Two Types of *Chlamydomonas* Flagellar Mutants Missing Different Components of Inner-Arm Dynein

Ritsu Kamiya,* Eiji Kurimoto,* and Etsuko Muto†

*Department of Molecular Biology, School of Science, Nagoya University, Nagoya 464-01, Japan; and †Department of Ultrastructural Research, The Tokyo Metropolitan Institute of Medical Science, Tokyo 113, Japan

Abstract. Two types of *Chlamydomonas reinhardtii* flagellar mutants (idaA and idaB) lacking partial components of the inner-arm dynein were isolated by screening mutations that produce paralyzed phenotypes when present in a mutant missing outer-arm dynein. Of the currently identified three inner-arm subspecies I1, I2, and I3, each containing two heterologous heavy chains (Piperno, G., Z. Ramanis, E. F. Smith, and W. S. Sale. 1990. *J. Cell Biol.* 110:379-389), idaA and idaB lacked I1 and I2, respectively. The 13 idaA isolates comprised three genetically different groups (idal, ida2, ida3) and the two idaB isolates comprised a single group (ida4). In averaged cross-section electron micrographs, inner dynein arms in wild-type axonemes appeared to have two projections pointing to discrete directions. In idal-3 and ida4 axonemes, on the other hand, either one of them was missing or greatly diminished. Both projections were weak in the double mutant idal-3 × ida4. These observations suggest that the inner dynein arms in *Chlamydomonas* axonemes are aligned not in a single straight row, but in a staggered row or two discrete rows. Both idal-3 and ida4 swam at reduced speed. Thus, the inner-arm subspecies missing in these mutants are not necessary for flagellar motility. However, the double mutants idal-3 × ida4 were nonmotile, suggesting that axonemes with significant defects in inner arms cannot function. The inner-arm dynein should be important for the generation of axonemal beating.

The inner and outer dynein arms in cilia and flagella are key protein assemblies that cause sliding between doublet microtubules (Gibbons, 1981). *Chlamydomonas* outer dynein arm contains three high molecular weight peptides (HMWs) with ATP-hydrolyzing activities, as well as several other proteins with lower molecular weights (Piperno and Luck, 1979; King and Witman, 1989). These proteins are assembled to form a three-headed, bouquet-like structure (Johnson and Wall, 1983; Witman et al., 1983; Goodenough and Heuser, 1984), and arranged at every 24 nm along the length of the outer-doublet microtubules. The inner arm, on the other hand, is more complex. It also contains several HMWs with the ATPase activities (Huang et al. 1979; Piperno and Luck, 1979, 1981; Piperno, 1988), but comprises two morphologically different species called dyads and triads (Goodenough and Heuser, 1985). Recently, Piperno et al. (1990) have presented evidence that there actually are three species of inner-arm dynein (called I1, I2, and I3), each containing two HMWs; there are six inner-arm heavy chains altogether. From structural analyses of mutants, they have suggested that the three species are aligned longitudinally in the order of I1-I2-I3, and the cluster of the three repeats every 96 nm.

Functional properties of the inner and outer dynein arms remain to be clarified. Specific removal of the outer dynein arm from sea urchin sperm axonemes results in a twofold reduction in reactivated motility and axonemal ATPase activity (Gibbons and Gibbons, 1973; Yano and Miki-Noumura, 1981). These observations support the view that the two types of arms perform almost identical functions in axonemal motility. However, studies with *Chlamydomonas* mutants have suggested that the two types of arms have somewhat different functions (Okagaki and Kamiya, 1986; Brokaw and Kamiya, 1987; Kamiya et al., 1989; Kagami and Kamiya, 1990). For example, mutations in inner arm lead to reduction in shear amplitude (amplitude of microtubule sliding) whereas those in outer arm result in reduction in the beat frequency (Brokaw and Kamiya, 1987).

For a definite comparison between the inner- and outer-arm functions, mutant axonemes devoid of either of the arms should be of great value. Mitchell and Rosenbaum (1985) and we (Kamiya and Okamoto, 1985; Kamiya, 1988) reported the isolation and characterization of outer-arm-missing *Chlamydomonas* mutants (pf28 and odas) that retained reduced motility. More recently, we isolated two types of motile mutants (idaA and idaB; previously named *ida* and *idb*, respectively) missing partial structure of the inner dynein arm.

---

Dr. Muto's present address is Aich Prefectural University of Fine Art, Aich 480-11, Japan.

1. *Abbreviations used in this paper.* HMW, high molecular weight peptides; I-projection, inner projection; O-projection, outer projection; wt, wildtype.
and described their properties briefly in a review (Kamiya et al., 1989). The present report describes the isolation and characterization of these mutants in more detail. We show that wild-type inner dynein arms appear to comprise two discrete rows of subunits, either one of which is predominantly missing in idaA and idaB axonemes.

Materials and Methods

Strains

Chlamydomonas reinhardtii 137e (wild type; wt), a mutant missing the entire outer dynein arm (oda1), and two types of inner-arm mutants (idaA [ida1-3] and idaB [ida4]) of both mating types were used. ida1 and ida4 were referred to as ida (or ida98) and idb, respectively, in previous papers (Brokaw and Kamiya, 1987; Kamiya et al., 1989). Isolation and characterization of oda1 have been described (Kamiya and Okamoto, 1985; Kamiya, 1989).

The inner-arm mutants ida1-3 and ida4 were isolated from nonmotile double mutants with oda mutant background in the following way. N-methyl-N'-nitro-N-nitrosoguanidine was added to oda1 cells, which had been grown in Tris-acetic acid-phosphate (TAP) medium (Gorman and Levine, 1965) to a late logarithmic phase under constant illumination, to give a final concentration of 20 μg/ml. After 15-45 min of the addition, the cell suspension was filtered, washed with buffer, and incubated in Tris-acetic acid-N-nitrosoguanidine by centrifugation and resuspension of the cells in fresh TAP medium. 1 ml aliquots were taken from this suspension and transferred to eight test tubes (~1 cm diam), each containing 3 ml of TAP medium. Cells were grown in these test tubes over a 12-h/12-h, light/dark cycle until the cells’ green color thickened at the upper part of the test tubes. Cells grown at the bottom of each test tube were then carefully transferred with a Pasteur pipette to another tube containing 4 ml of fresh TAP medium. This process was repeated three times over a period of 1-2 wk. The cells grown at the bottom of the last sets of test tubes were saved, diluted, and inoculated on 1.5% TAP agar plates. The agar plates were kept under constant illumination for 3-5 d. Colonies with heaped appearances were transferred with toothpicks to 96-well cell culture plates, each well containing 0.2 ml of liquid TAP medium. Usually 40 colonies were isolated from a single test tube. The plates were then placed under constant illumination for 1-2 d, after which time each well was observed with an inverted microscope. Completely nonmotile cells were saved and inoculated on TAP agar plates. 1 wk after the transfer to the agar plates, the cells were transferred to a gamete-inducing medium (Levine and Ebersold, 1960) and mated with wild-type gametes of the opposite mating type. Tetrams were obtained and analyzed according to standard procedures (Levine and Ebersold, 1960). The desired mutants were isolated from nonparental ditype tetrads, i.e., tetrads consisting of two daughter cells with oda phenotypes and two with an identical, slow-swimming phenotype. The latter two were saved and analyzed for the composition of flagellar axoneme. Mutants judged to be deficient in inner dynein arm were further mated with wild type, and daughter cells with both mating types were obtained. When two or more isolates from an initial test tube culture gave rise to daughter cells of similar flagellar composition, only one isolate was saved to assure that all the isolates were independent.

Double mutants were produced by a standard method of genetic crossing; desired mutants were obtained from nonparental-ditype tetrads containing two wild-type daughter cells.

Genetic Analysis

Newly isolated mutants were examined for allelism with other inner-arm mutants by using a standard method of tetrad analysis (Levine and Ebersold, 1960). Confirmation in temporary dikaryons (Starling and Randall, 1971) was not useful to determine whether a pair of mutants were allelic, because dikaryons between inner-arm mutants did not show good motility recovery even if the two mutants were nonallelic. A pair of mutants were judged to be allelic when cultures of >50 zygotes did not produce a wild-type daughter cell.

Isolation and characterization of ida1 mutants were described (Kamiya et al., 1989). The present report describes the isolation and characterization of ida1-3 and ida4. Mutants on various linkage groups were recovered even if the two mutants were nonallelic. A pair of mutants were examined for allelism with other inner-arm mutants by using a standard method of tetrad analysis (Levine and Ebersold, 1960). The desired mutants were isolated from nonparental ditype tetrads with both mating types.

EM

Flagella of wild type and mutants were isolated by the method of Witman et al. (1978), using dibucaine-HCl to detach flagella from the cell bodies. The isolated flagella were demembranated by suspending in a solution containing 10 mM Hepes, 5 mM MgSO4, 1 mM DTT, 1 mM EGTA, and 0.4% NP-40 (pH 7.4).

Electrophoresis

Compositions of dynein heavy chains were analyzed by SDS-PAGE using the method of Laemmli (1970), as modified by Jarvik and Rosenbaum (1980; see also Püster et al., 1982). A 3-5% polyacrylamide gradient and a 3-8 M urea gradient were used. Gels were stained with silver (Merril et al., 1981).
Results

Isolation of Inner-Arm Mutants

Soon after we started isolating dynein-arm mutants by screening cells that tended to grow at the bottom of a test tube culture, we came to notice that mutants deficient in inner dynein arms were more rarely obtained than outer-arm mutants; of ~100 slow swimmers obtained, only two were found to be inner arm-deficient mutants, whereas 35 were outer-arm-missing mutants of 10 different complementation groups (odas; Kamiya, 1988). We therefore used another method to isolate inner-arm mutants. This method, based on our presumption that a mutant with defects in both outer arm and inner arm would be nonmotile, comprised two steps: isolation of nonmotile double-mutants from mutagenized odal cells, and removal of the odal background by back-cross of the double mutants with wild type. This procedure proved to be effective for isolation of certain types of inner-arm mutants; of ~70 nonmotile double-mutants that underwent mating with wild type, at least 25 appeared to have defects in the inner arm in addition to the odal mutation. SDS-PAGE analyses (see below) of axonemes indicated that the second mutations in 15 of the 25 double mutants had stable phenotypes and can be classified into two types which we call idaA (13 strains) and idaB (2 strains). Some of the other inner-arm mutations found in these isolates had defects distinct from those in idaA or idaB. For example, one type of mutant was missing No. 4 heavy chain which are present in both idaA and idaB axonemes (for the nomenclature of heavy chains; Fig. 1). These mutants may have defects in a third inner-arm subspecies, although we were unable to assign the No. 4 HMW band to one of the three inner arm subspecies that have been identified with an SDS-PAGE system different from ours (Piperno et al., 1990). We did not use them in this study because of their somewhat variable phenotypes.

Genetic Analyses

Genetic crosses between these strains indicated that these idaA strains comprised three groups ida1–3, while the two idaB strains comprised one group ida4 (Table I). As reported before, ida1 is allelic to pf30, a similar mutant isolated by Huang, Ramanis, and Luck (Brokaw and Kamiya, 1987). Al-}

Table I. Classification of Inner-Arm Mutants

| Type   | Genetic group | Linkage group | Strains (isolation numbers) |
|--------|---------------|---------------|-----------------------------|
| idaA   | ida1          | XII           | 98,* 114, 116, 138, 146, 177, (pf30)* |
|        | ida2          | XV            | 124,* 131, 132, 133, 135     |
|        | ida3          | III           | 129, 206*                    |
| idaB   | ida4          | XII           | 166,* 168                    |

* Used as the exemplar strain for each group.

though pf30 had been mapped to linkage group IX (see Harris, 1989), we found ida1 and pf30 were linked with If2 on linkage group XII (ida1 × If2: parental di-type: nonparental di-type: tetra-type = 130:0:8). Also, ida2 was linked with nic1 on linkage group XV (58:0:3), and ida3 with acl17 (18:0:2) and pf15 (8:0:15) on linkage group III. Another type of inner-arm mutant ida4 was found to be closely linked with idal (20:1:0) and If2 (162:0:5). Therefore, idal, ida4, and If2 must be all located within a region of 2–3 cM at the left end of linkage group XII.

Axoneme Compositions

The SDS-PAGE pattern in Fig. 1 shows the difference in HMWs between wild-type and mutant axonemes. As reported previously (Piperno and Luck, 1979; Pfister et al., 1982), HMW bands of both inner- and outer-arm dyneins occurred in a small region corresponding to the molecular mass of 400–450 kD. (One or two strong bands often appeared just above these dynein bands, marked by m in Fig. 1. These are those with contaminating membrane proteins which tended to be strongly stained with silver.) The wild-type pattern contained three outer-arm bands (α, β, and γ), which were more intense than the inner-arm dynein bands. These bands were entirely absent from the oda pattern.

In ida1-3 axonemes, two bands (No. 1 and No. 2) between β and γ outer-arm bands are either missing or greatly diminished. The amount of residual No. 1 chain was slightly variable among different preparations of axonemes and among different strains, although it was constantly small compared with that of wild type. No. 2 band was missing from all of the ida1-3 axonemes. However, it was frequently difficult to identify this band in wild type and odal axonemes because of the overlap with the outer-γ chain and No. 3 inner-arm chain. On the other hand, ida4 axonemes had a reduced No. 5 band. In addition, this mutant appeared to lack another band (No. 3) that was overlapped with the outer-γ chain; in the double mutant ida4 and odal, this band was clearly missing. Three bands (No. 1, No. 2, and No. 5) were absent or weakened in the double mutant ida2 × ida4, although No. 1 band was not as totally absent as in the single mutant. The pattern of this double mutant was similar to that of the axoneme of a previously isolated mutant, pf23, except that the latter also lacked No. 4 band (Huang et al., 1979; Kamiya, R., unpublished observation). No. 3 band may also be missing in the double mutant; however, this was not confirmed because the triple mutant ida2 × ida4 × odal did not yield enough flagella for the analysis.

Besides the defects in the heavy chains, the inner arm

Figure 1. Portions of 3–5% SDS-urea-PAGE patterns showing the high molecular weight bands of mutant and wild-type axonemes. wt, wild type; ida2/ida4, recombinant between ida2 and ida4; odal/ida4, recombinant between odal and ida4; oda, odal. (*) indicate inner-arm bands missing or diminished in mutant axonemes. α, β, and γ indicate outer-arm heavy chains. (m) Bands with contaminating membrane proteins. Stained with silver.
mutants may well be missing some light chains and intermediate-sized chains. Although this possibility has not been made clear in the present study using one dimensional SDS-PAGE of axonemes, a study with fractionated dynein samples from mutants in fact indicates that ida1 lacks a peptide of 140 kD and ida4 lacks a peptide of ~30 kD (Kagami, O., and R. Kamiya, unpublished observations). More detailed analysis of the defects in lower molecular weight chains associated with the inner-arm mutations is now in progress.

**Electron Microscope Images of Mutant Axonemes**

Thin-section EM showed that both ida1-3 and ida4 axonemes retained inner-arm images, but that the inner-arm images in the mutants were weaker than in wild type (Kamiya et al., 1989). To look more closely at the inner-arm images, we averaged the cross-section images of doublet microtubules in mutant and wild-type axonemes using an image processor. Image averaging could be carried out by printing a cross-section image nine times, each time rotating the image by 360°/9° around the center of the axoneme (Markham et al., 1963). However, this procedure often results in a blurred image, because the outer doublets in cross-section micrographs are rarely positioned on an exact circle. We therefore averaged images of eight doublet microtubules (all the outer doublets but one that does not bear the outer dynein arm; Hoops and Witman, 1983) by superimposing the cross-section image of an outer doublet on that of another using a TV camera and an image processor (see Materials and Methods for details; see also Afzelius et al., 1990 for an improved technique).

Fig. 2 shows a gallery of cross-section micrographs of wt, ida1-3, ida4, and ida2 × ida4 axonemes together with the averaged doublet images. To show the extent of image variation, we present six sets of the original and processed images for each kind of axoneme. The micrographs used had been chosen only on the basis of whether all the nine outer doublets appeared clearly; no selection had been made on the basis of the appearance of the inner arm in the original and processed micrographs.

Such averaging clearly revealed differences between wild-type and mutant axonemes in their inner-arm images. In wt (Fig. 2 a), the inner arms appear to have two discrete projections, one pointing directly toward the adjacent doublet microtubule—outer (O)-projection—and the other to the inner part of the axoneme—inner (I)-projection. These two projections must not be artificial images resulted from the averaging, since they were frequently identifiable in individual outer-doublets in the original micrograph.

In ida1-3 (Fig. 3 b), the O-projection is missing or replaced by a thin fibrous structure, much weaker in intensity than the O-projection observed in wt axoneme. Because the thin fibres spans the entire distance between two adjacent outer doublets, we suppose its major part is the inter-doublet link, which can be observed in wt axonemes after extraction of dynein arms (Warner, 1983; Goodenough and Heuser, 1989). In contrast to ida1-3, ida4 axoneme lacks the I-projection (Fig. 2 c). The degree of the deficiency is slightly variable from an image of axoneme to another and some of the images show a low electron density in the I-projection region. However, the prominent globular appendage of the I-projection observed in wt and ida1-3 is consistently missing in all the micrographs of ida4 axoneme. In the double mutant ida2 × ida4 (Fig. 2 d), both of the two projections are weaker than the wt axoneme, although the image is somewhat variable. This kind of axoneme often has the O-projection in the ida4 axoneme. The occurrence of the O-projection may be because of the presence of the No. 1 band (see above) in this double mutant.

**Motility Characteristics**

All the mutants we isolated were motile (Table II). Therefore, neither the parts of inner arms missing in these mutants nor the entire outer arm (Kamiya and Okamoto, 1985; Mitchell and Rosenbaum, 1985) is necessary for flagellar motility. As Table II shows, ida4 had the same beat frequency as the wild type but its swimming velocity was ~2/3 of the normal value. This suggests that the ida4 mutation affected the flagellar waveform. Brokaw and Kamiya (1987) reported that ida1 flagella had a shear amplitude reduced to ~70% of the normal value. It is likely that ida4 also had a reduced shear amplitude, although this conclusion must await waveform analysis.

Since ida1-3 and ida4 were isolated by screening nonmotile double mutants with odal, the double mutants ida1-3 × odal and ida4 × odal were naturally nonmotile; these double mutants had short flagella that were almost completely motionless. Unexpectedly, however, the double mutants between the two types of motile inner-arm mutants (ida1-3 × ida4) were also found to have lost motility; their flagella showed only erratic twitching movements. This observation

| Table II. Motility and Flagellar Length in Dynein-arm Mutants |
|-------------|-----------------|-----------------|
| Flagellar length | Swimming velocity | Beat frequency |
| µm | µm/s | Hz |
| wt | 11.1 ± 1.9 | 1550 ± 23.3 | 63 |
| oda1 | 11.4 ± 2.0 | 58.0 ± 7.5 | 29 |
| ida1 | 10.9 ± 1.6 | 82.1 ± 13.8 | 50 |
| ida2 | 11.0 ± 1.2 | 77.7 ± 15.2 | 54 |
| ida3 | 10.2 ± 1.8 | 77.3 ± 11.0 | 45 |
| ida4 | 9.4 ± 1.6 | 102.2 ± 10.7 | 62 |
| ida2 × ida4 | 7.9 ± 3.4 | Nonmotile |
| ida1 × oda1 | 3.0 ± 0.8 | Nonmotile |
| ida4 × oda1 | 5.6 ± 2.4 | Nonmotile |

Flagellar lengths and swimming velocities are those averaged from more than 30 measurements and expressed as mean ± SD. Beat frequency was measured by analyzing the bodily vibration of cells with a fast Fourier transform analyzer (Kamiya and Hasegawa, 1987); the median value of the frequency peak is shown for each strain. Motility was measured at 25°C.

Figure 2. Gallery of cross-section axoneme images and averaged images of the outer doublets in these photographs. (a) wild type; (b; every two photographs from left to right) ida1, ida2, and ida3; (c) ida4; and (d) double mutant ida2 × ida4. Arrows in the upper left photograph indicate the I- and O-projections of the inner arm image. Note that the O-projection is greatly weakened in b, whereas the I-projection is weakened in c. Bars: (top) 0.1 µm; (bottom) 0.02 µm.
suggests that axonemes with large defects in the inner arm are incapable of beating.

Discussion

Mutants Missing Inner-Arm Components

In the present study we developed a new procedure for isolation of inner dynein arm mutations and thereby obtained two types of mutants (idaA and idaB) missing different subsets of HMWs. The 13 idaA-type strains isolated were found to comprise three genetically different groups, which were named idal, ida2, and ida3. idal was found to be the allele of pf30, a mutant isolated by Huang, Ramanis and Luck and used in previous studies on inner-arm structure and function (Piperno, 1988; Brokaw and Kamiya, 1987; Goodenough et al., 1987). The other two idaA-type mutants ida2 and ida3 are reported here for the first time. The two isolates of idaB comprised a genetically single group ida4, which has been described only briefly in a previous study (Kamiya et al., 1989).

Piperno et al. (1990) have reported that the inner arm is composed of three species I1, I2, and I3, each containing two HMWs, and that pf30 (idal) lacks I1 and pf23 both I1 and I2. Since the SDS-PAGE pattern of ida1-3 \times ida4 axoneme is similar to that of pf23, it is likely that idal lacks the I2 heavy chains. Piperno, G. (personal communication) has found this is actually the case; using an improved SDS-PAGE system, he has found that ida4 is missing one (chain 2'; Piperno, G., et al., 1990) of the two I2 heavy chains and has a reduced amount of the other (chain 2).

Inner-Arm Structure

The averaged cross-section images of wt and mutant axonemes (Fig. 2a) clearly demonstrated the presence of two distinct projections in the inner arms, one (O-projection) pointing directly toward the adjacent outer-doublet and the other (I-projection) toward the inner space of the axoneme. The former projection is greatly weakened in idal-3 (Fig. 2b) that lack the I1 subspecies and the latter in ida4 that lacks the I2 subspecies (Fig. 2c). These observations indicate that the three inner-arm subspecies are aligned not in a straight line within an axoneme, but rather in a staggered row, or in two discrete rows with the I1 subspecies in the outer row and the I2 subspecies in the inner row. In fact, a recent study using extensive series of thin-section tilt–series electron micrographs is suggesting that the inner dynein arms in Chlamydomonas do occur in two parallel rows (Muto et al., 1989; and manuscript in preparation).

Piperno et al. (1990) have indicated that the three inner-arm subspecies, I1, I2, and I3, are longitudinally aligned in the order of I2-I1-I3 repeating every 96 nm; they have shown that a gap is present at every 96 nm in the inner-arm row of wild type, and that this gap is wider in pf30 (idal) lacking I1 and still wider in another mutant pf23 lacking both I1 and I2. Although our present finding does not directly argue against their general conclusion, it suggests that a model with the three inner-arm species in line is too simple. Moreover, an alternative view that the three species are aligned in a staggered row seems to have some difficulty in explaining observed image intensities of the arms. In the micrographs of thin sections about 100-nm thick, which should contain all the dynein arm components within a repeat distance of 96 nm, the two inner-arm projections in wt are similar in electron density to each other and to the outer arm (Fig. 2a). On the other hand, if there are only three inner-arm subspecies aligned in a staggered row within the repeat distance, one of the two inner-arm projections seen in an axoneme cross section may well contain only one subspecies (containing two heavy chains) within the repeat unit, in which as many as four outer arms (each containing three heavy chains) are present. Thus, it would be difficult to explain the similar image intensity between the inner and outer arms by a simple staggered row model, unless we assume that some inner arm subspecies has a shape elongated in the direction of the axoneme length, or that the staining of the dynein arms is not linearly related with their actual mass.

The exact arrangement of the inner arm subspecies, however, must await further studies. One problem with our present study is that it gives us no clues about the localization of the third subspecies I3. The cross section micrographs of ida2 \times ida4 axoneme (Fig. 2d) show weakened inner-arm images which may represent the location of I3. However, this double mutant retains No. 1 HMW (a component of I1) (Fig. 1), and its inner-arm appearance is variable from a very faint one to the one similar to the image of ida4 inner arm (Fig. 2d). Thus the O-projection observed in this double-mutant might well be because of the presence of the No. 1 chain as well as of I3. Mutants missing both I1 and I2 completely, or those missing I3, will be necessary to locate I3.

We have tried to establish the inner-arm organization by observing the inner-arm images of mutants in longitudinal sections in the course of this study. However, the images we obtained varied so greatly from one axoneme to another that we were unable to reach a unified view. We think this difficulty is expected if the inner arm comprises two rows; if so, an inner-arm image in any longitudinal section should be that of the two rows superimposed, and this should vary greatly depending on the viewing angle. Hence, further studies using tilt series electron micrographs of various dynein mutants will be necessary to determine the organization of different inner-arm subspecies.

Motility in Inner-Arm Mutants

Both ida1-3 and ida4, as well as the outer-arm-missing mutant odal, were able to swim although more slowly than wt. Thus neither the part of the inner dynein arm missing in ida1-3 and ida4 nor the entire outer arm is necessary for generation of flagellar beating. However, the double mutant ida1-3 \times ida4 was paralyzed, as were double mutants ida1-3 \times odal and ida4 \times odal. These results indicate that the inner-arm subsets and the outer dynein arm are functionally independent and axonemes lacking a large part of the inner arm cannot beat. This idea agrees with the observation that a mutant (pf23) missing the I1 and I2 inner-arm subspecies is paralyzed (Huang et al., 1979).

Recently we have found that the ida2 \times ida4 axoneme can undergo sliding disintegration when perfused with protease and ATP, at a velocity as high as that in wt (Kurimoto and Kamiya, manuscript in preparation). Hence the lack of motility in the ida2 \times ida4 axoneme may be because of a defect in the mechanism that organizes microtubule sliding into regular axonemal beating, rather than a defect in the force.
production itself. It would be interesting to examine whether \( \text{id}a\text{d}3 \times \text{oda1} \) and \( \text{id}a\text{d4} \times \text{oda1} \) axonemes also can undergo sliding disintegration, and furthermore, to inquire by physiological experiments whether the defect in those mutants is in the initiation of bend formation or in the propagation of bend that is formed at the base.

Our studies on dynein-arm mutants have so far shown that no single species of axonemal dynein is essential to the generation of motility. Whether the third inner arm subspecies \( \text{i3} \) is essential has remained an important question.

We thank Drs. Hirokazu Hotani, Tomohiko Ito, Haruto Nakayama, and Nobuhiro Kamikie of Laboratory of Exploratory Research for Advanced Technology (Kyoto) for allowing us to use their IBAS image processor. Dr. Gianni Piperno (Rockefeller University, New York) kindly sent us a manuscript before publication. We are also grateful to him and Dr. Winfield Sale (Emory University School of Medicine, Atlanta, GA) for critically reading an early version of the manuscript, and to Dr. Elizabeth Harris (Duke University, Durham, NC) for sending us various mutants used for genetic analyses.

This study has been supported by grants-in-aid from the Ministry of Education, Science and Culture of Japan (01657001, 02239101) to R. Kamiya.

Received for publication 27 June 1990 and in revised form 16 October 1990.

References

Afzelius, B., P. L. Bellon, and S. Lanzavecchia. 1990. Microtubules and their protofilaments in the flagellum of an insect spermatozoon. J. Cell. Sci. 95:207–211.

Brokaw, C. J., and R. Kamiya. 1987. Bending patterns of Chlamydomonas flagella. IV. Mutants with defects in inner and outer dynein arms indicate differences in dynein arm function. Cell Motil. and Cytoskeleton. 8:68–75.

Gibbons, I. R. 1981. Cilia and flagella of eukaryotes. Nature (LoN.). 227:680–685.

Levine, R. P., and W. T. Ebersold. 1960. The genetics and cytology of Chlamydomonas. Ann. Rev. Microbiol. 14:197–216.

Kamiya, R., and M. Okamoto. 1985. A mutant of Chlamydomonas reinhardtii that lacks the flagellar outer dynein arm but can swim. J. Cell. Sci. 74:181–191.

Kamiya, R., E. Kurimoto, H. Sakakibara, and T. Okagaki. 1989. A genetic approach to the function of inner and outer arm dynein. In Cell Movement Vol. I. P. F. Warner, P. Satir, and I. R. Gibbons, editors. Alan R. Liss, Inc., New York. 209–218.

King, S. M., and G. B. Witman. 1989. Molecular structure of Chlamydomonas outer arm dynein. In Cell Movement Vol. I. P. F. Warner, P. Satir, and I. R. Gibbons, editors. Alan R. Liss, Inc., New York. 61–75.

Merrill, C. R., P. Satir, and J. S. Wall. 1983. Structure and molecular weight of the ciliary dynein outer arm. J. Cell Biol. 95:207–217.

Muto, E., R. Kamiya, and S. Tsukita. 1989. Structure of inner dynein arms in Chlamydomonas flagella. J. Muscle Res. Cell Motil. 10:269.

Okagaki, T., and R. Kamiya. 1986. Microtubule sliding in mutant Chlamydomonas devoid of outer or inner dynein arms. J. Cell Biol. 103:1895–1902.

Pfister, K. K., R. B. Fay, and G. B. Witman. 1982. Purification and polypeptide composition of dynein ATPase from Chlamydomonas flagella. Cell Motil. and Cytoskeleton. 2:525–547.

Piperno, G. 1988. Isolation of a sixth dynein subunit adenosine triphosphatase of Chlamydomonas axonemes. J. Cell. Biol. 106:133–140.

Piperno, G., and D. J. L. Luck. 1987. A motile Chlamydomonas flagellum devoid of outer arm dynein. J. Cell Biol. 107:1228–1234.

Starling, D., and J. S. Wall. 1983. Fine structure and molecular weight of the outer dyneins of Chlamydomonas. J. Submicrosc. Cytol. 15:193–198.

Yano, Y., and T. Miki-Noumura. 1981. Recovery of sliding ability in arm-depleted flagellar axonemes after recombination with extracted dynein 1. J. Cell Sci. 48:223–239.