Symbiont-Induced Changes in Host Actin during the Onset of a Beneficial Animal-Bacterial Association

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The influence of bacteria on the cytoskeleton of animal cells has been studied extensively only in pathogenic associations. We characterized changes in host cytoskeletal actin induced by the bacterial partner during the onset of a cooperative animal-bacteria association using the squid-vibrio model. Two-dimensional sodium dodecyl sulfate-polyacrylamide gel electrophoresis and Western blot analysis revealed that Vibrio fischeri induced a dramatic increase in actin protein abundance in the bacteria-associated host tissues during the onset of the symbiosis. Immunocytochemistry revealed that this change in actin abundance correlated with a two- to threefold increase in actin in the apical cell surface of the epithelium-lined ducts, the route of entry of symbionts into host tissues. Real-time reverse transcriptase PCR and in situ hybridization did not detect corresponding changes in actin mRNA. Temporally correlated with the bacteria-induced changes in actin levels was a two- to threefold decrease in duct circumference, a 20% loss in the average number of cells interfacing with the duct lumina, and dramatic changes in duct cell shape. When considered with previous studies of the biomechanical and biochemical characteristics of the duct, these findings suggest that the bacterial symbionts, upon colonizing the host organ, induce modifications that physically and chemically limit the opportunity for subsequent colonizers to pass through the ducts. Continued study of the squid-vibrio system will allow further comparisons of the mechanisms by which pathogenic and cooperative bacteria influence cytoskeleton dynamics in host cells.

Studies of beneficial and pathogenic bacteria have demonstrated that bacteria can dramatically influence the morphology, biochemistry, and molecular biology of host tissues with which they associate (13). Principal targets of such modifications are the polarized epithelial cells that line mucosal surfaces, the sites of first and often persistent interaction with bacterial cells (18). Among the changes in host epithelial cells that result from interaction with bacterial pathogens, perhaps the most conspicuous and well studied are the modifications of the host cell cytoskeleton (1, 8). By altering the host cytoskeleton, often through the activity of a secreted toxin or a factor injected into the host cell, various pathogens increase their intimacy with the host tissues, i.e., gain entry into cells, increase membrane contact with the cell surface, or invade tissues by disrupting junctions between cells. For example, enterohemorrhagic Escherichia coli rearranges the cytoskeleton of the host intestinal epithelium, which results in the formation of apical surface pedestals where the pathogen resides extracellularly (9).

Most studies of the interactions of bacteria with polarized epithelia have been restricted to the influence of pathogens on cultured mammalian cells (18). However, as the community of biologists increasingly recognizes the importance of beneficial bacteria in the health and disease of humans and other animals (15, 33, 34), models are being developed to characterize the nature of the interplay between the epithelial cells and their coevolved bacterial partners (10, 14, 20). One such model is the exclusive partnership between the host squid Euprymna scoloplos and the luminous bacterium Vibrio fischeri. This system offers a relatively simple experimental model for the study of persistent colonization of epithelia by extracellular bacteria in an intact association (23, 40). Both partners can be cultured independently under laboratory conditions, and molecular genetic methods have been developed for the bacterial symbiont.

Similar to the acquisition by mammals of their microbiota following birth, each generation of host squid acquires V. fischeri from the environment within hours of hatching from the egg (31). During embryogenesis, superficial ciliated fields that potentiate the colonization process develop on either side of the incipient light organ (Fig. 1A) (26, 31). At hatching, the cells of these fields secrete mucus within which V. fischeri aggregates for the first 2 to 3 h (32). The harvested V. fischeri are suspended by the cilia of the epithelial cells above a series of pores at the base of the appendages (Fig. 1B). At 3 to 4 h posthatching, amassed V. fischeri cells follow a path through these pores, travel down ciliated ducts, and colonize deep epithelium-lined crypts on each side of the light organ (Fig. 1C and D).

Beginning with passage through the ducts and the onset of colonization of the crypt spaces, V. fischeri induces a sequence of widespread developmental changes in the host light organ over the first hours to days (40). Using antibiotics to cure the light organ of V. fischeri, these modifications of host tissue have been found to be either reversible, i.e., constant exposure to V. fischeri is required to maintain these changes, or irreversible, i.e., a transient interaction with the symbiont is sufficient to induce morphological changes. Specifically, the crypt epithelium, which interfaces directly with the symbionts, undergoes a reversible increase in the microvillar density of the apical sur-

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Bar, 25 μm. (B) Higher-magnification confocal image of one side of the organ in the region of the ducts. The most medial duct (⊙) was used for all analyses in the present study. Bar, 25 μm. (C) Histological section of an uninfected juvenile light organ. Each pore (p) is an opening to a ciliated, microvillous duct that leads (arrows) to one of three epithelium-lined crypts (c) on each side of the light organ in the crypts, the bacteria interact with the epithelium and a population of macrophage-like hemocytes, which appear in this image as central in a crypt space. The dashed line indicates the position in the duct at which all confocal analyses were performed. Bar, 25 μm. (D) Transmission electron micrograph demonstrating the density of cilia (cil) along the duct (d = duct epithelial cell). Bar, 2 μm.

faces of the crypt cells (21) and an induction of edema in these cells (41). In contrast, the superficial ciliated fields, which are remote from the symbionts occupying the crypt spaces, undergo a 4-day program of regression that only requires a 12-h exposure to the symbionts (4, 27). In addition to these initial changes, the epithelial cells lining the ducts present high levels of oxidative stress conferred by an abundance of halide peroxidase and nitric oxide synthase (36; S. K. Davidson, R. Kossmehl, and M. J. McFall-Ngai, Abstr. 101st Gen. Meet. Am. Soc. Microbiol., abstr. N-107, 2001).

This study reports symbiont-induced changes in the actin cytoskeleton of the duct cells and provides evidence that these changes may work in concert with the other duct defenses to restrict entry into host tissues. Specifically, we (i) characterized actin gene expression at the level of transcription and translation in the E. scolopes light organ ducts during the onset of the association with V. fischeri; (ii) localized changes by immunocytochemistry and in situ hybridization; (iii) determined whether such changes are reversible or irreversible; and (iv) correlated these changes with symbiont-induced changes in duct dimensions and duct cell architecture. The results of these studies support the use of the E. scolopes-V. fischeri model for the study of bacteria-altered host cytoskeleton in an intact, beneficial association.

MATERIALS AND METHODS

General procedures. Juvenile squid were obtained from egg clutches that were maintained in open-flow seawater tables as previously described (4). Newly hatched squid were made symbiotic by incubating them in offshore seawater to which ~10^9 V. fischeri ES114 cells/ml were added for 12 h, followed by maintenance in Hawaiian seawater that did not contain sufficient quantities of V. fischeri to result in colonization. Squid were maintained in this offshore seawater immediately upon hatching to maintain them as aposymbiotic. All animals were kept individually in glass scintillation vials, and the seawater was changed daily. To confirm the presence or absence of symbiosis in host animals, the luminescence of the animals, which results from induction of luminescence in colonizing V. fischeri, was assessed daily by measuring light output with a photometer (model 20/20; Turner Designs, Sunnyvale, Calif.). In experiments where light organs were cured of their symbionts, the antibiotic chloramphenicol (CM) was used at 20 μg/ml of seawater (4). A CM-resistant strain of V. fischeri ES114(pV08), was used as a control for any pharmacological effects that CM treatment may have on the light organ tissues (39). Chemicals and antibodies were purchased from Sigma-Aldrich Chemical Co. (St. Louis, Mo.) unless otherwise stated.

Confocal microscopy. All confocal experiments were performed on a Zeiss LSM 510 system. In comparisons of fluorescence intensity, a ratio of duct fluorescence intensity over the fluorescence intensity of background tissue was calculated for each specimen. Dimensions of tissue features, as well as the intensity of cross-reactivity of fluorochromes, were determined using Zeiss LSM 510 software version 2.8. All measurements were conducted on the medial duct (Fig. 1B). The ducts develop sequentially over embryogenesis (26). At hatching, the medial duct is the most fully developed and leads to the most fully developed crypt. The less mature ducts and crypts have more anatomical variation than the medial duct and crypt. For each determination of fluorescence intensity within a duct cross-section, a 20-μm length of the brightest area was measured. All measurements were conducted below the level of saturation of the detector.

2-D PAGE. To prepare extracts of squid tissue for two-dimensional polyacrylamide gel electrophoresis (2-D PAGE) of unlabeled proteins, the juvenile squid were anesthetized in 2% ethanol in seawater. The light organs were dissected from the mantle cavity of the animal and accumulated in 50 mM Tris buffer (pH 7.2) on ice and then homogenized with a glass micromortar and pestle. The homogenate was centrifuged at 14,000 × g for 15 min at 4°C to remove bacterial cells, animal membranes, and squid ink. The soluble fraction was removed, and
the protein concentration was determined spectrophotometrically. Twenty micrograms of soluble protein sample was prepared for isoelectric focusing by adding four parts of lysis solution (8 M urea, 1% dithiothreitol, 2% Pharmalyte 3-10, 0.05% Triton X-100, 0.14% phenylmethylsulfonyl fluoride; pH 7.4) to one part protein sample. We used the Pharmacia Multiphor II 2-D electrophoresis system with precast Immobilon dry strip isoelectric focusing gels (linear pH gradient of 4 to 7 or 4.5 to 5.5) and ExcelGel (12 to 14% gradient) sodium dodecyl sulfate (SDS)-polyacrylamide gels. The gels were run and stained as previously described (5). For 2-D PAGE of microinjected proteins, prior to labeling squid were permeabilized for 20 min at 30°C to inhibit protein synthesis of the bacterial symbionts. Squid were then incubated for 2 h with ION TransS-label, which contains 70% [35S]methionine. Thirty micrograms of label was used per squid. Squid were first rinsed twice for several seconds and then once for 1 h in seawater supplemented with 8 mM methionine. 2-D PAGE gels of the soluble proteins were run as described above, and proteins that had incorporated radio-label were visualized by autoradiography. Experiments were done in triplicate to confirm reproducibility of the results.

Western blot analyses. For protein-antibody hybridizations, relevant sections of the 2-D PAGE gels were electrophoretically transferred to nitrocellulose membranes (0.45-μm pore size) at 15°C for 1 h at 200 mA in transfer buffer (25 mM Tris-HCl, 200 mM glycine, 20% methanol; pH 8.3). The membranes were then blocked overnight in Tris-buffered saline (TBS; 20 mM Tris with 0.5 M NaCl; pH 7.4) with 3% bovine serum albumin (BSA), followed by incubation with a 1:50 dilution of a polyclonal antibody (catalog no. A2066; Sigma-Aldrich Chemical Co.) to the conserved C-terminal 11 amino acids of the actin molecule in 1% BSA–Tween-20 (TTBS) overnight. They were then rinsed three times for 10 min in TBS and incubated for 45 min in goat anti-rabbit antibody conjugated to horseradish peroxidase (1:10,000 in 1% BSA–TTBS at room temperature and then rinsed three times for 10 min in TTBS at room temperature. Cross-reactivity with the antiactin antibody was visualized by chemiluminescence (Renaissance Western blot chemiluminescent kit; NEN Life Science Products, Boston, Mass.). One hundred nanograms of control actin from chicken was loaded onto the second dimensions of the 2-D gels as a positive control.

Immunocytochemistry. Squid were anesthetized in 2% ethanol in seawater and fixed in 4% paraformaldehyde in marine PBS at 4°C. Light organs were dissected from the mantle and rinsed three times for 10 min in 0.1% Tween 20–marine PBS (PTw). Specimens were fixed for 1 h at room temperature in 4% paraformaldehyde in PTw and then rinsed five times for 10 min in PTw. Specimens were incubated for 10 min at room temperature in hybridization buffer (50% formamide, 100 μg of salmon sperm DNA/mL, 1% SDS, 0.1% Tween 20, 50 μg of heparin/mL, and 5× SSC [1× SSC = 0.15 M NaCl and 0.015 M sodium citrate; pH 7.4]). Each specimen was incubated overnight at 60°C in hybridization buffer. Digoxigenin-labeled probes (50 ng/μL of stock) were denatured at 90°C in hybridization buffer for 10 min. After denaturation, probes were added to fresh hybridization buffer at a concentration of 0.1 ng/μL. Specimens were incubated overnight at 80°C, rinsed at 60°C for 20 min in 75% hybridization buffer–25% 2× SSC. They were then placed for 15 min in 50% hybridization buffer–50% 2× SSC, followed by 15 min in 25% hybridization buffer–75% 2× SSC, then 15 min in 100% 2× SSC, and finally twice for 30 min in 0.05× SSC. Samples were then rinsed at room temperature for 10 min in 75% 0.05× SSC–25% PTw, followed by 10 min in 50% 0.05× SSC–50% PTw, 10 min in 25% 0.05× SSC–25% PTw, and 10 min in 100% PTw. After rinsing, samples were blocked overnight at 4°C in marine PBS with 1% Triton X-100, 0.5% BSA, and 1% goat serum. Specimens were then incubated in alkaline phosphatase-conjugated sheep anti-digoxigenin Fab fragments (Boehringer-Mannheim, Indianapolis, Ind.) at a 1:500 dilution overnight at 4°C, washed three times for 10 min at room temperature in marine PBS with 0.2% Triton X-100 and 0.1% BSA, washed three times for 10 min in HNPP detection buffer (2-hydroxy-3-naphtoic acid-2-phenylanilide phosphate [Roche Applied Sciences, Indianapolis, Ind.] in 100 mM Tris-HCl, 100 mM NaCl, 10 mM MgCl2, pH 8.5), and then incubated for 2 h in HNPP-Fast Red TR substrate at room temperature according to the manufacturer’s instructions. Samples were then rinsed four times for 10 min in marine PBS, mounted in 2% DABCO (1,4-diazabicyclo[2.2.2]octane) in 50% glycerol–marine PBS, and viewed by confocal microscopy.

Labeling with phallolidin. Animals were anesthetized in 2% ethanol and fixed in 4% paraformaldehyde in marine PBS for 1 h at room temperature. Light organs were dissected into fresh marine PBS, rinsed twice for 10 min in marine PBS, and then permeabilized in 1% Triton X-100 in marine PBS for 20 min. Permeabilized samples were stained overnight in 2 mg of rhodamine phallolidin/ml in 1% Triton X-100 from marine PBS, then rinsed three times for 10 min in marine PBS, and viewed by confocal microscopy.

Measurements of duct dimensions and determinations of duct epithelial cell number and shape. To stain duct cells in living animals, live specimens were placed for 30 min in a 5-nm solution of acridine orange in seawater. Animals were then rinsed in seawater and anesthetized in 2% ethanol. The animals were ventrally dissected to reveal the light organ, and the circumference of the medial ducts was determined by confocal microscopy.

To investigate possible cellular mechanisms for change in duct dimensions, animal tissues of 48-h apysymbiotic and symbiotic animals were stained with rhodamine phallolidin (as described above) to study the number of cells lining the duct space and the shape of these cells.

RESULTS

Actin synthesis increased during the onset of the symbiosis. No reproducible differences in the steady-state profile of soluble proteins of 12-h apysymbiotic and symbiotic organs were detected in the present study (Fig. 2A), as previously shown (5). However, 2-D PAGE of 35S-labeled proteins from 12-h light organs detected several newly synthesized proteins, some of which were different between apysymbiotic and symbiotic organs and some of which showed similar levels of synthesis (Fig. 2B). Strong labeling of two proteins at an M, of ~42 and ~50 kDa suggested that they may be newly synthesized actin. Immunoblotting of the total soluble protein run in the

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same pH range of these gels, i.e., pH 4 to 7, confirmed that one or both of these proteins were actin (Fig. 2C). Immunoblotting in which proteins were resolved across a narrower pH range (i.e., 4.5 to 5.5) indicated that only one of these two proteins was actin (Fig. 2D). At this resolution, the proteins ran at slightly different pIs.

Increases in actin abundance were localized to the apical surfaces of the duct epithelia. Immunocytochemistry was performed to determine where symbiont-induced changes in actin levels were occurring in the light organ. Although the antibody used in immunocytochemistry analyses recognized both muscle and cytoskeletal actin in the host light organ, differences in antibody cross-reactivity, as detected by fluorescence intensity, were not observed between aposymbiotic and symbiotic animals in the labeling of the muscles or other portions of the light organ (data not shown). However, significant differences were observed between aposymbiotic and symbiotic animals in actin levels in the epithelial cells lining the light organ ducts within 6 h of exposure of the symbiotic animals to V. fischeri (Fig. 3).

The changes indicated an increase in actin in the microvillar surface and terminal web (Fig. 3A), the actin-rich region of the apical area of polarized epithelia. Beginning at the 6-h time point and continuing through 48 h, significant differences in immune cross-reactivity were detected in the duct between aposymbiotic and symbiotic animals (Fig. 3B). A significant increase in cross-reactivity was detected between 6- and 12-h symbiotic animals; thereafter, no significant differences were detected among the symbiotic animals at time points between 12 and 48 h. At 60 h, although the level of cross-reactivity for symbiotic ducts was on average slightly higher, it was not significantly different from that of aposymbiotic animals. No sig-

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**FIG. 2.** Actin protein was more actively synthesized in 12-h symbiotic light organs. (A) Sectors of silver-stained, 2-D SDS-PAGE gels of soluble proteins (20 μg loaded on each gel) of 12-h aposymbiotic and symbiotic light organs. No reproducible differences in protein profiles could be detected by this method. (B) Autoradiograms of 35S-labeled soluble proteins (20 μg loaded on each gel) of the light organs of aposymbiotic and symbiotic animals. Proteins at a pI and M₉ consistent with actin (1, 2) were more actively synthesized in symbiotic animals. (C) Upper panel: sector of a silver-stained, 2-D SDS-PAGE gel of soluble proteins from the light organs of symbiotic animals resolved in the pI range from 4.0 to 7.0. Numbers (1, 2) correspond to those in panel B. Lower panel: immunoblot of the above 2-D gel using antiactin antibodies. Control actin (arrow) was loaded onto the second dimension as a control. (D) Upper panel: sector of a silver-stained, 2-D SDS-PAGE gel of soluble proteins from the light organs of symbiotic animals resolved in the pI range from 4.5 to 5.5. Numbers (1, 2) correspond to those in panel B. Lower panel: immunoblot of the above 2-D gel using antiactin antibodies. Antibody labeling demonstrated that the more acidic of the two proteins (1) at ~42 kDa was actin. Biotinylated standards were used to determine the M₉. Molecular mass markers for A and B and for C and D appear on the left and right, respectively. pI markers are along the top. apo = aposymbiotic; sym = symbiotic.

**FIG. 3.** The V. fischeri-induced changes in actin protein levels occurred in the duct epithelia. (A) Confocal micrographs of cross-sections of ducts of 12-h aposymbiotic and symbiotic light organs. Light organs were labeled with antiactin rhodamine-conjugated goat anti-rabbit antibody. The medial ducts (arrows) were used for fluorescence quantification. (B) Relative fluorescence of sites cross-reactive with antiactin antibodies in the medial ducts of aposymbiotic and symbiotic light organs over the first 60 h. For each condition, n = 15. Data are the mean ± the standard error of the mean. # indicates a significant difference (analysis of variance followed by a Tukey's pairwise comparison; P < 0.05) between aposymbiotic and symbiotic samples. Bar, 25 μm. apo = aposymbiotic; sym = symbiotic.
significant differences in actin cross-reactive sites were detected among the aposymbiotic cohorts. Incubation of animals in secondary antibody alone resulted in no detectable cross-reactivity above background.

Because the antiaactin antibody recognizes both globular and filamentous actin, we used labeling with phalloidin, which recognizes only the latter, at the 48-h time point to determine whether the levels of filamentous actin changed during the onset of the symbiosis. This analysis provided evidence for a twofold increase in filamentous actin in symbiotic animals, suggesting that some of the change in actin protein was due to an increase in filamentous actin (Student’s t test; $P < 0.05$; for an example of phalloidin staining patterns, see Fig. 7, below).

No changes in actin mRNA levels were detectable during the onset of the symbiosis. To determine whether symbiont-induced changes in actin were the result of changes at the level of transcription, we studied actin mRNA levels in whole light organs by real-time RT-PCR and Northern analysis and localized actin mRNA in light organs by in situ hybridization. No significant differences in actin mRNA levels were detected by real-time RT-PCR (Fig. 4A), Northern blot analyses (data not shown), or in situ hybridization in aposymbiotic and symbiotic light organs at 12 h (Fig. 4B). These data revealed that the significant changes in actin protein synthesis at 12 h in symbiotic animals were not coincident with detectable changes in message levels. Because symbiont-induced changes in mRNA levels may have occurred hours before changes in actin protein could be detected, we also performed in situ hybridization to localize actin mRNA in the light organs of 3- and 6-h aposymbiotic and symbiotic animals. No differences in actin mRNA could be detected in any part of the light organs, including the duct epithelia, at these time points (Fig. 4B). The sense probe controls did not bind to the tissues to produce fluorescence above background levels.

Persistent colonization with V. fischeri was required to maintain the changes in light organ actin. To determine whether V. fischeri delivers a signal that directs an irreversible change in the duct epithelia or produces a signal that must be constantly present to maintain the change, we performed experiments in which we cured the light organs after symbiont-induced changes had been induced (Fig. 5). CM was used to cure juveniles of their bacterial symbionts after 12 h of infection; the subsequent abundance of antiaactin cross-reactivity in the duct epithelia at 48 h was not significantly different from those of the aposymbiotic juveniles at that time. CM treatment alone had no apparent pharmacological effect; the intensity of fluorescence due to cross-reactivity in hatchling squid infected with the CM-resistant strain of V. fischeri and subsequently treated

FIG. 5. Constant exposure to V. fischeri was required to maintain the symbiont-induced changes in actin. A representative experiment showing relative fluorescence of cross-reactive sites to antiaactin in the medial ducts of 48-h juvenile light organs is shown. White bars indicate animals that had no exposure to CM. Solid bars indicate animals that were treated continuously with 20 μg of CM/ml in seawater starting at 12 h. Cohorts of animals that had been inoculated with a CM-resistant strain of V. fischeri [ES114(pVO8)] were used to control for effects of CM. For each condition, $n = 15$; data are the mean ± standard error of the mean. A * indicates a significant difference (analysis of variance followed by a Tukey’s pairwise comparison; $P < 0.05$) between aposymbiotic animals (± CM) and symbiotic animals colonized by either wild type (± CM) or CM-resistant (± CM) strains of V. fischeri.
with CM was not significantly different from that of juveniles exposed to the wild-type strain. Taken together, these data demonstrated that constant exposure to *V. fischeri* is required to maintain the symbiont-induced changes in actin observed in the early hours of the symbiosis.

**Increases in duct actin correlated with constriction of the ducts.** We sought to determine whether a corresponding change in the architecture of the duct occurred with the observed increase in actin. Specifically, we asked whether duct circumference, epithelial cell number interfacing the duct space, and epithelial cell shape changed at various time points during the onset of the symbiosis. Commencing at 12 h, the average medial duct circumference of symbiotic animals was significantly smaller than that of aposymbiotic animals (Fig. 6). Antibiotic curing experiments revealed that this change in the ducts required persistent interaction with *V. fischeri*. However, the reversal of duct circumference to aposymbiotic levels after antibiotic treatment at 12 h was not apparent until 60 h, unlike the reversible changes in actin protein levels, which were apparent by 48 h (Fig. 6B).

The average number of cells lining the duct lumina of 48-h symbiotic animals was significantly lower than that of aposymbiotic animals (Fig. 7). Further, in aposymbiotic animals, the duct epithelia were a regular array of parallel columnar cells, whereas the shape of many of the duct cells in symbiotic animals was highly irregular, with narrow apical surfaces. These data suggested that the symbiont-induced decrease in duct circumference is a result of a reduction in the number of cells circumscribing the duct lumen and that reduction is due to displacement of cells away from the population comprising the duct epithelium.

**DISCUSSION**

The results of this study provide evidence that, similar to the action of some pathogenic microorganisms, a beneficial bacterial partner markedly influences the actin cytoskeleton of host cells. Beginning with the first exposure and continuing for the early days of the symbiosis, *V. fischeri* cells induced an increase in actin synthesis in the ducts, the passageway through which the symbionts enter during initial colonization and are expelled each day as part of the natural diel rhythm of the symbiosis. The increase in duct actin correlated with a decrease in the circumference of the duct. Thus, the symbionts enter the tissues and, by some unknown mechanism, narrow the entry way behind them.

Bacterial pathogens or symbionts usually affect the cytoskeletons of the cells with which they directly associate (8, 18). In contrast, the light organ ducts of *E. scolopes* are adjacent to, but not in direct contact with, the population of *V. fischeri* cells.
in the crypts, except at the onset of the symbiosis and with venting each morning. Thus, either these brief, direct interactions are enough to cause the observed changes in the actin cytoskeleton or the products released from the symbiont population in the crypts are swept past the duct epithelial cells by the cilia lining the ducts. Alternatively, these cytoskeletal modifications in the duct cells could be signaled indirectly through the adjacent crypt epithelia, where the bacteria colonize. At present, it is not possible to identify experimentally which of these mechanisms is naturally occurring.

The changes observed in the duct appear to involve a complex array of factors, including an increase in actin synthesis and a reduction of duct circumference, accompanied by changes in duct cell shape and number. One interpretation of these data might be that the increase in actin protein in the apical surfaces of the ducts reflects a condensation or focusing of the preexisting actin pool. However, the evidence presented here supports an absolute change in actin synthesis. First, our 2-D gel data (Fig. 2) demonstrated such an increase, which was only detectable in the apical surfaces of the duct cells (Fig. 3). Second, significant increases in actin protein were first observed at 6 h, before a significant decrease in duct circumference was detectable (Fig. 3 and 6). And third, upon curing, the loss of actin cross-reactivity from the apical surfaces of the cells did not coincide in time with the reversal of duct dimension (Fig. 5 and 6).

We found that the increase in actin protein synthesis was not preceded by an increase in actin mRNA levels. Instead, our data suggest that the symbiont-induced changes are the consequence of posttranscriptional modifications. With rare exception (17), microbe-induced changes in actin gene transcription are unusual, as reported both in focused analyses of host cytoskeleton (for reviews, see references 1, 8, and 18) and in the more comprehensive microarray studies (16, 35, 38). However, it is possible in many such studies that still yet biologically significant changes have been disregarded because the actin gene is often used as a control gene, which may not always be a sound assumption (22).

Actin-mediated changes in tubule dimensions have been noted in the development of tissues of both invertebrates and vertebrates. For example, in the development of the tracheal system in Drosophila melanogaster, similar to the E. scolopes light organ ducts, the dimensions of the tracheal lumina change concomitantly with alterations in actin cytoskeleton in the apical surfaces of the tracheal epithelial cells (2). However, the mechanisms appear to differ. While the data suggest that changes in cell shape and a decrease in cell number are responsible for mediating the constriction of the light organ ducts, the developmental change in the tubules of the Drosophila tracheal system could be accounted for entirely by constriction in the apical cell surfaces of the tubule cells. Similarly, in a study of the development of salamander kidney tubules, a decrease in cell number also did not correlate with alterations in tubule size (6). The extent to which the symbiont-induced developmental program of the squid ducts is unusual, i.e., being mediated by a change in cell number, will require further comparative analyses.

These symbiont-induced developmental changes in the ducts add to the other known biomechanical and biochemical constraints, such as the activity of the cilia lining the duct (25) and the high concentration of toxic oxygen species (36; Davidson et al., Abstr. 101st Gen. Meet. Am. Soc. Microbiol., 2001), that the duct places on entry into host tissues. Further, this modification in the ducts is only one of a suite of effects that result from symbiont-directed shutdown of activities associated with the infection process. Once the symbiont has resided in the crypts for several hours, secretion of mucus from the superficial ciliated field ceases and, thus, further harvesting of potential symbiont cells no longer occurs (32). In addition, the symbionts induce the complete regression of the superficial ciliated field over 4 to 5 days (27). Together, the cessation of harvesting, loss of the superficial field, and constriction of the duct appear to act in concert to render the organ less susceptible to subsequent colonization.

The results of this study present two principal questions: (i) what is the bacterial signal that induces changes in the duct cells, and (ii) by what mechanisms do bacteria change the host cytoskeleton? The bacterial effectors known to induce changes in the cytoskeleton of animal hosts are most often specific toxins (1). The genome sequence of V. fischeri has revealed the presence of a gene encoding an ortholog of RTX, an actin cross-linking toxin in Vibrio cholerae (7), and may reveal orthologs of other toxins known to influence the actin cytoskeleton. Characterizations of the duct cell responses will provide an interesting comparison of the ways in which host cells respond to pathogens and beneficial symbionts. Most often, bacteria are reported to alter preexisting actin protein by directly or indirectly, covalently or noncovalently, modifying the actin cytoskeleton (1, 3, 18). Among other effects, pathogens are known to ADP-ribosylate actin in a covalent modification of the molecule, affect actin nucleation through the ARP 2/3 complex, or modulate actin cytoskeleton through modifying Rho GTPases (1). It is possible that similar molecular responses will be involved in the squid-vibrio system, but because outcomes of pathogenic and beneficial interactions are diametrically opposed, additional or different mechanisms may be operating. The ongoing production of expressed sequence tag libraries from the host light organs, characteristic of different times in the developmental program and of development under symbiotic and aposymbiotic conditions, should be immensely helpful in our quest to understand such complex interactions. With these molecular tools, we should be able to identify candidate genes and characterize their role in the dynamics of the symbiosis between the host squid and its beneficial bacterial partner.

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