild type and in fractions 1 to 4 and 22 to 25 of _oda2-t_. Similarly, LC2 is present in both the wild type and _oda2-t_, in the fractions that contain the α and β DHCs but not in the fraction that contains only the γ DHC (fractions 1 to 4). These results and the SDS-PAGE patterns of the low-molecular-weight bands (Fig. 4B) are consistent with the previous report that LC1 and LC4 are, respectively, bound with the head and tail domains of the γ DHC, while other LCs are associated with the αβ DHCs or with the ICs. These observations also indicate that the truncated γ DHC remains stably associated with the αβ complex.

Electron microscope observations. In thin-section electron micrographs (Fig. 5A), _oda2-t_ axonemes bear an average of 6.7 ± 1.5 (standard deviation) outer arms per nine outer doublets (n = 100). This is about 85% of the number in wild-type axonemes (7.9 ± 0.3, measured in samples prepared at the same time; Fig. 5B). Thus, _oda2-t_ appears to have a slightly impaired ability to assemble outer dynein arms on the outer doublets. A similar reduction in the outer-arm number has been observed in _oda4-s7_, a mutant having outer arms with a truncated β DHC; in this case, the axoneme has been reported to have an average of 5.7 ± 1.1 outer arms (40).

When viewed in cross section, the outer arms in the _oda2-t_ axonemes appeared somewhat thinner than those in the wild-type axonemes (Fig. 5A). To examine the difference in a more quantitative manner, we classified and averaged the individual outer-arm images using an image-clustering protocol. Two typical image groups of _oda2-t_ axonemes are shown in Fig. 6A, together with the density difference from the averaged image of the wild-type outer arm. One group displays a major density difference in the inner lobe of the outer arm, while the other group shows two areas of density difference, one in the inner lobe, and the other in the outer side of the arm. In all groups of _oda2-t_ outer-arm images, the density in the inner lobe (indicated by the green arrows in Fig. 6) was consistently found to be reduced, suggesting that the motor domain of the γ DHC is...
The CT61 antibody detected LC4 in fractions where the SDS-PAGE analysis. The order was observed for the flagellar beat frequency, as determined from the power spectra of swimming cells (Fig. 7C). The spectrum of oda2-t has a peak at about 50 Hz, while the spectra of oda4-s7 and oda2 have peaks at about 30 Hz. These results suggest that outer-arm dyneins lacking γ DHC motor domains can function to some extent, while outer arms missing β DHC motor domains are almost completely nonfunctional.

**Motility.** The swimming velocity and flagellar beat frequency in oda2-t, oda2, oda4-s7, oda11, and the wild type are compared in Fig. 7. The swimming velocity of oda2-t is much lower than that of the wild type, but interestingly, it is significantly higher than that of oda2, a mutant lacking outer arms (Fig. 7A). A comparison of these five strains revealed that their average swimming velocities are in the following order, from highest to lowest: the wild type, oda11, oda2-t, and oda4-s7 and oda2, which were approximately equal (Fig. 7B). The same order was observed for the flagellar beat frequency, as determined from the power spectra of swimming cells (Fig. 7C). The

**FIG. 4.** Purification and composition of outer-arm dyneins from oda2-t. (A) The elution patterns of extracts from the wild-type and oda2-t axonemes. Axonemes were extracted with 0.6 M KCl, and the extracts were fractionated on a Uno Q column. Elution was carried out with a linear 80 to 300 mM KCl gradient. Note that the γ DHC peak (arrows) and the αβγ DHC peak are smaller for the oda2-t extracts than for those of the wild type. (B and C) SDS-PAGE patterns and Western blotting of the peak fractions (Fr.). (B) Heavy chains, intermediate-sized chains, and LCs were separated on a 5 to 20% gel (the central portion of the gel is not shown). LC1 is absent from the wild type. (B and C) SDS-PAGE patterns and Western blotting of the peak fractions (Fr.). (B) Heavy chains, intermediate-sized chains, and LCs were separated on a 5 to 20% gel (the central portion of the gel is not shown). LC1 is absent from the wild type. (B and C) SDS-PAGE patterns and Western blotting of the peak fractions (Fr.). (B) Heavy chains, intermediate-sized chains, and LCs were separated on a 5 to 20% gel (the central portion of the gel is not shown). LC1 is absent from the wild type.
axonemes in solution displayed beating with similar asymmetrical waveforms (Fig. 7D). At 10^{-4} M Ca^{2+}, ~60% of the wild-type axonemes were motile and all displayed beating with a symmetrical waveform moving straight in the medium at a velocity of 140.2 ± 29.3 μm/s (average ± standard deviation for 20 moving axonemes). In contrast, only ~10% of the _oda2_ axonemes beat and did so with a symmetrical waveform of very small amplitude (Fig. 7D). These data are consistent with the observations of Mitchell and Rosenbaum (29). The beating of _oda2_ axonemes was so ineffective that it did not result in the efficient propulsion of the axoneme; the velocity of movement was only 3.5 ± 1.3 μm/s (n = 20), i.e., about 1/40 of the wild-type speed. The movement of the _oda2_ axonemes at 10^{-4} M Ca^{2+} was much the same as that of _oda2_; the axonemes moved at 5.74 ± 1.4 μm/s (n = 20). These results suggest that outer-arm dyneins without the γ-DHC motor unit cannot contribute to the generation of symmetrical waveforms that cause effective propulsive force, even though LC4 is retained.

**ATPase activities.** The ATPase activities of the wild-type and mutant axonemes, either before or after extraction with 0.6 M KCl, are compared in Table 1. The values are expressed as the amount of phosphate liberated per unit time by a constant amount of axonemes. We used solution conditions in which axonemes displayed beating for longer than 10 min and checked that the phosphate liberation was linear with time for at least the initial 5 min. The axonemal ATPase activities thus measured were several times higher than those reported previously but consistent with the values obtained by Yagi and Kamiya (53) who used similar solution conditions. As reported previously (8, 19), the ATPase activity of axonemes is significantly higher than the sum of the activities in the high-salt extract and the extracted axonemes, indicating that the dynein ATPase is activated in situ. This activation is most likely caused by the interaction of dynein with the doublet microtubules.

Several features of the axonemal ATPase activities are noteworthy. First, as previously reported, ATPase activity was very low in axonemes that lack the entire outer arm (14, 19, 29). More than 75% of the ATPase activity in wild-type axonemes can be accounted for by the activity of outer-arm dyneins. Second, the ATPase activity in _oda4-s7_ was markedly lower than the activities of other mutant axonemes with partial
outer-arm defects, indicating the importance of the β DHC for the high ATPase activity in the axoneme. Finally, and most strikingly, the ATPase activity of the *oda2*-t axonemes was higher than those of the wild type.

**Lack of a suppressor activity.** Previously, a sup-*pf*-2 mutant was isolated that has an activity that suppresses the paralyzed-flagella phenotype of mutants deficient in the central pair microtubules or radial spokes (15). It was later shown to have a mutation in the γ DHC gene (38), and a number of additional alleles have been isolated (28, 38). An interesting feature in one of the sup-*pf*-2 alleles is that the number of outer arms in the axonemal cross section is reduced and the degree of the reduction is biased toward a subset of outer-doublet microtubules; outer arms are more often missing on doublets 3 and 6 to 9 than on other doublets (38; for doublet numbering, see the legend to Fig. 5 and reference 13). However, since the defects in the γ DHC genes of other sup-*pf*-2 alleles have not been determined, it is not understood whether the suppressor activity results from a specific class of γ DHC mutations or from a general loss of its motile activity in the background of unimpaired α and β DHCs. Nor is the relationship between the suppressor activity and the position of the outer-arm loss known.

To answer these questions, we examined whether *oda2*-t exhibits suppressor activity and counted the number of outer arms on specific outer doublets. We found that double mutants between *oda2*-t and *pf14*, lacking the radial spokes, and between *oda2*-t and *pf18*, missing the central pair, did not show flagellar beating. Therefore, *oda2*-t does not have a suppressor activity. On the other hand, the loss of outer arms in *oda2*-t was found to be biased among the outer doublets, such that doublets 6 to 9 tended to show greater loss than doublets 2 to 5 (doublet number 1 lacks outer arms even in the wild-type axonemes; 13) (Fig. 5C). Interestingly, a similar bias was also observed with the *oda4*-s7 γ DHC mutant. Therefore, doublets 6 to 9 may have a general tendency to lose outer arms more frequently than doublets 2 to 5 when the outer arm has an unstable structure. This bias could be a consequence of the beat pattern of *Chlamydomonas* flagella, which must produce asymmetric distortion within the axoneme.

**DISCUSSION**

In this study we isolated a novel *Chlamydomonas* mutant, the *oda2*-t strain, which expresses a truncated γ DHC consisting of the N-terminal third of the molecule. This mutant retains outer dynein arms containing the α and β DHCs, whereas the previously isolated *oda2* γ DHC mutant lacks the entire outer arm. This difference in phenotype clearly indicates that the N-terminal portion of the γ DHC, corresponding to the tail, is important for the assembly of the outer arm on the outer doublet microtubules. The *oda2*-t mutant is reminiscent of the
oda4-s7 mutant, which expresses a truncated β DHC of approximately one-third the normal size; it retains outer arms containing the α and γ DHCs, in contrast with the previously isolated oda4 mutant, which lacks the entire outer arm (40). Unlike these DHCs, the α DHC is dispensable for outer-arm assembly; oda11, which lacks this DHC, assembles outer arms that contain the β and γ DHCs (39). Consistent with this observation, the N-terminal sequence of the α DHC differs significantly from that of the β or γ DHC (30, 51; T. Yagi, unpublished results).

Electron microscopic observations of axonemal cross sections, in combination with cluster analysis, have clearly shown that the outer-arm images in oda2-t lack the electron density at the innermost lobe located near the A tubule. Some classes of images showed, in addition to the change in the innermost lobe, a wedge-shaped area of density change on the outer side of the entire outer arm. However, this latter difference could be explained by an alteration in the orientation of the entire arm, which might well be caused by the loss of the proximal structure. Thus, all observations are consistent with the γ DHC motor unit being located at the innermost lobe of the outer arm. This localization is as predicted from previous studies on mutants lacking the α and β DHCs (39, 40). Recent studies using cryoelectron tomography (17, 33) or three-dimensional reconstruction from cryoelectron microscopic images (34) have indicated that the motor domains of the three outer-arm DHCs are stacked against each other, such that the three AAA+ rings are aligned at a shallow angle to the wall of the A tubule. A striking feature is that the motor domain of the most proximal DHC, which is the γ DHC as judged from the images of the mutants, is located very close to the microtubule wall, almost attaching to it. Therefore, the presence of the γ DHC may be critically important for the attachment of the entire outer arm to the microtubule at a fixed angle.

The swimming velocity and flagellar beat frequency of oda2-t

FIG. 7. Motility analysis. (A) Swimming velocities of wild-type, oda2-t, and oda2 live cells (n = 50). The average velocity (standard deviation) for each strain (μm/s) was as follows: the wild type, 161.6 (9.2); oda2-t, 87.8 (9.4); and oda2, 50.6 (6.6). (B) Comparison of the average swimming velocities for the wild type and mutants deficient in outer-arm DHCs (n = 50). (C) Flagellar beat frequencies for the wild type and mutants deficient in outer-arm DHCs measured using the FFT method (22). (D) Images of isolated axonemes beating in a medium containing 1 mM ATP and 10−8 M or 10−4 M Ca2+. Six images are shown for a single axoneme with variable time intervals so as to illustrate the sequence of beat pattern. The beat frequency (standard deviation) measured for 20 axonemes under these conditions was as follows: at 10−8 M Ca2+, 69.5 (11.4) Hz for the wt, 48.4 (3.4) Hz for oda2-t, and 28.8 (2.8) Hz for oda2, and at 10−4 M Ca2+, 78.7 (13.2) Hz for the wt, 14.5 (2.7) Hz for oda2-t, and 15.1 (4.8) Hz for oda2. Temperature, 25°C. Bar, 10 μm.

TABLE 1. Axonemal ATPase activity in the wild type and mutants deficient in outer-arm DHCs

<table>
<thead>
<tr>
<th>Strain</th>
<th>Missing DHC</th>
<th>Axonemes</th>
<th>High-salt extract</th>
<th>Extracted axonemes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>1.59 (0.08)</td>
<td>0.41 (0.02)</td>
<td>0.12 (0.039)</td>
<td></td>
</tr>
<tr>
<td>oda11</td>
<td>α</td>
<td>1.30 (0.10)</td>
<td>0.67 (0.004)</td>
<td>0.054 (0.013)</td>
</tr>
<tr>
<td>oda4-s7</td>
<td>β</td>
<td>0.60 (0.06)</td>
<td>0.08 (0.02)</td>
<td>0.054 (0.006)</td>
</tr>
<tr>
<td>oda2-t</td>
<td>γ</td>
<td>2.03 (0.30)</td>
<td>0.27 (0.02)</td>
<td>0.052 (0.002)</td>
</tr>
<tr>
<td>oda2</td>
<td>αβγ</td>
<td>0.35 (0.04)</td>
<td>0.01 (0.001)</td>
<td>0.039 (0.010)</td>
</tr>
</tbody>
</table>

Values are the averages and standard deviations (in parentheses) of the results from three independent samples, given in units of μmol phosphate released/minute/mg axonemes.
were much lower than those of the wild type and *oda11* but significantly higher than those of *oda2* and *oda4*s7. In view of the finding that the loss of the γ DHC motor domain produces a large change in the alignment of the outer arms, it is somewhat surprising that this deficiency does not completely impair outer-arm function, whereas the lack of the β DHC motor unit in *oda4*s7 results in a total loss of motor activity. Thus, from a phenomenological point of view, the β DHC is the most important power generator among the three DHCs of the outer arm. Studies that compared outer-arm structure in the presence and absence of ATP/vanadate demonstrated that the outer arm undergoes a large structural change during the mechanochemical cycle (34, 49). In terms of the change in relative position upon nucleotide binding, the α DHC displays the largest change, the β DHC an intermediate change, and the γ DHC almost no change (34). The observed large positional alteration in structure may result from the concerted action of the three DHCs and the γ DHC positioned at the base may exert the greatest influence on the overall structure of the outer arm (see Fig. 6A). However, the present findings indicate that the outer arm can function to some extent even without the γ DHC.

Despite the presence of partially functional outer-arm dyneins with truncated γ DHCs that retain the calcium-binding LC4, *oda2*t axonemes, unlike wild-type axonemes, did not display effective beating with symmetrical waveforms at $10^{-4}$ M Ca$^{2+}$. This finding suggests that the γ DHC is important for axonemal beating at high Ca$^{2+}$ concentrations. Since LC4 has been suggested to play a role in the Ca$^{2+}$-dependent waveform conversion (44), this further suggests that there is a tight functional interaction between LC4 and the motor domain of the γ DHC. In other words, LC4 possibly controls waveforms through the DHC to which it binds.

Comparison of the ATPase activities in outer-arm-deficient axonemes revealed several interesting features. Most notably, the axonemes of *oda2γ* have higher ATPase activity than those of the wild type. This result suggests that in the wild-type outer-arm dynein, the γ DHC is somehow inhibiting the microtubule-activated ATPase activity of the α and β DHCs. Previously, Nakamura et al. (32) examined the ATPase activities of an 18S subparticle of the outer-arm dynein containing the α and β DHCs and a 12S subparticle containing the γ DHC and showed that the γ DHC suppressed the ATPase activity of the αβ DHC subparticle when the subparticles reassociate. Our present finding is in good agreement with their conclusion. Another interesting feature is that the ATPase activity of *oda11* axonemes is close to that of the wild type, and the activity in the *oda11* high-salt extract is even higher than that in the wild-type extract. Our observations are consistent with the previous report that the α DHC suppresses the ATPase activity of the β DHC in purified samples (37) and apparently controls ATP-dependent microtubule binding by the β and γ DHCs (42). Taken together, the ATPase activity of the outer-arm dynein, with or without activation by microtubules, must be regulated through the interactions between the different DHCs or their associated components. The observation that the *oda11* and *oda2γ* axonemes have high ATPase activities yet display much poorer motility than those of the wild type suggests that these axonemes have defects in mechanochemical coupling, which is likely to be regulated through interactions between DHCs as well as between dyneins and microtubules.

For an understanding of how an outer-arm dynein functions, we need to understand the properties of individual DHCs as motor proteins, as well as their structures and the manner of assembly of the arm. The use of *Chlamydomonas* mutants has been the most powerful approach to this end. However, recently, motility analysis using *Trypanosoma* in combination with gene knockdown by RNA interference has started to provide novel insights (1, 5). For example, the outer-arm dynein of this organism has been shown to play a key role in the switching of the beat propagation direction (1, 5). Combined studies using *Chlamydomonas* forward genetics and *Trypanosoma* reverse genetics will yield further important information as to how outer arms function, and the data obtained from these organisms will complement each other. The availability of *Chlamydomonas* mutants lacking the motor domain of any one of the three outer-arm DHCs implies that the use of appropriate biochemical techniques and single and double mutants will enable us to obtain outer-arm dyneins with any desired combination of motor domains. These altered dyneins will greatly facilitate studies aimed at understanding how the outer arm functions.

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