



The Variable Internal Structure of the *Mycoplasma penetrans* Attachment Organelle Revealed by Biochemical and Microscopic Analyses: Implications for Attachment Organelle Mechanism and Evolution

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ABSTRACT Although mycoplasmas have small genomes, many of them, including the HIV-associated opportunist *Mycoplasma penetrans*, construct a polar attachment organelle (AO) that is used for both adherence to host cells and gliding motility. However, the irregular phylogenetic distribution of similar structures within the mycoplasmas, as well as compositional and ultrastructural differences among these AOs, suggests that AOs have arisen several times through convergent evolution. We investigated the ultrastructure and protein composition of the cytoskeleton-like material of the *M. penetrans* AO with several forms of microscopy and biochemical analysis, to determine whether the *M. penetrans* AO was constructed at the molecular level on principles similar to those of other mycoplasmas, such as *Mycoplasma pneumoniae* and *Mycoplasma mobile*. We found that the *M. penetrans* AO interior was generally dissimilar from that of other mycoplasmas, in that it exhibited considerable heterogeneity in size and shape, suggesting a gel-like nature. In contrast, several of the 12 potential protein components identified by mass spectrometry of *M. penetrans* detergent-insoluble proteins shared certain distinctive biochemical characteristics with *M. pneumoniae* AO proteins, although not with *M. mobile* proteins. We conclude that convergence between *M. penetrans* and *M. pneumoniae* AOs extends to the molecular level, leading to the possibility that the less organized material in both *M. pneumoniae* and *M. penetrans* is the substance principally responsible for the organization and function of the AO.

IMPORTANCE *Mycoplasma penetrans* is a bacterium that infects HIV-positive patients and may contribute to the progression of AIDS. It attaches to host cells through a structure called an AO, but it is not clear how it builds this structure. Our research is significant not only because it identifies the novel protein components that make up the material within the AO that give it its structure but also because we find that the *M. penetrans* AO is organized unlike AOs from other mycoplasmas, suggesting that similar structures have evolved multiple times. From this work, we derive some basic principles by which mycoplasmas, and potentially all organisms, build structures at the subcellular level.

KEYWORDS *Mycoplasma*, cytoskeleton, electron microscopy, evolution, fractionation, mass spectrometry, transcriptomics

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Mycoplasmas, which are genomically reduced, cell wall-lacking bacteria that live parasitically or commensally in nature but can be cultured axenically, occupy a unique space within the field of bacterial cell polarization that makes them potentially informative concerning mechanisms by which polarization is achieved, as well as the evolutionary origins of such mechanisms. Some mycoplasmas have a differentiated prosthecal tip structure, known as an attachment organelle (AO), at one pole (reviewed in reference 1). The AO houses adhesin proteins for interactions with host cells and surfaces, as well as machinery that carries out unidirectional gliding motility, and appears to be physically associated with the chromosome in some cases (reviewed in reference 2). The interior of the AO contains a complex of structural proteins that serves as a cytoskeletal scaffold for AO assembly (reviewed in reference 2). In the sense that the AO is constructed at a cell pole and is involved in functions associated with polarity, this structure parallels other polar structures of bacteria, such as the *Caulobacter crescentus* stalk (2).

The majority of information regarding the components, assembly, and substructures of mycoplasma AOs comes from studies of two different species, *Mycoplasma pneumoniae* and *Mycoplasma mobile*. These two species are members of two distantly related phylogenetic clusters within the genus (3). Characterization of the structures and proteins that make up the interior of the AO of each of these species has shown them to be very different both ultrastructurally and compositionally, with no AO protein homologs shared between these organisms (4; reviewed in reference 1), suggesting that, like the *C. crescentus* stalk, these structures have evolved independently and recently, compared with other members of their respective phyla.

Mycoplasma penetrans is another species that has a polar AO, representing a third mycoplasma phylogenetic cluster (5–7). *M. penetrans* is best known for its association with AIDS patients, in whom it is commonly found in the urogenital tract, but it has also been isolated from an HIV-negative patient with antiphospholipid syndrome (5, 6, 8). The major focus of studies of *M. penetrans* has been its putative role as an AIDS cofactor (9–12). We showed previously that the *M. penetrans* AO contains material distinct in structure from that of *M. pneumoniae* and *M. mobile* (7). Furthermore, *M. penetrans* lacks any homologs of the AO structural proteins of *M. pneumoniae* or *M. mobile* (4, 13, 14). Special difficulty in working with this organism stems from its genetic intractability, leaving biochemical and cell biological routes as the best options for its study.

The ability to establish cellular polarization and to localize specific biomolecules to non-uniformly distributed subcellular positions is critical for many bacteria. Many fundamental processes, such as DNA segregation, cell division, and placement of specialized structures, rely on morphological polarity as a cue for appropriate spatial organization. The cytoskeleton, which is composed of a network of polymeric proteins, plays an integral role in maintaining the morphological and functional integrity of bacterial cells by establishing polarity (reviewed in reference 15). In addition to the widely phylogenetically distributed MreB and FtsZ, which form dynamic filaments that contribute to the nearly universal bacterial processes of peptidoglycan synthesis and cell division (reviewed in reference 16), there is increasing evidence of a diverse array of other cytoskeletal elements, with narrow phylogenetic distribution, that facilitate more recently evolved functions (reviewed in reference 17). In contrast to the highly dynamic MreB and FtsZ, many of these proteins form static structures. The α -helical coiled-coil-rich, intermediate filament-like protein crescentin from *C. crescentus* is associated with the establishment of cell shape (18, 19). Many bacterial scaffold proteins contain an abundance of coiled-coil domains, which are important for oligomerization and close protein-protein interactions and are referred to as coiled-coil-rich proteins (CCRPs) (reviewed in reference 20). CCRPs serve an array of functions, such as maintaining cell morphology and enabling cell motility, often by participating in polar localization of proteins (reviewed in reference 19). The widely distributed bactofilins are characterized by the conserved DUF583 domain and form stable nucleotide-independent filaments that serve as spatial landmarks (21, 22). Other bacterial cytoskeletal proteins include DivIVA and PopZ, which assemble into 2- or 3-dimensional

arrays at extant or nascent cell poles and interact with proteins involved in pole-associated processes (reviewed in reference 19). It is essential to understand processes that rely on bacterial cell polarization, and this understanding can be achieved by establishing an inventory of cytoskeletal proteins across bacterial phylogeny and determining their characteristics and how these properties enable them to carry out their functions.

We sought to identify the proteins of the interior of the *M. penetrans* AO and to examine their properties in relation to other proteins of mycoplasma AOs. Because mycoplasma AOs seem to have evolved in a convergent manner, we hypothesized that the proteins involved in the structure and assembly of the *M. penetrans* AO would be distinct from those of other mycoplasmas, and we predicted that their composition would reveal important information about the properties of the AO. To test these ideas, we investigated the protein content of the detergent-insoluble material of *M. penetrans*, which constitutes the AO interior, and characterized the structure of this material. We found that, although many of the proteins have some properties resembling those of the corresponding proteins of *M. pneumoniae*, the *M. penetrans* AO is quite distinct from its counterparts in other mycoplasmas, consistent with convergent evolution of AOs and of the mechanisms of polarization of bacterial cells. We also examined changes in gene expression of *M. penetrans* upon incubation with human cervical epithelioid HeLa cells, to test whether AO and other genes were subject to transcriptional regulation upon encountering the host, and we found that they were not.

(This work was performed in partial fulfillment of the doctoral dissertation requirements of S. L. Distelhorst.)

RESULTS

Internal structure of *M. penetrans* AO. As described previously, *M. penetrans* cells are elongated (Fig. 1B), with a polar tip structure that functions as an AO. The AO confers attachment and gliding motility, which contributes to cell division; dividing cells often take the form of two cell bodies linked by a membranous connecting filament (7) (Fig. 1A). *M. penetrans* cells attach to host cells by means of a pole, whose interior is distinctly differentiated from the cytoplasm of the rest of the cell despite being adjacent to it, as established previously by transmission electron microscopy (TEM) (5, 6). Treatment of *M. penetrans* cells attached to plastic with the nonionic detergent Triton X-100 results in solubilization of a large amount of cellular material, leaving behind discrete detergent-insoluble objects with a range of dimensions, as visualized by scanning electron microscopy (SEM) (Fig. 1C) (7). Unlike the AO structures obtained similarly from species of the *M. pneumoniae* cluster (23), treatment with DNase had no effect on the appearance of these objects (data not shown), indicating that they did not contain significant amounts of DNA.

Close examination of these Triton X-100-insoluble (TXI) structures revealed that, despite considerable heterogeneity in size and shape, they generally consisted of a wider, irregular, ball-like object from which emanated one or more rod-like filaments, which were often periodically punctuated with other material along the filaments. The structures had an average length of 320 ± 100 nm ($n = 70$) (Fig. 2A). This range of lengths was significantly $\sim 15\%$ smaller than that of the polar nucleoid-free zones of *M. penetrans* cells ($n = 84$; $P < 0.001$) (Fig. 2), as revealed by overlaying phase-contrast images and images of the DNA stained with 4',6-diamidino-2-phenylindole (DAPI) (Fig. 2G) (7). The mean length of the nucleoid-free zones was 380 ± 140 nm. In cells with single nucleoid-free zones, there was a positive correlation between the length of the cell and the length of the nucleoid-free zone ($n = 52$; $P < 0.0001$) (Fig. 2D), suggesting that growth of this structure is coupled to cell growth. In cells with two such zones at each pole, the length of the smaller one was correlated with cell length ($n = 16$; $P = 0.04$) (Fig. 2E), whereas that of the larger one was not ($n = 16$; $P = 0.52$) (Fig. 2F), consistent with the larger one being a preexisting structure whose net growth had slowed or stopped.

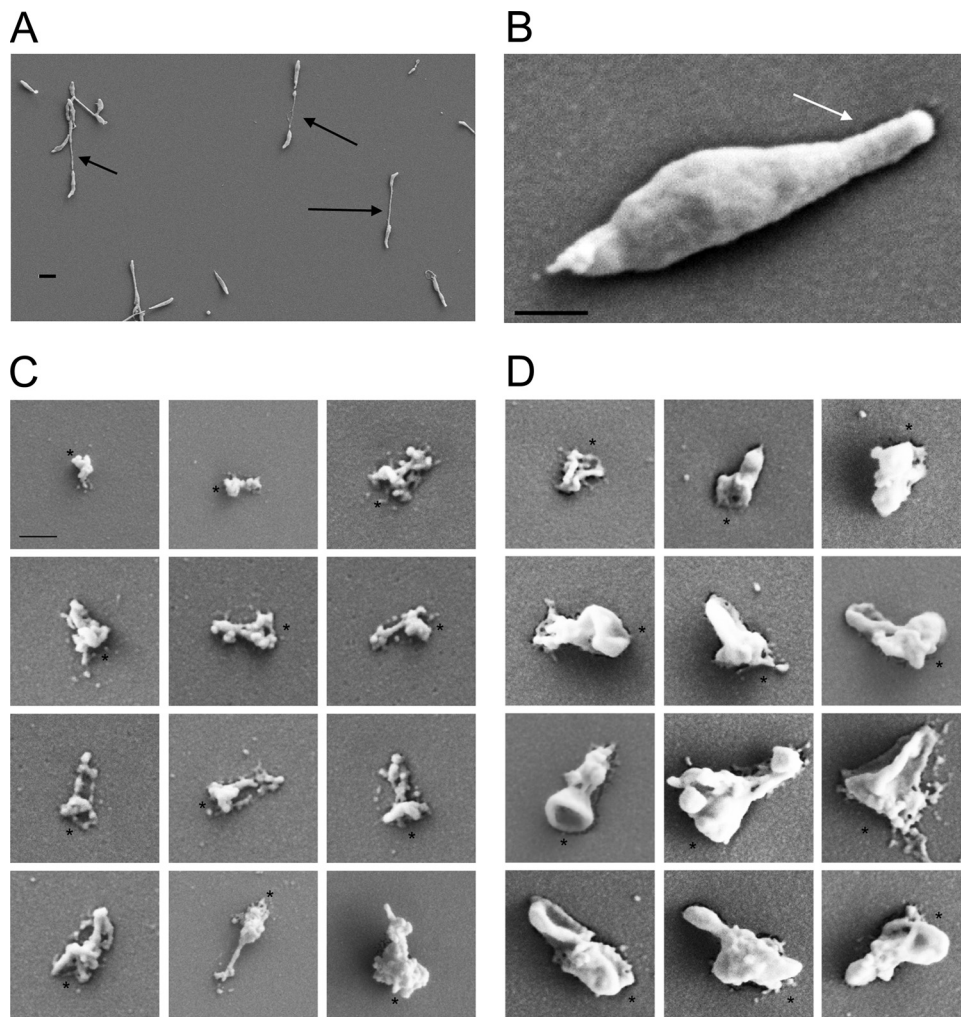


FIG 1 SEM images of *M. penetrans* whole cells and detergent-insoluble structures. (A) Field of *M. penetrans* cells. Arrows, connecting filaments. Scale bar, 1 μ m. (B) Individual *M. penetrans* cell. Arrow, AO. Scale bar, 200 nm. (C and D) Detergent-insoluble structures from *M. penetrans* cells extracted with Triton X-100 (C) or Tween 20 (D). Asterisks, wide ends of structures. Scale bar, 200 nm.

To determine whether another nonionic detergent might yield similar structures, we used SEM to examine the structures that remained following extraction of cells grown under identical conditions with the chemically distinct Tween 20 (Fig. 1D). These objects were generally similar to those obtained using Triton X-100, although they were larger and had a generally smoother appearance. Indeed, their mean length of 380 ± 130 nm was identical to that of the nucleoid-free zones ($n = 79$; $P > 0.8$) (Fig. 2C). The variability in the sizes and shapes of the detergent-insoluble objects was recapitulated in electron cryotomography (ECT) images, which showed ribosome-containing zones in the central area of the cell that were distinct from ribosome-free zones at both cell poles. These ribosome-free zones had irregular shapes and boundaries (Fig. 3A and B), often extending into the filament connecting cells undergoing division (Fig. 3C) (7). Given the similarities in size distributions, previous TEM images of similar material at *M. penetrans* poles (5, 6, 24), and the limited amount of space in an *M. penetrans* cell for large objects, we concluded that the polar nucleoid-free zones visualized with DAPI staining, the polar ribosome-free zones visualized with ECT, and the DNA-lacking, detergent-insoluble objects visualized with SEM are the same objects, with Triton X-100 solubilizing some component that Tween 20 does not. Interestingly, whereas the poles of whole *M. penetrans* cells, as visualized with SEM (Fig. 1B) or light microscopy (Fig. 2G),

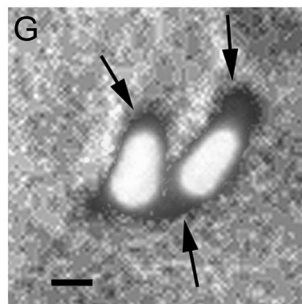
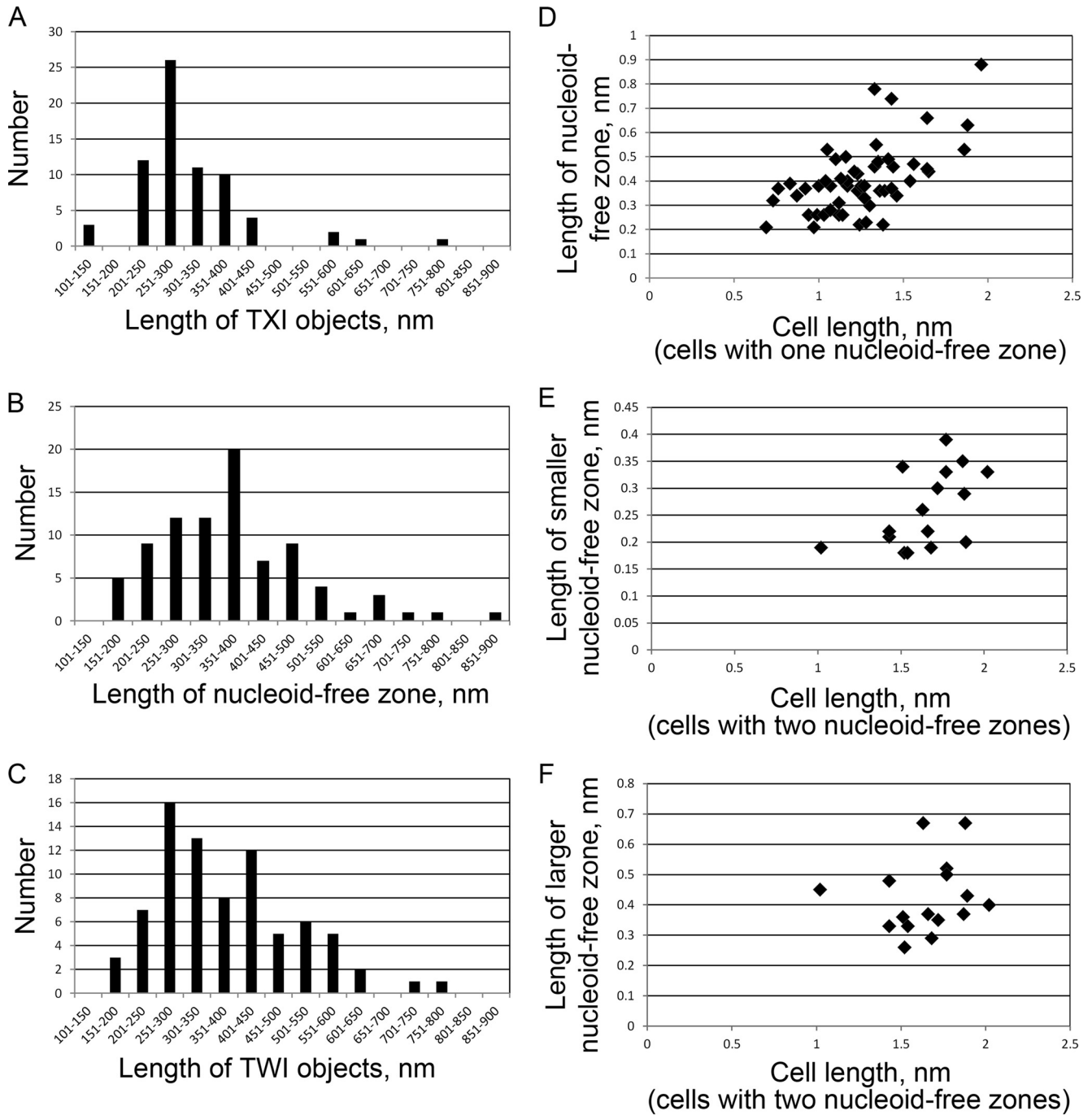


FIG 2 AO-associated objects and their lengths. (A) Distribution of the lengths of TXI objects observed by SEM. (B) Distribution of the lengths of nucleoid-free zones observed by light and fluorescence microscopy. (C) Distribution of the lengths of TWI objects observed by SEM. (D) Correspondence of the lengths of
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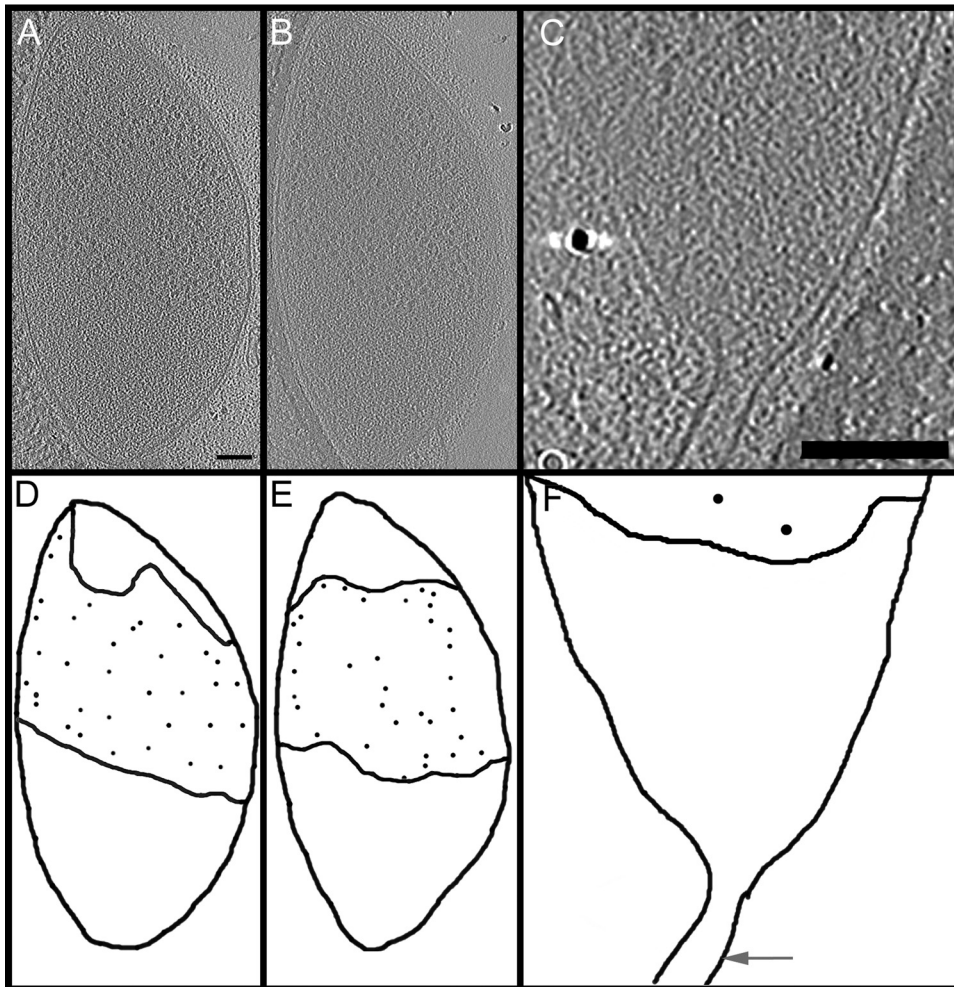


FIG 3 Internal organization of *M. penetrans* observed by ECT. (A to C) Sections of different whole cells. (D to F) Corresponding schematics. In panels A to C, ~20-nm granules are ribosomes (reviewed in reference 70). The insertion site of a connecting filament was observed at one cell pole (C and F). Black lines, cell membranes and boundaries of ribosome-free zones; circles, ribosomes (positions are schematic and do not represent actual ribosomes); arrow in panel F, location of connecting filament. Scale bars, 100 nm.

were distinctly narrower than the cell bodies, there was no abrupt narrowing at the poles in cells imaged by ECT, with the cells exhibiting a more ovoid shape (Fig. 3).

Identification and characterization of detergent-insoluble proteins. *M. penetrans* lacks homologs of the AO cytoskeletal proteins of other mycoplasmas (14). To determine which proteins make up the TXI and Tween 20-insoluble (TWI) material, we examined the protein profiles of whole-cell lysates, TXI proteins, and TWI proteins using SDS-PAGE. The detergent-insoluble material was generated by extracting attached *M. penetrans* cells in flasks and scraping the material that remained attached to the flasks, ensuring that it was identical to the material that was visualized by SEM, which consisted almost entirely of structures similar in shape but heterogeneous in size, as described above (Fig. 1C and D). Several proteins were prominent in the TXI and TWI fractions, and some were enriched, compared to the whole-cell lysate (Fig. 4). Consistent with the similar appearances of the structures, the profile of the TWI proteins was

FIG 2 Legend (Continued)

nucleoid-free zones to the fractions of the cell length occupied by the nucleoid-free zones in 52 cells with a single nucleoid-free zone. (E and F) Correspondence of the lengths of nucleoid-free zones to the fractions of the cell length occupied by the nucleoid-free zones in 16 cells with two nucleoid-free zones, a shorter one (E) and a longer one (F). (G) Example of nucleoid-free zones. Black, cell outline viewed by phase-contrast microscopy; white, nucleoid stained with DAPI. Arrows, nucleoid-free zones, measured along the long axis of the cell for panel B. Scale bar, 500 nm.

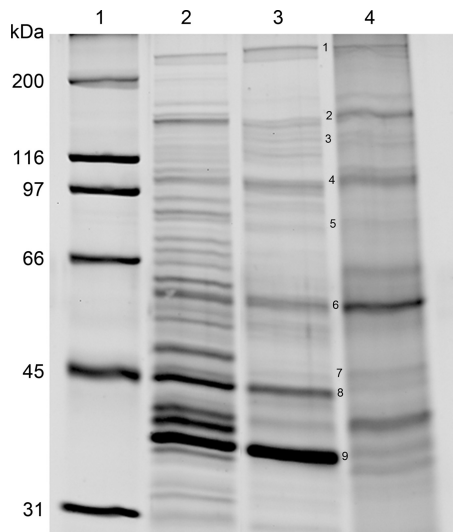


FIG 4 SDS-PAGE of the *M. penetrans* whole-cell lysate and TWI and TXI proteins. A total of 22 μg each of protein from the whole-cell lysate (lane 2), the TWI fraction (lane 3), and the TXI fraction (lane 4) was separated by SDS-PAGE for comparison and for excision of candidate bands chosen for identification. Lane 1 contains protein standards, with sizes indicated to the left. The 66-kDa band in lane 4 was present only in some preparations. Numbers in the gel indicate specific bands chosen for MALDI-TOF mass spectrometry and correspond to protein identifications in Table 1.

similar to that of the TXI proteins, with the principal exception being one prominent TWI band migrating at ~ 40 kDa that was absent in the TXI fraction (Fig. 4). In general, the profiles featured mostly the same proteins but often in different relative amounts.

We selected eight bands from SDS-polyacrylamide gels that appeared to be enriched in both detergent-insoluble fractions, compared to the whole-cell lysate, as well as the one band that was present in the TWI fraction but not the TXI fraction, for identification using matrix-assisted laser desorption ionization–time of flight (MALDI-TOF) mass spectrometry (Fig. 4). The 12 proteins that were identified were associated with a variety of functional characteristics, including four with well-established biochemical roles, three putative lipoproteins, and five of unknown function (Table 1). The four with characterized biochemical roles were lactate dehydrogenase, ribosomal protein S3, pyruvate dehydrogenase subunit E2, and OppF, an oligopeptide ABC transporter ATP-binding protein. Two of the three putative lipoproteins were identified as P35 and P42, both members of the P35 lipoprotein family; P42 was the protein that was present only in the TWI fraction. The third lipoprotein was a homolog of *Mycoplasma genitalium* G37 MG309. Of the five identified proteins that were uncharacterized, four were encoded by genes found very close to one another in the genome,

TABLE 1 TXI and TWI proteins identified by MALDI-TOF

Protein	Band no. in Fig. 4	Functional annotation	Size (kDa)
MYPE1530	3	NA ^a	122
MYPE1550	1	NA	386
MYPE1560	4	NA	99
MYPE1570	4	NA	93
MYPE4000	2	NA	99
MYPE5100	6	Pyruvate dehydrogenase subunit E2, PdhC	51
MYPE6630	8	P42 lipoprotein (P35 family)	42
MYPE6810	9	P35 lipoprotein (P35 family)	38
MYPE6960	3	Lipoprotein	138
MYPE8750	5	Oligopeptide ABC transporter ATP-binding protein, OppF	92
MYPE9640	9	Lactate dehydrogenase, Ldh	34
MYPE10090	7	Ribosomal subunit S3, RpsC	48

^aNA, not applicable.

TABLE 2 Comparison of AO protein features

Species and protein	Coiled coil	Molecular mass of >70 kDa	Extreme pI (<5 or >9)	Total no. of features
<i>M. pneumoniae</i>				
HMW1 (MPN447)	X	X	X	3
HMW2 (MPN310)	X	X	X	3
HMW3 (MPN453)		X	X	2
P24 (MPN312)				0
P41 (MPN311)	X		X	2
P65 (MPN309)	X		X	2
P200 (MPN567)		X	X	2
TopJ (MPN119)		X	X	2
<i>M. mobile</i>				
MMOB0150				0
MMOB1620				0
MMOB1630	X		X	2
MMOB1640				0
MMOB1650		X		1
MMOB1660				0
MMOB1670				0
MMOB4530			X	1
MMOB4860			X	1
MMOB5430			X	1
<i>M. penetrans</i>				
MYPE1530	X	X	X	3
MYPE1550	X	X	X	3
MYPE1560	X	X	X	3
MYPE1570	X	X	X	3
MYPE4000	X	X	X	3
MYPE5100				0
MYPE6630				0
MYPE6810				0
MYPE6960		X	X	2
MYPE8750	X	X	X	3
MYPE9640				0
MYPE10090				0

MYPE1530, MYPE1550, MYPE1560, and MYPE1570. Together with MYPE4000, they shared no specific sequence homology with other proteins except in a close relative of *M. penetrans*, namely, *Mycoplasma iowae* (25); however, the first four were labeled as predicted cytoskeletal proteins in the original annotation of the *M. penetrans* genome, on the basis of predicted extensive α -helical coiled-coil structures, reminiscent of *Mycoplasma pneumoniae* AO protein HMW2 (14), which is predicted to constitute a major structural element within the *M. pneumoniae* AO cytoskeleton (26–28).

The absence of sequence homology of these five proteins to the proteins of the AO cytoskeleton of *M. pneumoniae* (28) or the analogous structure in *M. mobile* (29), together with the disparity in the detergent-insoluble structures between *M. penetrans* and mycoplasmas from other phylogenetic clusters, is consistent with a model in which attachment organelles arose independently multiple times during the course of mycoplasma evolution (7, 30). We examined whether there were general features shared in common among the cytoskeletal proteins of *M. penetrans*, *M. pneumoniae*, and *M. mobile*. Interestingly, the five putative *M. penetrans* cytoskeletal proteins have molecular masses of ≥ 93 kDa, isoelectric points of ~ 4.5 , and predicted α -helical coiled-coil regions (extensive in all except MYPE4000). Many of the proteins of the *M. pneumoniae* attachment organelle, but not of the tip structure of *M. mobile*, share these features (Table 2).

Cotranscription of putative cytoskeletal genes. Examination of the genomic organization of the genes encoding proteins MYPE1530, MYPE1550, MYPE1560, and MYPE1570, as well as predicted structural and compositional similarities among the predicted protein products, suggested that, along with the genes encoding the struc-

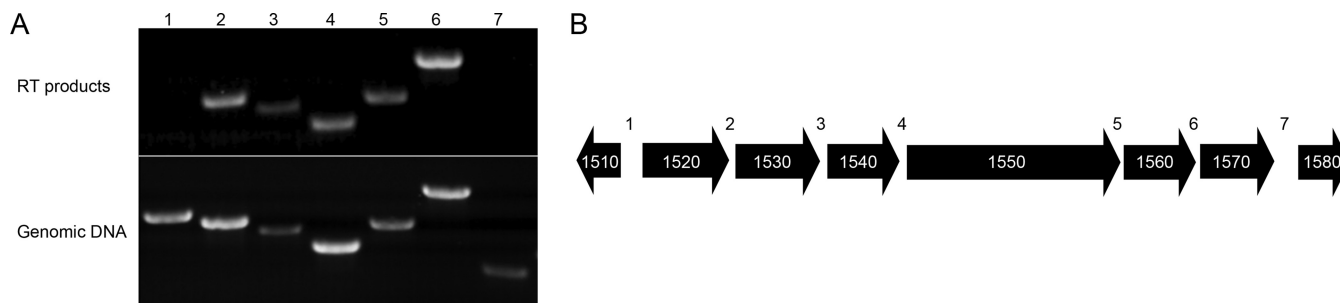


FIG 5 RT-PCR analysis of the putative cytoskeletal operon of *M. penetrans*. (A) Agarose gel containing PCR products from RT-generated transcripts that span the gene junctions numbered in panel B. Top, products amplified from reverse-transcribed RNA using a primer complementary to a region at the 3' end of *mype1570*; bottom, products amplified from genomic DNA used as a positive control. (B) Schematic of genes encoding MYPE1520 to MYPE1570, showing the genomic orientation and relative sizes, including the 5' and 3' flanking genes. Numbers 1 to 7 refer to the positions of the primer pairs used to span gene junctions for RT-PCR.

turally and compositionally similar proteins MYPE1520 and MYPE1540, they might form a transcriptional unit. We performed reverse transcription (RT)-PCR to test whether these genes, each separated by no more than 30 bp, were cotranscribed. We used a primer complementary to a region near the 3' end of MYPE1570, which is positioned downstream of the other genes in question, to create cDNA. From this cDNA, we amplified regions spanning the junctions between the six candidate genes (MYPE1520 to MYPE1570) as well as MYPE1510, which is ~180 bp away from MYPE1520 and on the opposite strand. We also made cDNA using a primer complementary to MYPE1580, which is also ~180 bp away, and we attempted to amplify cDNA linking MYPE1570 and MYPE1580. There were RT-PCR products for all of the junctions between MYPE1520 and MYPE1570 but no products for the MYPE1510-MYPE1520 junction (Fig. 5A). Amplification of the junction between MYPE1570 and MYPE1580 sometimes yielded a faint, poorly reproducible product, suggesting partial read-through (data not shown). These results suggest that MYPE1520 to MYPE1570 constitute an operon (Fig. 5B).

Transcriptomic analysis of *M. penetrans* grown with HeLa cells. Because the major function of the AO is adherence to host cells, which is expected to occur at the initial stages of infection, we hypothesized that the genes involved in AO formation and function, including those encoding the proteins identified in the TXI fraction, would be upregulated during the early stages of infection. The interactions between *M. penetrans* and HeLa cells, including the events that occur during invasion, have been documented (31–34). To identify genes that undergo differential expression upon interaction with host cells, we performed transcriptome sequencing (RNA-Seq) with *M. penetrans* cells grown alone and grown in the presence of HeLa cells, 6 h after infection (representing a time point at which cytoadherence and invasion had occurred [33]), and we compared global gene expression profiles between the two conditions. Surprisingly, there was almost no statistically significant difference in gene expression between the two conditions, including the AO genes, with the exception of three genes that were upregulated and one gene that was downregulated 2-fold (see Data Set S1 in the supplemental material). The three upregulated genes were MYPE1035, MYPE3985, and MYPE8260, all of which encode small hypothetical proteins of less than 70 amino acids. The single downregulated gene was MYPE20130, which codes for a tRNA for arginine and whose corresponding codon is predicted to be commonly used; its expression was 3-fold lower in mycoplasmas incubated with HeLa cells.

DISCUSSION

Convergent evolution of attachment organelles. The overall organization of the AO-associated structures of *M. penetrans* was quite distinct from those of *M. pneumoniae* (35, 36), *Mycoplasma mobile* (29), and *Mycoplasma insons* (30), which represent three other phylogenetic lineages or clusters within the genus. Despite superficial similarities in the morphology and function of the AOs of *M. pneumoniae* and *M. mobile*, dissimilarities in both the organization and the composition of their internal structures

are well documented (37). In contrast, the structures and compositions of AO cytoskeletal structures are shared among the close relatives of *M. pneumoniae* in the *M. pneumoniae* cluster (23). These differences have been taken to suggest that the AOs of *M. pneumoniae* and *M. mobile* arose through convergent evolution (2), which is not altogether surprising, considering that the two species are separated by one of the deepest phylogenetic divisions within the genus, with *M. pneumoniae* being in the pneumoniae group and *M. mobile* in the hominis group (3). However, the subsequent finding that *Mycoplasma insons* (a species in the pneumoniae group that is not a close relative of *M. pneumoniae*) is unidirectionally motile but lacks a distinct tip structure and instead contains cytoskeletal elements throughout its entire rod-shaped body (30) raised the possibility that even within the pneumoniae group there has been independent evolution of polar attachment and motility. If so, then these structures have an even more recent evolutionary origin. Alternatively, the *M. insons* organization could represent a variation of the *M. pneumoniae* organization whose homology is not obvious, pending further studies of this organism. *M. penetrans* is also in the pneumoniae group but within a different cluster than either *M. pneumoniae* or *M. insons* (38). The AO of this species, whose motility is driven by a fundamentally different mechanism than that of *M. mobile* (39), provided an opportunity to test whether there were detectable conserved elements of AO organization within the pneumoniae group.

The absence of homology between *M. penetrans* and *M. pneumoniae* AO proteins strongly suggests that these parallel systems evolved independently. *M. mobile*, with no apparent equivalent of a ribosome-free space (29) and with its cytoskeletal components lacking the hallmarks of the *M. pneumoniae* and *M. penetrans* AO proteins identified in the present work, appears to have evolved a different approach to generating a protrusive and therefore superficially similar structure. *M. insons* achieves polarization of adherence and motility without a protrusive structure (30), representing a completely different model for polarization.

Properties of *M. penetrans* AO interior. As described previously (7), there can be little doubt that the detergent-insoluble, DNA-free objects observed by SEM, whose abundance on coverslips is routinely parallel to that of unextracted *M. penetrans* cells grown from the same inoculum (data not shown), occupy the space in the nucleoid-free zones of *M. penetrans* cells and represent the material observed at the cell poles in thin sections and also observed here by ECT. Indeed, the material underlying the cell poles in ECT images, which closely resemble the material at the cell poles observed in TEM images of thin sections (5, 6, 24), are strikingly reminiscent of ribosome-free PopZ-containing structures at the poles of *C. crescentus* cells when PopZ is overproduced (40). Our deeper study of these objects has revealed that the organization of the TXI and TWI structures of the *M. penetrans* AO is completely unlike that of the *M. pneumoniae* structures. Whereas the dimensions of the *M. pneumoniae* structures appear to be tightly constrained (23), the dimensions are highly variable in *M. penetrans*, as visualized in a number of ways. SEM images of these variable objects revealed some commonalities, however, including a wider portion and a narrower portion, the latter often containing one or more flexible rod-like elements. This heterogeneity, which is distinct from the great regularity of the *M. pneumoniae* AO structures, suggests the possibility of growth by accretion of material onto, or absorption of material into, a preexisting structure. ECT images suggest that the two poles contain the same differentiated material, and this material even extends into the filaments connecting dividing *M. penetrans* cells. These facts are consistent with a model in which nucleation and biogenesis of the AO underlying structure are linked to the cell cycle, perhaps with new material being deposited into spaces in the cytoplasm vacated during chromosome condensation and segregation. The broader size distribution of the nucleoid-free zones and the detergent-insoluble structures suggests that growth of the *M. penetrans* AO, though capped, is not capped at a uniform size.

In addition to the heterogeneity in size, the pleomorphy of the detergent-insoluble objects is striking. We noted that, whereas our SEM images of whole *M. penetrans* cells

and previously described TEM images of thin sections (5–7, 24) showed that the region of the cell at the pole is constricted along the long axis, this was not the case in ECT images. One possible interpretation of this discrepancy is that the material within the *M. penetrans* AO is highly hydrated and therefore subject to shrinkage during the dehydration steps associated with standard SEM and TEM processing techniques, whereas ECT lacks a dehydration step. Conceivably, this hydration results from attraction of water molecules to the highly negatively charged proteins present in the structure. If the structural elements are principally organized parallel to the long axis, then dehydration would result predominantly in a decrease in width rather than length. Another possibility is that, because the cell bodies visualized by ECT are not attached firmly to surfaces but instead are present in the holes of the carbon-coated grids, they are in a relaxed conformation. In either case, we predict that the material within the AO is flexible like a gel.

***M. penetrans* AO composition and gene expression.** Of the proteins we identified in the TXI and similar TWI fractions of *M. penetrans* by MALDI-TOF mass spectrometry, four have characterized biochemical roles. Whether any or all of them are part of the internal structure of the AO is unclear. Pyruvate dehydrogenase subunit E2 normally functions as part of a large structure that could be insoluble in nonionic detergents; however, none of the other subunits of pyruvate dehydrogenase was identified, suggesting a possible alternative or additional role for this protein, which also functions as a surface adhesin in other mycoplasmas (41). OppF is normally a peripheral membrane component of the oligopeptide ABC transporter (42), including in mycoplasmas (43). The substrate-binding protein component of this transporter, OppA, doubles as an adhesin in *Mycoplasma hominis* (44, 45), raising the possibility that *M. penetrans* OppF anchors the material underlying the AO to OppA, acting as an adhesin. However, none of the proteins of this complex other than OppF was identified in the *M. penetrans* detergent-insoluble fractions.

Surprisingly, three lipoproteins were identified in the detergent-insoluble fraction. Two of these lipoproteins are members of the P35 lipoprotein family, which comprises 44 total *M. penetrans* genes (14). The P35 lipoprotein and its paralogs, which are distributed across the surface of *M. penetrans* cells, are immunodominant (46–49). The expression of genes encoding these proteins undergoes extensive antigenic variation, which is thought to be a major factor for immune evasion (50). The presence of two of these P35 homologs in the detergent-insoluble fractions may suggest that they indirectly interact with the AO interior, although it is also possible, given their extensive distribution across the cell surface, that they aggregate or form multiple interactions with each other that prevent their solubilization in nonionic detergents. Indeed, the difference in appearance between the TWI and TXI fractions might be entirely attributable to the presence of P42 in the former. The third lipoprotein identified, MYPE6960, has no known function, and its distribution and presence at the cell surface of *M. penetrans* cells is unknown. However, this protein is a homolog of MG309, a protein in *M. genitalium* that binds and activates Toll-like receptors (51), making MYPE6960 a particularly interesting candidate for further study with respect to pathogenesis. Conceivably, this lipoprotein plays a role in the adherence of *M. penetrans* to host cells and is associated with AO proteins.

The remaining five proteins that were identified in this study had no predicted function or conserved domains, making them ideal candidates for principal structural elements of the AO. The genes encoding four of these proteins are located very close together in the genome of *M. penetrans* and were found to be transcribed as a polycistronic message with two other genes. Three of these, MYPE1550, MYPE1560, and MYPE1570, were among the most abundant proteins in the detergent-insoluble fractions. All six of these proteins, as well as MYPE4000, share an unusual set of characteristics, including predicted α -helical coiled-coil regions, molecular masses of >90 kDa, and pI values of approximately 4.5 (Table 2). Interestingly, these characteristics are shared with many of the *M. pneumoniae* AO cytoskeletal proteins (Table 2) despite the

lack of sequence homology. It is highly unlikely, given that the *M. penetrans* proteins lack characteristics of *M. pneumoniae* AO proteins such as proline-rich domains and sequence elements such as the EAGR box, that the *M. pneumoniae* and *M. penetrans* AO proteins share a common ancestry. The shared properties are intriguing, however, when the functional analogy of the structures that they build is considered. These shared properties suggest that one solution to forming a polar cytoskeletal structure, arrived at independently in *M. pneumoniae* and *M. penetrans*, involves the assembly of large, highly charged proteins that interact with each other over an extensive surface area via α -helical coiled coils.

Although another group reported changes in the expression of a specific set of phase-variable genes in response to infection of a host (52), this study is the first, to our knowledge, to report on the expression of an entire mycoplasma transcriptome upon exposure to host cells. Particularly striking was the absence of major differences in the transcript profiles of *M. penetrans* cells grown independently and grown in the presence of HeLa cells for 6 h, which is enough time for interactions to occur between the two cell types, including invasion (31, 33). We anticipate that, after 6 h in an environment filled with signals from host cells, *M. penetrans* cells will have had ample opportunity to adjust their gene expression profiles to prepare for interactions, whether or not all *M. penetrans* cells have adhered to and invaded the HeLa cells. Therefore, even if, as a consequence of the use of HeLa cells at 70 to 80% confluence, only ~75% of *M. penetrans* cells interacted with host cells, we would expect to see significant changes in the overall gene expression pattern if a change were occurring. The absence of transcriptional regulation is a formal but unlikely possibility, as mycoplasmas, despite having limited machinery for transcriptional regulation, nonetheless exhibit it (53–55). A more likely explanation is that *M. penetrans*, which naturally occurs as an obligate commensal or parasite like all mycoplasmas, has no evolutionary adaptation to living in the absence of host cells and therefore is always in a mode in which it is deployed for interaction with the host. Relatedly, it is possible that the fetal bovine serum endows SP-4 broth with all of the relevant factors provided by HeLa cells.

Implications for mechanisms of AO function. Unlike the *M. penetrans* AO, which consists entirely of a poorly organized matrix, the corresponding *M. pneumoniae* feature is visually dominated by a dense, highly organized structure flanked by a ribosome-excluding zone in contact with the cell membrane (35). The structure, which has been called the electron-dense core because of its prominent staining with metal-based dyes (56), has been the focus of considerable study (reviewed by Balish [2]). Because the ribosome-excluding zone is directly adjacent to the core, even in cells in which the core is detached from the membrane (35, 57, 58), it is likely formed by some fine substance linked to the core, probably regions of the same proteins that make up the core itself. Indeed, the failure to identify any non-core-associated proteins in the *M. pneumoniae* AO (58) supports this model. In contrast to *M. pneumoniae*, the *M. penetrans* AO consists solely of material whose lack of obvious organization is more reminiscent of the lucent area of *M. pneumoniae* than the core. Because both AOs, one with a core and one without, function in adherence and motility, it is possible that the core itself is dispensable for these functions in *M. pneumoniae*, as suggested previously (23, 59), although this assertion is in opposition to some models (35, 58). Therefore, we propose that the less organized material, constituting the ribosome-excluding zone in *M. pneumoniae* and perhaps the entire AO interior in *M. penetrans*, is the principal organizer of the AO, driving the concentration and organization of adhesins at the AO in both species and leaving motility to be carried out directly by adhesins with motor properties, as described for the more distantly related *M. mobile* (37).

Regarding *M. pneumoniae*, if it is the lucent material that organizes the AO, then the core may be dispensable for adherence and motility. Superficially, this statement appears to be counter to observations (reviewed in reference 60). However, if the lucent and core regions of the *M. pneumoniae* AO are actually composed of different regions of the same proteins, then mutants in which core proteins are lost would also be

defective in the functions associated with the lucent region. Indeed, if this model is correct, then *M. pneumoniae* mutants affected only in the lucent area and not in the core have never been studied. The physical association of DNA with the core in *M. pneumoniae* relatives (23) raises the possibility that the *M. pneumoniae* core itself is primarily involved in chromosome segregation. In *M. pneumoniae*, a newly formed AO is displaced from the preexisting pole to the pole created during binary fission during cell division (61), possibly carrying the newly replicated chromosome with it to the daughter cell (2). Because *M. penetrans* cells appear to acquire a mass of AO material opposite the pole of the functioning AO, and not adjacent to it, during the process of division and because DNA is not associated with the *M. penetrans* detergent-insoluble material, it is reasonable to propose that the *M. penetrans* AO is not directly involved in chromosome segregation.

MATERIALS AND METHODS

Bacterial culture. Hyperadherent *Mycoplasma penetrans* strain HP88 cells (39) were grown to the mid-log phase at 37°C in plastic tissue culture flasks (Thermo Scientific BioLite) containing SP-4 broth (62).

Detergent extraction and protein analysis. Cultures of *M. penetrans* were decanted and washed twice with warm potassium-free phosphate-buffered saline (PBS) (145 mM NaCl, 2.83 mM NaH₂PO₄·H₂O, 7.20 mM NaHPO₄·7H₂O [pH 7.2]). For detergent extraction, Triton X-100 or Tween 20 in 20 mM Tris-HCl-150 mM NaCl (pH 7.2) was added to the cultures to a final concentration of 1% in 25 ml PBS. Detergent extracts were then incubated for 30 min at 37°C, scraped, and centrifuged for 20 min at 17,050 × *g*. Whole-cell lysates were treated in the same way without the addition of detergent. Following centrifugation, the pellets were washed three times and resuspended in PBS. Protein concentrations were determined using a bicinchoninic acid protein assay kit (Pierce). Equal amounts of protein of each fraction were subjected to 9% SDS-PAGE (63) and stained with Coomassie brilliant blue or Sypro Ruby (Molecular Probes). Detergent-extracted proteins were separated by SDS-PAGE, and select bands, chosen on the basis of being abundant or enriched, compared to the total-cell lysate, were excised, reduced and alkylated, digested with trypsin, and analyzed by MALDI-TOF mass spectrometry (AB Sciex 4800 MALDI-TOF/TOF analyzer) at the University of Cincinnati Proteomics Laboratory, as described previously (64). MultiCoil (65) was used to test predicted amino acid sequences for the presence of coiled-coil domains; the presence of regions with >50% probability of forming α -helical coiled coils was recorded as positive.

Scanning electron microscopy. Cells were prepared for SEM as described previously (66). Briefly, cells were grown on glass coverslips for 6 to 24 h at 37°C in SP-4 medium supplemented with 3% gelatin. For examination of TXI structures or TWI structures, detergent in 20 mM Tris-HCl (pH 7.5)-150 mM NaCl was added to cells at a final concentration of 1%, and cells were incubated for 30 min at 37°C. The samples were fixed for 30 min at room temperature in 1.5% glutaraldehyde-1% paraformaldehyde-0.1 M sodium cacodylate (pH 7.2). Following fixation, the coverslips were washed four times in 0.1 M sodium cacodylate (pH 7.2) and dehydrated through a series of ethanol washes from 25% to 100%. The coverslips were then critical point dried, gold sputter coated, and viewed on a Zeiss Supra 35 FEG-VP scanning electron microscope at the Miami University Center for Advanced Microscopy and Imaging.

Electron cryotomography. *M. penetrans* was grown at 37°C on Quantifoil Au-finder grids (Quantifoil Micro Tools GmbH) in SP-4 broth to the mid-log phase, which was indicated by a color change in the medium from red to orange. The electron microscopy grids with attached *M. penetrans* cells were removed from the medium, loaded onto tweezers, and washed with fresh medium. Colloidal gold (10 nm) was added to the grids, and the grids were blotted (Whatman grade 40 filter paper) prior to being plunge-frozen in liquid ethane on a gravity plunger freezer. Grids were loaded into a 300-kV FEI Polara G2 electro-cryotransmission electron microscope equipped with a field emission gun, a lens-coupled Gatan UltraCam (4,000- by 4,000-pixel resolution), and a Gatan imaging filter (GIF). Samples were maintained at liquid nitrogen temperature as tilt series of whole cells were captured. The tilt series were recorded from -60° to +60° at 1° increments, with a 10- μ m defocus, using Legikon (67). All tilt series were collected through the GIF around zero-loss energy with a slit width of 20 keV, for a cumulative dose of 180 e⁻/Å² for each tilt series.

Measurements and analysis. To measure the lengths of nucleoid-free zones, *M. penetrans* cells were grown overnight on coverslips, fixed as described previously (7), and mounted on slides with Vectashield containing DAPI (Vector Laboratories). Fields were imaged using a 100× objective, as described previously (7). Phase-contrast and DAPI fluorescence images were overlaid, and the lengths of 84 polar nucleoid-free zones were measured using SPOT software. To measure the lengths of TXI and TWI objects, SEM images of the objects from multiple fields, captured as described above, were measured along their long axes. Objects that appeared to be clusters of individual objects were excluded. For TXI objects, 70 were measured; for TWI objects, 79 were measured. Statistical comparisons of lengths were performed by one-way analysis of variance (ANOVA), after a Shannon-Wilk test was used to establish that the distributions of lengths were not normal. Statistical comparisons of the lengths of the cells and the nucleoid-free zones were performed using least-squares regression analysis.

RNA extraction. RNA was harvested as described previously (52). *M. penetrans* cells were harvested by centrifugation at 17,400 × *g* for 20 min, followed by three washes in PBS. The cell pellet was resuspended in 1 ml of TRI Reagent solution (Ambion). Total RNA was then purified using the Direct-zol

TABLE 3 Primers used for RT-PCR

Primer designation	Sequence (5' to 3')	Target	Product size (bp)
1510end(up)	AATGAGCAGTAAGTGCTAGC	MYPE1510-MYPE1520	603
1520begin(down)	GTCTTAGGAATTTGAACAGGTG		
1520end(up)	GGAAAAAGAAGCTGTACACAG	MYPE1520-MYPE1530	566
1530begin(down)	GGTTGATTGTCAGCAGAACC		
1530end(up)	TTCCAGCTCCTGCATATGAC	MYPE1530-MYPE1540	527
1540begin(down)	GTCAGTTCCTGAAATTGTTT		
1540end(up)	CTAAATTGTTCCAAATTTGAATAATCGC	MYPE1540-MYPE1550	454
1550begin(down)	ATAGATGAGATTCTTTATCAAAGTTCTC		
1550end(up)	ATCAAACCCCAAGCGATTCTG	MYPE1550-MYPE1560	541
1560begin(down)	TCAACATCGTCTTGTCTTGG		
1560end(up)	TGCTCCTAGTAGTGAGTTGC	MYPE1560-MYPE1570	696
1570begin(down)	CTTCTCTAGCTTGAAGTTGG		
1570end(up)	TTTGAAGACCAGCAGGAAG	MYPE1570-MYPE1580	380
1580begin(down)	GTGTTGCCAGGTCTAAGATC		
1580RT(down)	CATAGCCGTTTGAACAGTGT	NA ^a	NA

^aNA, not applicable.

RNA MiniPrep kit (Zymo Research), according to the manufacturer's instructions. The remaining DNA was removed from the sample by treatment with DNase I twice using the DNA-free kit (Applied Biosystems), according to the manufacturer's instructions. Quantitative PCR was performed to ensure that the RNA was free of DNA. The quality of the RNA was then examined using an Agilent 2100 bioanalyzer (Agilent Technologies).

RNA sequencing. *M. penetrans* cells were grown for 6 h at 37°C in SP-4 broth, either alone or together with HeLa cells grown to 70 to 80% confluence, at a multiplicity of infection (MOI) of 100. Three biological replicates were used for each of the conditions. For each biological replicate, RNA was extracted as described above and divided into three technical replicates. Purified bacterial RNA was enriched by performing poly(A) depletion using the NEBNext poly(A) mRNA magnetic isolation module (New England BioLabs) on all samples, as described previously (52). The RNA was then depleted of both prokaryotic and eukaryotic rRNA by using the Ribo-Zero Gold rRNA removal kit (Illumina), as directed by the manufacturer's instructions, and was cleaned and concentrated by using the Zymo RNA Clean and Concentrator-25 kit (Zymo Research), as described previously (52). The RNA was eluted in 25 μ l RNase-free water, examined for quality using an Agilent 2100 Bioanalyzer (Agilent Technologies), and used for cDNA synthesis.

cDNA libraries were created using the Illumina TruSeq stranded mRNA library preparation kit (Illumina), following the manufacturer's protocol, as described previously (52). The cDNA libraries were examined for quality using the Agilent 2100 Bioanalyzer and then were quantified using a library quantification kit (Kapa Biosystems). Libraries were normalized to 10 nM, pooled, diluted, denatured, and loaded onto a MiSeq reagent kit v3 cartridge, as directed by the manufacturer (Illumina). The samples were sequenced on a MiSeq system using a 75-bp paired-end approach.

Data from the RNA-Seq experiment were analyzed as described previously for *Mycoplasma gallisepticum* data (52). Briefly, the fastq files were assembled, mapped, and analyzed for differential gene expression using Rockhopper (<http://cs.wellesley.edu/~btjaden/Rockhopper>) (68), with the *M. penetrans* HF-2 genome as the reference genome. Parameters were as described previously (52). For comparison of gene expression values, the data were first normalized by determining the ratio of reads per kilobase per million (RPKM) for mycoplasmas grown with HeLa cells to those for mycoplasmas grown axenically and then the fold change was calculated by taking the log₂ transformation of the RPKM difference between the two biological samples.

Reverse transcription-coupled PCR. The RT reaction was carried out as described previously for *Mycoplasma iowae* (69). Briefly, cDNA synthesis was performed with 100 ng of total RNA from *M. penetrans* and random hexamers, using the Verso cDNA synthesis kit (Thermo Scientific), according to the manufacturer's instructions. One microliter of cDNA was then used as the template in a 50- μ l PCR mixture with *Taq* polymerase (New England BioLabs), using EasyStart PCR tubes (Molecular Bioproducts), according to the manufacturer's instructions. The primers used in each reaction were designed to span the adjacent gene junctions (Table 3). Reactions were amplified by incubation at 94°C for 2 min, followed by 30 cycles of 94°C for 30 s, 53°C for 1 min, and 68°C for 1 min, with a final extension at 68°C for 5 min. The PCR products were subjected to gel electrophoresis. Reactions with genomic DNA and without the addition of reverse transcriptase or template were performed as positive and negative controls, respectively.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/JB.00069-17>.

SUPPLEMENTAL FILE 1, XLS file, 0.6 MB.

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