Why Are Two Different Cross-linkers Necessary for Actin Bundle Formation In Vivo and What Does Each Cross-link Contribute?

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Abstract. In developing Drosophila bristles two species of cross-linker, the forked proteins and fascin, connect adjacent actin filaments into bundles. Bundles form in three phases: (a) tiny bundles appear; (b) these bundles aggregate into larger bundles; and (c) the filaments become maximally cross-linked by fascin. In mutants that completely lack forked, aggregation of the bundles does not occur so that the mature bundles consist of <50 filaments versus ~700 for wild type. If the forked concentration is genetically reduced to half the wild type, aggregation of the tiny bundles occurs but the filaments are poorly ordered albeit with small patches of fascin cross-linked filaments. In mutants containing an excess of forked, all the bundles tend to aggregate and the filaments are maximally crossbridged by fascin. Alternatively, if fascin is absent, phases 1 and 2 occur normally but the resultant bundles are twisted and the filaments within them are poorly ordered. By extracting fully elongated bristles with potassium iodide which removes fascin but leaves forked, the bundles change from being straight to twisted and the filaments within them become poorly ordered. From these observations we conclude that (a) forked is used early in development to aggregate the tiny bundles into larger bundles; and (b) forked facilitates fascin entry into the bundles to maximally cross-link the actin filaments into straight, compact, rigid bundles. Thus, forked aligns the filaments and then directs fascin binding so that inappropriate cross-linking does not occur.

Key words: actin • Drosophila • bristles • cross-links • fascin • forked

Bundles of actin filaments are key components in many eukaryotic cells. Obvious examples include microvilli, the acrosomal process of many invertebrate and lower vertebrate sperm, the stereocilia of the inner ear, the bristles and hairs of Drosophila, the sertoli cells of the testes, the ring canals and nurse cell bundles in insect oocytes, the fertilization cone, stress fibers, filopodia, and growth cones. Studies in the 1970s and 1980s focused our attention on the packing of actin filaments into bundles and identified and characterized some of the macromolecular cross-linking components involved. Details on how these cross-linkers were positioned in order to maximize the packing of actin filaments into bundles—no small feat as each actin filament is a helical polymer—followed and are reviewed by DeRosier and Tilney (1982). From the beginning, a number of descriptive studies on how the bundles are generated in vivo were published from which steps in bundle formation could be identified.

Experiments followed that aimed to reproduce the in vivo observations by studying the assembly of actin bundles in vitro using purified actin filaments and a single cross-bridge. Accordingly, Stokes and DeRosier (1991) investigated factors that control bundle order (the packing of actin filaments in each bundle), bundle size, and bundle number, using fascin, a cross-link isolated from sea urchin eggs. They related these variables to the relative concentrations of actin and fascin.

A careful comparison of the in vivo and in vitro results show that there are serious discrepancies not least of which is the time necessary to form an ordered actin filament bundle in vitro which takes 4–8 d for a small bundle (Stokes and DeRosier, 1991) yet occurs in less than 1 h for the formation of large bundles in vivo (Tilney et al., 1996). Furthermore, there are discrete stages in bundle formation in vivo involving reordering of the filaments in the bundles; these do not occur in vitro. And finally, subsequent studies have made us aware that in most precisely ordered biological bundles formed in vivo there are two and sometimes three macromolecular cross-linkers. Thus, in microvilli, villin and fimbrin are used, in Drosophila bristles fascin and the forked proteins are used (Tilney et al.,
two separate species of cross-links per bundle. These links for each bundle and why it is essential biologically to hold the bundles in a partially completed state so that fascin or the forked proteins are eliminated (Tilney et al., 1995). The only case known where there is only a single cross-link is in Limulus sperm, but this, judging by the re- construction of the bundle, is a special case, as the cross- link, scrin, binds to multiple subunits on each actin fila- ment (Schmid et al., 1994; Bullitt et al., 1988).

Why are different cross-links necessary for actin bundle formation in vivo? One idea (Tilney et al., 1996) based upon the time of appearance of the two cross-linkers in bristles during their growth (Wulfkuhle et al., 1998) and on our EM description of stages in bundle formation (Til- ney et al., 1995, 1996) is that one cross-link may be used early in bundle formation to tie the filaments together in small bundles so that they can subsequently be zippered together into a precise hexagonally packed unit by fascin. This idea is consistent with earlier observations on the de- velopment of microvilli (Chambers and Grey, 1979) and stereocilia (Tilney and DeRosier, 1986).

In this report we have tested the sequential cross-link idea and expanded it using mutants, developmental time points, and extraction methods. From these studies we are beginning to understand why two species of cross-links are essential in vivo and what each contributes during the elongation of a bristle. Our study provides the basis for fu- ture in vitro studies using purified cross-links and actin. What we find is that the forked proteins are used early in development to align the filaments into tiny bundles and to aggregate these tiny bundles into the seven to eleven larger bundles found in fully formed bristles. Then, forked proteins somehow orchestrate fascin entry into the bund- les which leads to maximally cross-linked, straight, com- pact and rigid bundles. In essence, the forked proteins hold the bundles in a partially completed state so that fascin can quickly add to generate large bundles with maxi- mal cross-linking. Our study has direct relevance to under- standing why other actin bundles (e.g., in microvilli and stereocilia) also contain two or more, albeit different, cross- links for each bundle and why it is essential biologically to have two separate species of cross-links per bundle.

Materials and Methods

Drosophila Stocks

The Oregon-R strain of Drosophila melanogaster was used as the wild type in these studies. The singed stock and the forked stock were obtained from the Drosophila Stock Center (Indiana University, Bloomington, IN) and maintained as viable homozygotes. The singed and singed stocks were generously supplied by L. Cooley (Yale University, New Haven, CT). The Inn1-dl-49, snKL chromosome was maintained over the Inn1Msd, Msd chromosome. Male third instar larvae hemizy- gous for the Inn1-dl-49, snKL chromosome were selected under a dissecting microscope (where the developing male gonad can be identified) and allowed. The snKL chromosome was maintained in males in a stock with C(1)RM-attached X females. Male third instar larvae hemizy- gous for snKL were obtained as above. A viable and fertile stock containing two copies of the wild-type forked gene and four transgenic copi- es (for a total of six copies per diploid female genome) was generously supplied by N. Petersen (University of Wyoming, Laramie, WY) (Petersen et al., 1994). Female larvae containing 50% of the normal forked protein concentration were used as heterozygotes. Elles were maintained on standard cornmeal-molasses-yeast food at 25°C, 60–70% relative hu- midity, with a 12-h day/night cycle. Complete descriptions of genes and symbols can be found in Lindsley and Zimm (1992) and on FlyBase (Fly- Base Consortium, 1998).

Developmental Staging

All animals were staged at the point of puparium formation, an easily rec- ognizable and brief stage lasting 30 min at the beginning of metamorpho- sis (Bainbridge and Bowens, 1981). White prepupae (0 time) were col- lected and placed on double-stick scotch tape in a Petri dish that was put back in the incubator at 25°C. Bristle elongation takes place during the in- terval of 32–48 h after pupariation (Tilney et al., 1996). At the appropriate time the Petri dishes were removed from the incubator and the pupae were dissected.

Dissection of Pupae

We modified our earlier procedure (Tilney et al., 1995). Under ideal mois- ture conditions, the outer pupal case could be removed with a pair of fine forceps since it is crunchy and tears easily. First, we removed the opercu- lum, and then with one tip of the forceps, tore the outer pupal case down the length of the pupa. The torn sides of the outer pupal case were then stuck down to the double-stick scotch tape. The pupae were lifted free by grabbing the tip of the abdomen, and the pupae were then stuck to an- other piece of double-stick scotch tape, ventral side down. The pupae were then covered with PBS. With a 5-mm scalpel (Accurate Surgical and Scientific Inc./Kaiser Medical Industrial Co., Westbury, NY), a small inci- sion was made between the developing eyes extending down to the mouth parts. With curved Gill’s Welsh Vannes micro scissors (Storz Co., St. Louis, MO), a cut was made from the incision down the sides of the head along both sides of the thorax above the wing buds, and then down the sides of the abdomen. A cross-wise cut along the anterior top of the abdo- men joined the two side cuts. The cut piece of tissue or dorsal surface of the thorax was removed, placed on is back, and the internal organs and fat bodies were removed with fine forceps. The flight muscles were not re- moved at this time, since they help the tissue maintain its integrity. Once cleaned, the tissue was fixed for light or electron microscopy.

Fixation and Processing for Light Microscopy

The cut and cleaned thorax was transferred to 4% paraformaldehyde in PBS for 20 min. The tissue was then transferred to 4% paraformaldehyde containing 0.1% Triton X-100 in PBS for an additional 20 min, washed three times in 0.1% Triton X-100 in PBS, and then placed in 0.1% Triton X-100 containing 10⁻⁷ M phallolid conjugated to rhodamine (Sigma Chemical Co., St. Louis, MO) for 30 min. The sample was then placed on a slide and mounted in glycerol Citiflour (Ted Pella, Inc., Redding, CA). A coverslip was applied, and the preparation was sealed with nail polish. Slides were examined either with a universal fluorescent microscope (Carl Zeiss, Inc., Thornwood, NY) or a model TCS 4D confocal microscope (Leica, Heidelberg, Germany).

To establish the presence or absence of fascin or the forked proteins af- ter extraction with potassium iodide (KI),1 we took the detergent or deter- gent/KI-extracted thoraces and fixed them for 5 min in 4% formaldehyde in PBS and then blocked by PBS-1% BSA and washed three times in PBS. The specimens were incubated in the primary antibody, diluted in PBS containing 1% BSA for 3 h or overnight at 4°C. The thoraces were then washed three times in PBS-1% BSA for 30 min each time and then incu- bated in secondary antibodies conjugated with fluorescein for 3 h. The specimens were washed and mounted as described above. Confocal im- ages were examined and printed using Adobe Photoshop (Adobe Sys- tems, Inc., Mountain View, CA).

The primary antibodies to the forked proteins were obtained courtesy of N. Petersen and the anti-Drosophila fascin antibodies courtesy of L. Cooley. They were used at concentrations of 1:500 and 1:10, respectively. Secondary antibodies conjugated with fluorescein were obtained from Sigma Chemical Co. (St. Louis, MO).

1. Abbreviation used in this paper: KI, potassium iodide.
Fixation and Methods for Transmission Electron Microscopy

Two types of fixatives were used. The most successful was immersion in 2% glutaraldehyde (from an 8% stock purchased form Electron Microscope Sciences, Fort Washington, PA) in 0.05 M phosphate buffer, pH 6.8. Fixation for 1 h was begun at room temperature and the sample was put at 4°C. After 1 h the tissue was then placed in 1% OsO₄ in 0.05 phosphate buffer, pH 6.2, at 4°C for 45 min. In the second procedure, fixation by immersion was carried out for 45 min at 4°C. The fixative, which was made just before use, consisted of 1% glutaraldehyde, 1% OsO₄, in 0.05 phosphate buffer, pH 6.2. This procedure was designed for fixing actin filaments in situ. A rationale for this fixation is outlined in Tilney and Tilney (1994). After fixation by either method the specimens were washed three times in water at 4°C to remove the phosphate and en bloc stained with 1% uranyl acetate and lead citrate, and then examined with a Philips 200 electron microscope (Philips Scientific, Mahwah, NJ). Because it is essential to have perfect transverse and longitudinal sections through the bristles, an appendage that curves in space, it was necessary to reorient the block, an extremely tedious undertaking. Photographs were taken of sections mounted on uncoated grids.

Scanning Microscopy

Adult Drosophila were fixed for several hours by immersion in 70% alcohol. They were then dehydrated completely, air dried, placed upon stubs, coated with tungsten-platinum, and then examined with a scanning microscope (AMR 1000; Amray Inc., Bedford, MA).

Results

Manipulating Forked Protein Concentration

Background. We have included this section because to understand the significance of the new data presented in this paper the reader should be cognizant of the data presented in our earlier studies. Unfortunately the significance of some of this background data was not appreciated when the earlier papers were written and thus we have emphasized the important points in this section.

There are three discrete stages in bundle formation that take place as the bristle elongates (Tilney et al., 1996). First, tiny bundles (less than 12 filaments each) appear at the tip of an elongating bristle (phase 1). Most of these tiny bundles are located near the plasma membrane (e.g., Tilney et al., 1996, Fig. 12 a shows 27 cortical bundles and 11 internal bundles). Second, these tiny bundles aggregate into the seven to eleven larger bundles that are attached to the plasma membrane (phase 2) (Tilney et al., 1996). More filaments must appear later as bundles of 180–265 filaments can be seen (e.g., Tilney et al., 1996, Fig. 12 b). Interestingly, the filaments are not hexagonally packed but display liquid order and, in longitudinal section, no 12-nm periodicity indicative of fascin is found. (For further details of why the 12-nm fascin period occurs see Tilney et al., 1995). Not surprisingly, when bristles are stained with antibodies, fascin is not yet present in the bundles, unlike forked which is present as predicted.

New Observations. No Forked Protein, Newly Emerged Bristles. The bristles first emerge in the 34-h pupae. Transverse sections through emerging bristles of the mutant forked₃₁₀ mutant 48 h after puparium formation. Of interest are the size of the seven actin bundles. The number of filaments per bundle are indicated on the micrograph. One of the bundles is enlarged to show that the filaments are hexagonally packed. Reprinted from Tilney et al. (1995).
only one internal bundle but throughout the cytoplasm there appear to be individual actin filaments. All filaments are oriented parallel to the elongating axis of the bristle as is the large population of microtubules.

**No Forked Protein, Fully Elongated Bristles.** From Fig. 1 we see that the diameter of the bundles at the tip of a fully elongated bristle are smaller than the wild type. In this micrograph there are 12, 18, 20, 16, 12, 9, and 3 actin filaments per bundle.

In other transverse sections through bristles the numbers of filaments per cross section are also small in number like that seen at the tip. As mentioned in Tilney et al. (1996) elongation of a bristle occurs at the tip. Therefore, a bundle near the base of a fully elongated bristle has had approximately 14 h to grow, differentiate, and expand by addition of more filaments. We have managed to cut transverse sections through two bristles at their base so we could count actin filament numbers. (We should note that obtaining perfect transverse sections through a bristle is no small feat since each bristle is curved and thus, the block must be continuously readjusted before sectioning; if the angle is off by more than a few degrees one cannot count the number of actin filaments.) In one of these sections we counted the number of filaments per bundle (Fig. 3). The values were 24, 47, 51, and 47 compared with 500–700 filaments per bundle in the wild type. As expected, the filaments are hexagonally packed because fascin is present. The bundles tend to be rectangular in shape.

To summarize, in the absence of the forked proteins, many tiny bundles form at the tip of the bristle but most fail to aggregate and subsequently disappear. As the bris-
tle matures some additional filaments become incorporated and are cross-linked by fascin into the bundles. However, the size of the mature bundles is considerably smaller in forked mutant bristles (50 filaments/bundle) when compared with wild type (e.g., 500–700 filaments/bundle).

Reduced Forked Protein Concentration, Fully Elongated Bundles. By constructing forked36a/1 females we could examine animals with a ~50% reduction in the amount of forked proteins.

By immunofluorescence of phalloidin-stained bristles this mutant contains the appropriate number of bundles per bristle and the bundles generally run parallel to each other (Fig. 4). Furthermore, the units or modules that by being attached end to end form each bundle (Tilney et al., 1996) are usually found in transverse register. What is different is that the bundles are more variable in thickness with many appearing somewhat swollen and less compact than in the wild type (Fig. 4). Some of the thinner bundles are also wavy in profile (Fig. 4).

The swollen, less compact nature of the bristles can be understood from thin transverse sections cut through bristles (Fig. 5). The actin filaments making up the bundles display liquid order. There are small patches in these bundles where the filaments are hexagonally packed (e.g., Fig. 5 B) but elsewhere there are spaces in the pattern where the filaments are less ordered. In many cases the diameter of the bundles is greater than that of the wild type due to the more disordered arrangement of the filaments. Nevertheless, the number of filaments per bundle is always less than in bundles in wild-type bristles of the same diameter. In longitudinal section through the bundles often we see the 12-nm period (Fig. 6). A careful comparison of this pattern of crossbridges relative to the wild-type is very instructive because it reveals that the crossbridges only span across a few filaments and not across all the filaments in the bundle. These areas correspond to the few areas in the transverse sections where the filaments are hexagonally packed (Fig. 5 B, arrows). Particularly interesting is that in longitudinal section the 12-nm period extends between the cross-linked pairs of filaments for a long distance. Thus if fascin comes in, it zippers adjacent filaments together all along their length.

Increased Forked Protein Concentration. We also examined a mutant containing six copies of the forked gene (Petersen et al., 1994). By scanning microscopy the macrochaetes and some of the microchaetes often had a fish hook appearance (Petersen et al., 1994) (Fig. 7, A and B). At higher resolution, the grooves in the main shaft terminated at the barb of the fish hook and a new set at an acute angle to the first group appeared in the barb (Fig. 7 C).

Thin sections cut through the barb portion of bristles from this mutant were especially revealing (Fig. 8 A). The bundles were very large and often irregular in shape. The filaments within these bundles were hexagonally packed (Fig. 8 B) and in longitudinal section showed the 12-nm period indicative of fascin (Fig. 8 C). However, there are gaps in all the bundles where no filaments could be found giving the bundles a moth-eaten appearance (Fig. 8, A and B). We also find in this mutant that in addition to the seven to
eleven membrane-associated bundles, there are internal bundles composed of hexagonally packed filaments (Fig. 9 A). These internal bundles tend to aggregate together on one half of the bristle shaft. In longitudinal section these internal bundles display the 12-nm period of fascin. Longitudinal sections also revealed that many membrane-associated bundles appear to be composed of closely applied subbundles each containing \( \approx 50-100 \) filaments (Fig. 9 B). Thus, many of the gaps seen in transverse section are in reality areas where subbundles do not join perfectly into a single hexagonally packed bundle and/or that additional filaments have not been added to fill in spaces between subbundles as they aggregate during development. Thus, from our thin sections, bristles in this mutant appear to contain more bundles than in the wild type but still have only seven to eleven membrane-associated bundles. These membrane-associated bundles are much larger due to the aggregation of many more smaller bundles than occurs in wild-type bristles. Even so, in all cases the filaments making up the bundles seem to be maximally cross-linked by fascin. There seems to be no liquid ordered filaments.

**Altering the Fascin Cross-link**

**Background.** Cant et al. (1993) analyzed the **singed** \(^{X2}\) and **singed** \(^{x}\) mutations. Although neither appears to be a complete null, immunofluorescence with antibodies directed against the **singed** product—fascin—showed that the mutant bristles did not show any appreciable fluorescence. Although the length of the mutant microchaete bristles were \( \approx 90\% \) that of the wild type (Tilney et al., 1995) all the bristles appeared twisted, often curled, or singed. The packing of the filaments was no longer hexagonal, but instead showed a liquid order (Tilney et al., 1995). In longitudinal section through the bundles the 12-nm cross striations were no longer present due to the absence of fascin. Nevertheless, actin filaments are present in bundles albeit the number of filaments per bundle is considerably less than the wild type. The number of bundles and the asymmetry in the position of the bundles was unaffected. What is particularly striking is that by immunofluorescence the actin bundles are not straight but are twisted (e.g., Tilney et al., 1996, Fig. 14), which we presume accounts for the twisted appearance of the bristles seen by scanning microscopy (e.g., Tilney et al., 1995, Fig. 7).

**New Observations. No Fascin, Early Stages in Bristle Elongation.** As in the wild-type and **forked** mutants tiny bundles of actin appear near the plasma membrane at the tip of elongating **singed** \(^{X2}\) bristles. There are also a few bundles and some solitary actin filaments in the center of the extension. By 37 h the tiny bundles begin to aggregate into the seven to eleven larger bundles seen in the mature bristles. This process of aggregation is particularly striking.
in the *singed* mutants. What one sees are ribbon-shaped aggregations of filaments (Fig. 10); these aggregations are adjacent to the plasma membrane. Each linear aggregate is composed of individual bundles of approximately a dozen filaments attached side by side. That the packing of the actin filaments in each tiny bundle or in the aggregate lacks hexagonal packing as would be expected. Thus, in the mutant where the forked proteins but little or no fascin are present, aggregation occurs in association with the plasma membrane, but in resulting bundles the filaments are poorly ordered.

**Fully Elongated Bristles in a Mutant Lacking Functional Fascin.** Recently we have examined the *singedS289N* allele described by Cant and Cooley (1996). Although this mutant produces nearly normal amounts of protein, the mutant protein appears to be completely inactive. When we examined surviving homozygous male pupae we see that both macrochaetes and microchaetes are twisted and the grooves which one sees by scanning microscopy also twist along their length. These grooves are deep and often split or fuse with each other (Fig. 11).

But what is interesting about this mutant are the actin bundles as visualized in thin section. These bundles, albeit small, are rectangular in shape and are usually associated with the membrane. As expected, the filaments within each bundle are not hexagonally packed but display a liquid-like arrangement (Fig. 12, *insets*). Of particular interest is that in this mutant there are also internal bundles, but in contrast to the *forked* mutant which overproduces the forked protein, each bundle is generally associated with or connected to a vesicle or within an infolding of the plasma membrane. We found a number of instances in which the peripherally located rectangular bundles are actually positioned along an infolding of the plasma membrane rather than the surface proper (Fig. 12). In longitudinal section part of the bundles are attached to the plasma membrane and half extend inward into the cytoplasm proper. Moreover, in longitudinal section, each bundle seems to be an aggregate of many smaller bundles as if after aggregation of the bundles there is an imperfect filling in of space between the surface of these bundles.

**Chemical Removal of Cross-links Affects Bundling**

To further determine the exact function of individual cross-links we extracted fully elongated wild-type bristles with a variety of reagents to selectively remove one or the other cross-link and then analyzed the morphology of the bristles and their actin bundles. An agent that was particularly revealing was potassium iodide (KI).

Our procedure was as follows. We dissected out the dorsal thorax, then treated the preparation with detergent followed by extraction with KI. The thoraces were then fixed and stained with fluorescently labeled phalloidin and/or antibodies against fascin or the forked proteins labeled with a different fluorochrome. What we noticed is that the actin bundles could still be detected in most bristles even in concentrations of KI as high as 0.6 M, but unlike the controls the bundles are twisted (Fig. 13A). The bundles were swollen at concentrations of KI above 0.3 M and in some cases the bundles were reduced in number, but 0.65 M was necessary to break down all the filaments.

An analysis of the effects of varying concentrations of KI on the presence of fascin or the forked protein is summarized in Table I. At 0.1 M all of the fascin was removed and apparently redistributed within the bristle. With 0.2 M
the bundles are fascin free. In contrast, the forked proteins remain at 0.1 M. At higher KI concentrations, forked proteins are extracted from the bundles but concentrate into spots both along and between bundles. These forked protein-positive spots are present even at high KI concentrations, up to 0.6 M.

In thin sections through bristles in 0.6 M KI the actin filaments in the bundles show liquid order (Fig. 13, B and C) and in longitudinal section do not display the 12-nm cross striations. These observations are consistent with the antibody studies that show fascin is removed under these circumstances. If the fascin is removed, then the filaments are only held together by the forked protein or aggregates of the forked proteins (the fluorescent spots) or by a yet unidentified cross-link. This in turn leads to poorly ordered filaments whose bundles are twisted.

Discussion

Small Imperfect Bundles Form in the Presence of the Fascin Cross-link

Using one cross-link, fascin, isolated from sea urchin eggs, and purified actin filaments, Stokes and DeRosier (1991)

Figure 9. Thin section through the fully elongated bristles from a mutant containing six copies of the forked gene. (A) In this transverse section through a bristle we find that the actin bundles are larger than the wild type and tend to aggregate together. Arrows indicate eight of the membrane-associated bundles. These bundles now associate with some internal bundles to form large masses of filaments. Of interest is that most of the bundles are located in the lower half of this cell. The dendrite (D) of the nerve is on the opposite side where there are no bundles. (B) Longitudinal section cut through an actin bundle. Notice that it is composed of a number of subbundles.

Figure 10. Thin section through a newly emerging bristle from a singed animal 36 h after pupation. Associated with the limiting plasma membrane are nine bundles of actin filaments. Of great interest are that the bundles are rectangular in shape. Arrows, linear series of tiny bundles.
What is the Precise Function(s) of the Forked Proteins?

In mutants that lack fascin (singed mutants) rectangular-shaped bundles are present in which the filaments are poorly ordered. Here the filaments are cross-linked together by a different crossbridge, presumably the forked proteins. A similar situation occurs when detergent-treated fully formed bristles are extracted with KI. From antibody staining of KI-extracted bristles it is clear that fascin is completely removed yet the forked proteins remain. Filament order changes from a precise order to a liquid order reminiscent of the singed mutants, which lack fascin at all stages. Thus the forked proteins, directly or indirectly via other components, hold the filaments together but the filaments are poorly ordered.

We also know from the studies of Wulfkuhle et al. (1998) (which we have confirmed) that in wild-type flies, forked proteins appear in the bundles of newly emerged bristles well before fascin. Thus, the tiny filament bundles at the newly emerging bristle tip are first joined together by the forked proteins. The filaments in these bundles are poorly ordered because fascin has not entered the bundles. Subsequently these tiny bundles aggregate into the seven to eleven larger bundles. Only after aggregation is fascin incorporated into the enlarging bundles. This ultimately leads to the filaments becoming maximally cross-linked. When we examine early stages in bundle formation in singed mutants that lack fascin but contain forked protein, in 36-h pupae we see long ribbon-like bundles attached to the limiting plasma membrane in transverse section. Thus in the fascinless mutant there has been a lateral aggregation of the tiny discrete bundles into wider ribbon-like bundles as in the wild type. At later developmental times, shorter, thicker bundles replace the long ribbons as if the ribbons collapse upon themselves. Over the same time scale, in mutants that lack forked proteins but contain wild-type amounts of fascin, aggregation of the tiny bundles does not occur. Accordingly, a second function of the forked proteins must be to aggregate small bundles into larger bundles in phase 2.

The above conclusion is consistent with observations on mutants overexpressing the forked proteins. In this mutant much larger bundles form. They are not flat, and unlike the singed mutant they bulge inward forming large triangular-shaped bundles. Each of these large bundles is composed of numerous subbundles which are aggregated together, presumably by the overexpression of forked protein.

Besides aggregating the actin filaments together into tiny bundles at the bristle tip, and somehow helping in the aggregation of the tiny disordered bundles into the larger disordered bundles, the forked proteins have a third function. This function (the last) must be to somehow, and we do not yet know precisely how, cooperate in reordering the disordered bundle into a large, ordered, compact, non-twisted, hexagonally packed, maximally cross-linked bundle. In short, the forked proteins must facilitate fascin binding. This conclusion stems from the observation that if only fascin is present in vitro or in vivo (e.g., in forked mutants) or the amount of the forked protein is reduced (e.g., forked+ heterozygote), large ordered bundles do not form although there are tiny regions of well-ordered fully crossbridged filaments in the heterozygote. Furthermore, if there is an excess of forked protein the filaments are always fully crossbridged by fascin even though there are more bundles produced. Further studies will be necessary.
to determine how the forked protein facilitates fascin binding.

To maximally cross-link filaments together into a three-dimensional bundle is a much more complex problem than one might expect because what is key is to inhibit inappropriate fascin cross-links so that the resultant bundle has no gaps in it or regions with liquid order between regions of more precise order. If liquid order were to occur the bundles would twist and not maintain strict linearity. Furthermore, the forked protein must somehow manage to increase the efficiency of adding additional filaments in an orderly way to the reordered bundles as they increase dramatically in filament number (Tilney et al., 1996).

We know from the work of Stokes and DeRosier (1991) that actin and fascin form tiny bundles in vitro composed of well-ordered filaments that can increase somewhat in filament number and/or diameter over 4–8 d. In bristles we know that the time it takes to go from a tiny bundle (34-h pupae) to large well-ordered bundles (37-h pupae) occurs in only 3 or 4 h and not the 4–8 d required in vitro. In addition, the ultimate size of the bundles formed in vitro versus in vivo differs by over an order of magnitude. These dramatic differences in time and number must be somehow attributable to the forked proteins as the forked proteins were not present in the in vitro studies.

The Fascin Cross-link Revisited: What Additional Function Exists for This Cross-link?

From earlier studies analyzing the cross-linking of actin filaments with a single crossbridge both in vitro and in vivo (DeRosier and Tilney, 1982) we know that the molar ratio of fascin/actin or fimbrin/actin in stereocilia is 1:4.6. This cross-linking is essential in forming a rigid bundle because if the cross-links are lost, e.g., during noise damage to stereocilia (Tilney et al., 1982) the rigidity of the stereocilia are greatly diminished. Furthermore, in the singed mutants that lack fascin or in bristles treated with 0.2 M KI, characteristically the bristles are/become bent and twisted and lie against the surface of the thorax. Not only does fascin contribute markedly if not almost exclusively to the rigidity of the bristles but it also inhibits the formation of sharp bends and/or twisted profiles. These observations are consistent with diffraction patterns of fascin/actin in in
vitro paracrystals (DeRosier and Tilney 1982; DeRosier and Censullo, 1981) or thin longitudinal sections through these crystals or through bristles (Tilney et al., 1995) where the filaments do not twist over each other but instead run perfectly parallel to each other. Thus, not only does fascin maximize the stiffness of the bristles but it also insures that the bristles are not twisted or bent, a situation key for a mechanoreceptor.

### Why Are Two Cross-links Essential to Form a Bundle In Vivo?

By comparing our in vivo results on wild-type bristles with those obtained on mutants that lack one or the other cross-link or with the in vitro observations of Stokes and DeRosier (1991), there are a number of reasons why two or more cross-links rather than one make more sense biologically. First, bristles up to 400 μm in length form in less than 14 h. These bristles contain up to 1,000 filaments per bundle. Thus, the time required for one cross-link to form in vitro (8 d for a small bundle of 50–100 filaments) is just not compatible with in vivo formation using two cross-links (14-h) which also results in much larger bundles (>1,000 filaments). Second, and this point is discussed briefly in Stokes and DeRosier (1991) and in Tilney et al. (1983), the way to form a bundle that is not twisted is to maximally cross-link adjacent actin filaments together. This results in a rigid bundle that will provide a cell extension such as a bristle with its characteristic shape. It is obvious that pairs of filaments maximally cross-linked together can be easily formed but to maximally cross-link filaments in three dimensions as in a bundle requires that the cross-links be made and broken until maximum order is achieved. Thus the fascin cross-link, intellectually at least, should have a weak binding constant so the fascin links can come off and go on at more appropriate positions. To hold the filaments to-

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**Table I. KI Treatment Selectively Removes Actin Cross-links**

| Antibody   | KI (M) | Uniform | None | Particulate | % Bristles stained | n |
|------------|--------|---------|------|-------------|-------------------|---|
| Anti-forked| 0      | 100     | 3    | 78          |                   | 71 |
|            | 0.1    | 97      | 6    | 83          |                   | 78 |
|            | 0.2    | 58      | 42   | 73          |                   | 73 |
|            | 0.3    | 75      | 25   | 4           |                   | 4  |
|            | 0.4    | 78      | 23   | 26          |                   | 26 |
|            | 0.5    | 100     | 4    | 4           |                   | 4  |
|            | 0.6    | 100     | 6    | 6           |                   | 6  |

| Anti-singed| 0      | 100     | 3    | 3           |                   | 3  |
|            | 0.1    | 100     | 31   | 31          |                   | 31 |
|            | 0.2    | 92      | 8    | 66          |                   | 66 |
|            | 0.3    |         |      |             |                   |    |
|            | 0.4    |         |      |             |                   |    |
|            | 0.5    |         |      |             |                   |    |
|            | 0.6    | 100     | 5    | 5           |                   | 5  |

Bristles were detergent extracted, treated with KI, fixed, stained with an antibody directed against one cross-link, and then visualized by confocal microscopy. The actin bundles were either stained evenly (Uniform), not at all (None), or exhibited a very nonuniform speckled appearance (Particulate).
gether during this ordering phase may require a different species of cross-link that at the same time does not inhibit fascin binding but instead facilitates appropriate fascin cross-linking and/or inhibits inappropriate cross-linking.

We are currently trying to determine how the forked proteins facilitate fascin binding or inhibit inappropriate interactions. One possibility is for the forked proteins to change the twist of the actin filaments and thus make every fascin–actin connection appropriate. Such a model would be consistent with the forked+/ heterozygotes where patches of filaments are maximally cross-linked together by fascin. These cross-links run the full length of the filaments.

**Two or More Cross-links per Bundle Is a Common Scenario in Biological Systems**

In microvilli of intestinal epithelial cells there are three actin cross-links per bundle, villin, fimbrin, and espin (Bartles et al., 1998), in nurse cell actin bundles there are two cross-links, villin and fascin (Cant et al., 1994; Mahajan-Miklos and Cooley, 1994), in the actin bundle in each stereocilium of the cochlea there is fimbrin and espin (Tilney et al., 1989) (Bartles, J.R., and L.G. Tilney, unpublished observations), and in bristle and hair bundles there is fascin and the forked protein. Significantly in large bundles such as those in stereocilia and in bristles, the formation of a maximally cross-linked bundles albeit by fimbrin or fascin forms in very similar ways. In both cases small bundles of poorly ordered (liquid order) filaments first appear and with time the order increases so that at the end one has maximally cross-linked, hexagonally packed bundles which display the 12-nm period in longitudinal sections. The same developmental sequence also occurs in small bundles. Examples include intestinal microvilli (Chambers and Grey, 1979) and Drosophila nurse cell bundles (Guild et al., 1997). In the latter case a poorly ordered bundle is present in the microvillus and as the bundle moves inward it becomes hexagonally packed. Furthermore, in the intestinal microvilli and in bristles the cross-links appear sequentially. Thus, the forked proteins precede fascin in bristle bundles just as villin precedes fimbrin in intestinal microvillar bundles (Shibayama et al., 1987; Ezzel et al., 1989). There is an additional parallel. Just as bristles that lack the forked protein form but are shorter and have poorly organized actin bundles, microvilli that lack villin also form in villin-deficient mice but they are also shorter and exhibit poorly bundled actin cores (Pinson et al., 1998).

**Conclusions**

What we proposed to do was to try and understand why two or more cross-links are necessary to form a maximally cross-linked and thus maximally rigid actin bundle. Such bundles could then be used by a cell as cytoskeletal units, the bones of a cell. To answer why questions is difficult, if not pretentious, so what we have done is to try and determine what role each cross-link plays in forming a mature bundle. But what the results from this paper actually do is raise more questions than were initially asked and thus opens new areas for future investigation.

For example, how do the forked proteins facilitate fascin cross-linking and how in the same cell at the same time do cross-links form between actin filaments sequentially (first forked proteins appear on the forming bundles then fascin appears). Furthermore, bristles have mature bundles at the base and less mature or early stage bundles towards the tip. Thus, in the same cytoplasm at the same time all the constituents are present and, yet miraculously, they somehow perform their individual job(s) and interact sequentially to produce straight, rigid, maximally cross-linked bundles.

Although there is abundant evidence that the forked proteins may be a second cross-link, up to now no one has been able to show that purified forked proteins bundle actin in vitro, an experiment that would show conclusively that forked is a cross-linking protein. The reason this information is not available is because recombinant forked protein is insoluble and to purify forked from Drosophila is just not practical. Notwithstanding there is genetic evidence that forked may be a crossbridge as demonstrated by this and earlier papers from our lab and others. Furthermore, antibodies to forked decorate bristle bundles and, if fascin is removed by KI, forked remains on the bundles presumably holding actin filaments together as it does in actin patches early in bundle development (Tilney et al., 1996; Wulfkuhle et al., 1998). Lastly, unpublished results cited in Wulfkuhle et al. (1998), indicate that certain regions of the forked protein can bind to actin filaments in vitro.

There are two additional puzzles that should be aired. First, we know that besides fascin there must be at least one additional cross-link holding the filaments together into bundles; presumably it is the forked protein or proteins associated with forked, but there may be other cross-links as well. Evidence that more than two may be present comes from observations that in the forked-singed double mutant (a mutant lacking both fascin and forked) tiny membrane-associated monolayers—we refer to them as rafts—appear (Tilney et al., 1995) and stain with phalloidin (Wulfkuhle et al., 1998). Second, before we can speculate or better determine exactly how a second and/or third cross-link interacts with fascin to form a bundle we need to know the stoichiometries of each relative to actin. This must be a topic for further study because at present we cannot determine this value in vivo and only by in vitro studies using the two or three cross-links can we determine this value. Nevertheless to better understand how these cross-links work together to facilitate fascin binding will require this knowledge.

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