Host’s guardian protein counters degenerative symbiont evolution

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Microbial symbioses significantly contribute to diverse organisms, where long-lasting associations tend to result in symbiont genome erosion, unculturability, extinction, and replacement. How such inherently deteriorating symbiosis can be harnessed to stable partnership is of general evolutionary interest. Here, we report the discovery of a host protein essential for sustaining symbiosis. Plataspid stinkbugs obligatorily host an uncultivable and genome-reduced gut symbiont, Ishikawaella. Upon oviposition, females deposit “capsules” for symbiont delivery to offspring. Within the capsules, the fragile symbiotic bacteria survive the harsh conditions outside the host until acquired by newborn nymphs to establish vertical transmission. We identified a single protein dominating the capsule content, which is massively secreted by female-specific intestinal organs, embedding the symbiont cells, and packaged into the capsules. Knockdown of the protein resulted in symbiont degeneration, arrested capsule production, symbiont transmission failure, and retarded nymphal growth, unveiling its essential function for ensuring symbiont survival and vertical transmission. The protein originated from a lineage of odorant-binding protein-like multifunctional family, shedding light on the origin of evolutionary novelty regarding symbiosis. Experimental suppression of capsule production extended the female’s lifespan, uncovering a substantial cost for maintaining symbiosis. In addition to the host’s guardian protein, the symbiont’s molecular chaperone, GroEL, was overproduced in the capsules, highlighting that the symbiont’s degrading functionality is compensated for by stabilizer molecules of host and symbiont origins. Our finding provides insight into how intimate host–symbiont associations can be maintained over evolutionary time despite the symbiont’s potential vulnerability to degeneration and malfunctioning.

Microbial symbioses are ubiquitously found in nature, intricately interwoven into the adaptation, ecology, and evolution of almost all life forms (1, 2). In diverse animals encompassing invertebrates and vertebrates, specific microorganisms are present on their body surface, in their alimentary tract, within their body cavity, or even inside their cells (3). In obligate and long-lasting host–symbiont associations, the symbiont genomes often exhibit drastic size reduction and massive gene losses, which are attributed to relaxed natural selection acting on many symbiont genes unnecessary for the intrahost life as well as to irreversibly accumulated deleterious mutations due to strong population bottleneck and restricted horizontal gene acquisitions imposed by continuous vertical transmission (4, 5). Due to the accumulated genetic load, such degenerative symbionts should suffer cellular malfunctioning and instability, which would finally lead to symbiont genome erosion, unculturability, extinction, and/or replacement (4–7). In this context, both host and symbiont are expected to evolve some molecular mechanisms for stabilizing the symbiotic association that is inherently vulnerable to degeneration and malfunctioning. For the symbiont side, many genome-reduced endosymbiotic bacteria were reported to overproduce molecular capacitors such as heat shock proteins and molecular chaperones, thereby stabilizing molecular structure and functioning of the degenerative bacterial cells (8–10). For the host side, by contrast, little has been known about such molecular mechanisms underpinning the stability of symbiosis. Here, we report the discovery of such a key molecule essential for sustaining symbiosis.

Results and Discussion

Survival of Genome-Eroded Symbiont in Mother-Made Capsules. Stinkbugs of the family Plataspidae host an uncultivable, genome-reduced (~0.7 Mb), and nutrition-provisioning (=essential amino acids) bacterial symbiont Ishikawaella in a specialized region of the midgut (11–13). Upon oviposition, adult females deposit “symbiont capsules,” each containing around 107 Ishikawaella cells (14, 15), with eggs on plant surfaces (Fig. 1 A and B), where the symbiont capsules are exposed to intense sunshine for 7 to 10 d until the eggs hatch and the newborn nymphs orally acquire the symbiont (12, 15). Here, it should be noted that Ishikawaella cells must be fragile because, like the 0.6 Mb reduced genome of the aphid endosymbiont Buchnera (11, 16), the Ishikawaella genome is devoid of major cell wall–related genes. How such a genome-reduced symbiont can survive the harsh...
Fig. 1. Gut symbiotic system of M. punctatissima and PMDP. (A) An adult female of M. punctatissima laying eggs on a host plant bud. (B) Underside view of an egg mass. Yellow arrowheads indicate symbiont capsules. (C and D) Dissected alimentary tracts of adult male (C) and female (D). (E) Gene and structure of PMDP. (F and G) Expression levels of PMDP in dissected tissues (F) and in whole insects at different developmental stages (G). Values of PMDP complementary DNA (cDNA) copies per Rpl19 cDNA copy obtained by qPCR are regarded as relative expression levels of PMDP gene. For dissected tissues, the average value for SSM is shown as 1, whereas for whole insects, the average value for adult females is shown as 1. The number on the top indicates the number of samples analyzed. Different alphabetical letters (a, b, and c) indicate statistically significant differences (Tukey honestly significant difference [HSD] test: \( P < 0.05 \)). (H and I) SDS-PAGE of whole proteins (H) and immunoblot detection of PMDP (I) from dissected tissues of adult females. Red arrowheads indicate PMDP bands. M1, midgut first section; M2, midgut second section; M3, midgut third section; M4, midgut fourth section with crypts (= symbiotic midgut); M4B, M4 bulb.

and fluctuating environment outside the host and successfully establish transmission to the next generation is mysterious.

In plataspid and other stinkbugs, their alimentary tract is differentiated into structurally distinct regions, of which the posterior M4 region is specialized for hosting symbiotic bacteria (17, 18) (Fig. 1C). In adult females of plataspid stinkbugs, notably, the voluminous M4 region was further differentiated into specialized sections: main symbiotic midgut (MSM), a long stretch of symbiotic region; and enlarged end midgut (EEM), a dark-colored enlarged region (Fig. 1D). Previous studies regarded these midgut regions as female-specific, capsule-producing organs (12–14), but it has been unknown what genes and molecules are functioning there.

A Single Host Protein, Posterior Midgut Dominant Protein, Predominant in Symbiont Capsules. We performed RNA sequencing of these midgut regions and found that the majority of transcripts represented a single gene in the female-specific regions MSM and EEM in all plataspid species we examined (SI Appendix, Table S1). In Megacopta punctatissima, for example, the transcript was 829 bp in size, contained a 729-bp open-reading frame, and encoded a protein of 243 amino acid residues, in which the following domains were identified: a secretion signal sequence at the N terminus, an intrinsically disordered region with no predicted stable secondary structure, and an odorant-binding protein (OBP)-like domain at the C terminus with sequence similarity to OBPs of various insects (Fig. 1E). qRT-PCR confirmed highly specific and predominant expression of the gene in SSM and EEM of adult females (Fig. 1F and G). Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotting detected a 28-kDa predominant protein band specifically in SSM, EEM, and symbiont capsules (Fig. 1H and I and SI Appendix, Fig. S1A). Liquid chromatography-mass spectrometry (LC-MS), Southern blotting, and qPCR verified that the 28-kDa protein is derived from the gene that is single copy in the insect genome (SI Appendix, Figs. S1 B–D). The protein accounted for the majority of total proteins in the symbiont capsules (Fig. 1H and SI Appendix, Fig. S1), which is hereafter referred to as posterior midgut dominant protein (PMDP). Here, we note that, besides the host-derived PMDP, the symbiont-derived molecular chaperone GroEL was the second most abundant protein in the symbiont capsules (SI Appendix, Fig. S1E), whose extreme overproduction has been observed in many genome-reduced endosymbiotic bacteria for stabilizing degenerative protein structure and functioning (8, 10).

Symbiont Cells Embedded in PMDP and Packaged into Capsules. We histologically investigated the detailed processes of capsule formation in the female-specific midgut organs, which uncovered structural and functional specialization of the female’s alimentary tract for production of symbiont capsules: MSM and SSM retain the symbiotic bacteria, SSM and EEM produce and supplement the PMDP-containing secretion to the symbiotic bacteria, and EEM excretes
Fig. 2. Formation process of symbiont capsules in female’s posterior midgut organs. (A) Whole sectioned image of the female’s posterior midgut organs for production of symbiont capsules. Many microscopic images are combined to obtain the whole picture. PMDP is immunohistochemically visualized in dark brown, whereas cell nuclei are counterstained in purple. Ishikawaella cells are not stained but cytologically recognized. The morphologically and functionally differentiated regions, MSM, SSM, CJM (constricted junction midgut), EEM, and hindgut, are highlighted by different frame colors. (B) The MSM–SSM junction. The symbiont cells from MSM are combined with the PMDP-containing secretion from the epithelial cells of SSM. (C) Magnified image of the symbiont cells mixed with the PMDP-containing secretion in SSM. (D) Magnified differential interference contrast (DIC) image of the symbiont–secretion mixture in SSM, in which round symbiont cells are seen within the PMDP-positive secretion. (E) Transmission electron microscopic (TEM) image of the PMDP-secreting epithelium of SSM. Well-developed microvilli, electron-dense PMDP-containing secretion, and the symbiont cells (asterisks) are seen. (F) TEM image of the PMDP-secreting epithelium of SSM. Well-developed microvilli, electron-dense PMDP-containing secretion, and the symbiont cells (asterisks) are seen. (G) The SSM–CJM junction. (H) Magnified image of CJM, whose narrow inner cavity is filled with a homogeneous mixture of the symbiont cells and the PMDP-containing secretion. (I) The CJM–EEM junction. The upstream folds of EEM (highlighted in J) are wide, secreting much PMDP with cuticular sheets, whereas the downstream folds of EEM (highlighted in L) are narrow and deep, mainly secreting cuticular sheets with some PMDP. The main duct is voluminous, storing a large amount of the symbiont–secretion mixture. (J) Magnified image of the upstream folds of EEM, which secrete much PMDP with cuticular sheet. (K) Magnified DIC image of an upstream fold cavity of EEM. (L) Magnified image of a downstream fold of EEM, secreted cuticular sheets in the fold cavity are seen. (M) TEM image of a downstream fold of EEM. Secreted cuticular sheets in the fold cavity are seen. (N) EEM-hindgut junction in the process of capsule formation. A mass of the symbiont–secretion mixture surrounded by the cuticular shell is being squeezed out to form a symbiont capsule. (O) Structural and functional differentiation of the posterior intestinal regions for production of symbiont capsules in adult plataspid females. ep, epithelium or epithelial cell; fc, fold cavity; md, main duct; mp, Malpighian tubule. Yellow arrowheads indicate the border of the intestinal regions, whereas red arrows indicate the flow direction in the intestinal main duct.
the cuticular layered shell and stores the capsule components consisting of the symbiont-secretion mixture and the capsule shell. Passing through the serially arranged gut regions, the symbiotic bacteria are embedded in the PMDP-containing secretion and finally packaged into the symbiont capsules (Fig. 2 and SI Appendix, Fig. S2 and Note S1). Symbiont cytometry and in vitro culturing of the dissected gut regions with $^{15}$N-labeled glutamine revealed that the symbiont’s metabolic activities are high in MSM, low in SSM, and almost undetectable in EEM (Fig. 3), indicating that the symbiont cells are transformed into a dormant status during the capsule formation process.

**PMDP Knockdown Stops Capsule Production.** Considering the abundant supplementation and intimate packaging of PMDP with the symbiont cells in the symbiont capsules, some important biological functions were suspected for the protein. Hence, we performed RNA interference (RNAi) knockdown of PMDP expression. When adult females were injected with double-stranded RNA (dsRNA) representing a partial PMDP gene region (see SI Appendix, Fig. S1B), the PMDP gene expression was suppressed by two orders of magnitude within 3 d, and the suppression continued for over 2 wk (SI Appendix, Fig. S3A). The RNAi knockdown of PMDP did not affect the number of the eggs, the clutch size of the egg masses, and the hatch rate of the eggs (SI Appendix and Fig. 3 B–D) but drastically suppressed the capsule production. Upon laying egg masses, the control adult females deposited around one capsule per three eggs, where the capsules were large in size and dark in color, throughout the experimental period for 2 wk (Fig. 4 A, Top and Fig. 4 B–D and Movie S1). By contrast, the PMDP-knockdown females started to produce abnormal capsules, which were smaller in size and paler in color, around 7 d after dsRNA injection (Fig. 4 A, Middle). Coincidently, the number of capsules started to decline (Fig. 4 B and C). Subsequently, size and number of the capsules became smaller and smaller. Around 18 d after dsRNA injection and on, no or only a few tiny translucent capsules were produced by the PMDP-knockdown females (Fig. 4 A, Bottom and Fig. 4 B and C). These females laid eggs normally but deposited few capsules, thereby producing the egg masses devoid of the symbiont capsules (Fig. 4E and Movie S2).

**PMDP Knockdown Causes Symbiont Degeneration.** Detailed histological inspection of the midgut organs provided insight into how and why the PMDP-knockdown females stopped producing the symbiont capsules. The dark-colored material in the main tract of SSM and the whole EEM, which represented the capsule components produced and stored in these organs, disappeared following the PMDP dsRNA injection (Fig. 4F, “Dissected capsule-producing midgut” column). On the day of dsRNA injection as well as in the control insects, the symbiont cells were morphologically normal; in the fold cavities of SSM, the symbiont cells were densely packed, whereas in the main tract of SSM and within EEM, the symbiont cells were embedded in the PMDP-containing secretion (Fig. 4F, “0 day after injection” row). Three days after dsRNA injection, the PMDP-containing secretion decreased, and the symbiont cells looked degenerate in the main duct of SSM and in EEM (Fig. 4F, “3 days after injection” row). Two weeks after dsRNA injection, the PMDP-containing secretion almost disappeared in SSM and EEM, and the symbiont cells were highly degenerate in the main duct of SSM and completely disintegrated in EEM (Fig. 4F, “14 days after injection” row). It is notable that, deep in the fold cavities of SSM where little PMDP was secreted, the symbiont degeneration was not conspicuous (Fig. 4F, “Fold cavity” column), suggesting a connection between the PMDP depletion and the symbiont degeneration.

**PMDP Knockdown Disrupts Symbiont Transmission, Affects Nymphal Behavior, and Retards Nymphal Growth.** How does the maternal RNAi knockdown of PMDP, which causes symbiont degeneration and arrested capsule production, affect the next generation of the host insect? The eggs laid by the PMDP-knockdown females hatched normally, like the eggs laid by the control females (SI Appendix, Fig. S3D). However, the hatchlings exhibited strikingly different behaviors between the treatment groups. The newborn nymphs from the egg masses of the control females immediately sucked the symbiont capsules for around 1 h and then aggregated and became quiescent near the eggshells as previously reported (19) (Fig. 5A and Movie S3). By contrast, the newborn nymphs from the egg masses of the PMDP-knockdown females became neither aggregated nor quiescent but wandered around and dispersed (Fig. 5B and Movie S4). Plausibly, the nymphs could not acquire the symbiotic bacteria from the egg masses without intact symbiont capsules and thus continued to search for them. This idea was supported by quantification of symbiont titers in the nymphal insects. From the egg masses with intact symbiont capsules, the nymphs acquired symbiotic bacteria from the egg masses and could become normal and aggregated (Fig. 5C, Left). From the egg masses with small pale capsules (Fig. 4A, Middle), the nymphs acquired very low levels of the symbiotic bacteria from the egg masses without intact symbiont capsules and thus continued to search for them. This idea was supported by quantification of symbiont titers in the nymphal insects. From the egg masses with normal dark capsules (Fig. 4A, Top), the nymphs acquired considerable titers of the symbiotic bacteria, which were comparable to the symbiont titers in the control nymphs (Fig. 5C, Middle). From the egg masses with no or a few tiny translucent capsules (Fig. 4A, Bottom), the nymphs acquired no symbiotic bacteria (Fig. 5C, Right). In accordance with the symbiont transmission failure, the nymphs from the egg masses

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**Fig. 3.** Symbiont morphology and metabolism during capsule formation process. (A) Symbiont cell sizes in MSM, in the fold cavity of SSM, in the main duct of SSM, and in EEM measured on transmission electron microscopic images. Different letters (a, b, and c) indicate statistically significant differences (Tukey HSD test; $P < 0.05$). (B–D) Ligated and dissected preparations of MSM (B), SSM (C), and EEM (D) for the metabolic measurement. (E) Metabolic activity of symbiont cells evaluated by synthesis of essential and semi-essential amino acids, which was analyzed by in vitro culturing and LC-MS of dissected MSM, SSM, and EEM in the presence of $^{15}$N-glutamine. Averages and SEs of four replicate assays are shown.
Fig. 4. Effects of PMDP RNAi on capsule production and symbiotic bacteria. (A) Egg masses and symbiont capsules produced by PMDP-knockdown females: top row, a control egg mass, with normal capsules dark brown in color; middle row, an egg mass produced about 10 d after PMDP dsRNA injection, with abnormal small pale capsules; bottom row, an egg mass produced about 20 d after PMDP dsRNA injection, with abnormal tiny translucent capsules; left column, underside images of whole egg masses; and right column, magnified images of symbiont capsules indicated by arrowheads. (B) Time course of arrested capsule production in PMDP-knockdown females: (Top) number of normal capsules per egg; (Middle) number of abnormal capsules per egg; and (Bottom) total number of capsules per egg. Each colored symbol indicates an egg mass: magenta, egg masses laid by PMDP dsRNA-injected females; blue, egg masses laid by β-lactamase dsRNA-injected control females; and green, egg masses laid by control females without dsRNA injection. (C) Statistical test for arrested capsule production in PMDP-knockdown females. From left to right are shown the data of 1 to 7, 8 to 14, 15 to 28, and 29 to 42 d after injection, in which the number of capsules was significantly suppressed by PMDP dsRNA injection except for the first interval (1 to 7 d) (Tukey HSD test, \( P < 0.001 \)). n.s., not significant. (D) A control female laying an egg mass with symbiont capsules. Also see Movie S1. (E) A PMDP-knockdown female laying an egg mass devoid of symbiont capsules. Also see Movie S2. (F) Effects of PMDP RNAi on symbiotic bacteria and PMDP-containing secretion in the capsule-producing midgut organs. ep, epithelial cell; fc, fold cavity; md, main duct.
laid by the PMDP-knockdown females exhibited drastic growth retardation (Fig. 5D–G). Notably, when the egg masses devoid of the symbiont capsules laid by the PMDP-knockdown females were experimentally supplemented with normal symbiont capsules, the nymphs restored symbiont acquisition and growth (Fig. 5C, D, and H). These results indicate that RNAi knockdown of PMDP causes symbiont degeneration and arrested capsule production, which disrupt vertical transmission of the essential symbiont and, consequently, result in fatal fitness defects of the offspring.

**Trade-Off between Capsule Production and Maternal Survival.** Notably, the experimental adult females exhibited intriguing mortality patterns. While the control females died almost constantly throughout the experimental period, the PMDP-knockdown females stopped dying around 10 d after dsRNA injection (Fig. 5I), which was coincidental with the timing of when the capsule production started to decline in the PMDP-knockdown females (Fig. 4B). The significantly lower mortality of the PMDP-knockdown females in comparison with the control females continued until around 36 d after dsRNA injection (Fig. 5I). These patterns suggest that the capsule production may incur substantial cost for the egg-laying females. In other words, the reproductive females are producing the symbiont capsules for their offspring at the expense of their own survival, which highlights a significant parent–offspring trade-off mediated by investment in successful vertical transmission of the genome-eroded symbiont.
Fig. 6. Molecular, structural, and evolutionary features of PMDP. (A) Various plataspisid stinkbugs (Top) and their symbiont capsules (Bottom, arrowheads). (B) Aligned PMDP sequences of six plataspisid species representing three genera. In the OBP-like region, deduced secondary structures including six cysteine residues are shown. Support probabilities are indicated at each node in the order of maximum likelihood/maximum parsimony/neighbor-joining. Sequence accession numbers are in brackets. Note that the PMDP phylogeny is unrooted due to the absence of appropriate outgroup sequences. (C) A structural model for PMDP predicted by C-QUARK (24). Colors indicate the average prediction errors: 3, blue; 10, red; and 17, yellow. (D) A comparison of PMDP phylogeny and mitochondrial phylogeny. Support probabilities are indicated at each node in the order of maximum likelihood/maximum parsimony/neighbor-joining. Sequence accession numbers are in brackets. Note that the PMDP phylogeny is unrooted due to the absence of appropriate outgroup sequences. (E) Comparison of PMDP phylogeny and mitochondrial phylogeny. Support probabilities are indicated at each node in the order of maximum likelihood/maximum parsimony/neighbor-joining. Sequence accession numbers are in brackets. Note that the PMDP phylogeny is unrooted due to the absence of appropriate outgroup sequences.
Evolutionary Origin of PMDP. All stinkbug species of the family Plataspidae thus far examined were reported to deposit symbiont capsules upon oviposition (12, 20), which was also confirmed by our observation in this study (Figs. 1 A and B and 6A). Meanwhile, no other stinkbug groups exhibit such a peculiar trait of symbiont capsule production (21, 22). When and how, then, has PMDP evolved in the evolutionary course of plataspid stinkbugs? A comparison of PMDP sequences among six plataspid species representing three genera showed that the N-terminal signal peptide region and the C-terminal OBP-like region were conserved, whereas the disordered region was not (Fig. 6B). The PMDP phylogeny and the mitochondrial gene phylogeny exhibited perfect congruence in topology (Fig. 6C), which strongly suggests that PMDP was acquired by the common ancestor of extant plataspid species and has evolved in parallel with speciation and diversification of the Plataspidae. The computational structural calculation of PMDP predicted the OBP-like region as a globular domain consisting of six α-helices, typical of OBPs (23), and the disordered region as a polypeptide stretch without a fixed three-dimensional structure (24) (Fig. 6D). The OBP-like region of PMDP exhibited some sequence similarity (<36% for amino acid sequences) to OBP-like sequences of other insects, including the stinkbugs in the databases. Meanwhile, the disordered region of PMDP yielded no significant hits, and no OBP-like sequences from other insects including stinkbugs accompanied a stretch of the PMDP-like disordered polypeptide. These observations suggest that PMDP is a newly evolved OBP-like protein that acquired an extra disordered segment in the lineage of the Plataspidae. Many OBP-like genes were identified in the transcriptomic/genomic data of the plataspid stinkbugs and other stinkbug species (SI Appendix, Fig. S4 and Table S2), illustrating the dynamic evolution of OBP-allied multigene families as observed in Drosophila and other insects (25, 26). While conventional OBPs were reported to function in antennae, bind to small hydrophobic odorant/pheromone molecules, and mediate chemosensory perception, most OBPs and allied proteins are not restricted to olfactory tissues and probably involved in...
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6. G. M. Bennett, N. A. Moran, Heritable symbiosis: The advantages and perils of an established. We speculate that PMDP may bind to the symbiont survival outside the host is still to be established. We speculate that PMDP may bind to the symbiont surface and stabilize the fragile bacterial cell, which should be verified by biochemical and microbiological experiments using purified or recombinant PMDP protein. Besides PMDP, minor proteinous and other capsule ingredients may also have important biological roles, which deserve future studies. The evolutionary origin of the symbiont capsule as well as the female-specific capsule-producing organ is still an enigma. The stinkbug family Plataspidae embraces some 530 species and 56 genera in the world (31), and a wider survey of the world’s plaspid diversity would further light on the origin and evolution of PMDP and the symbiont capsule. Finally, we point out the possibility that mass production of PMDP and an understanding of its molecular function would lead to novel approaches to cultivation and/or preservation of fastidious microorganisms (7, 32).

Materials and Methods

Laboratory-maintained strains and wild-caught individuals of the common plaspid stinkbug M. punctatissima were mainly used in this study. The laboratory strains were reared on soybean plants (Glycine max), pea pods (Pisum sativum), or kidney bean pods (Phaseolus vulgaris) at 25 °C under a long-day condition of 16 h light and 8 h dark. The field-collected insects were either immediately used for experiments, preserved in an ultracold freezer at –80 °C until use, or reared as described above during experiments. M. punctatissima and other plaspid stinkbugs used in this study are listed in SI Appendix, Table S3. More detailed methods are described in SI Appendix, S1 Materials and Methods.

Data Availability. Nucleotide sequence data have been deposited in DNA Database of Japan (PMDP gene sequences, LC516888–LC516893 (33); RNase reads, KR222736–KR222737 (34); OBP-like gene sequences, ICQ01000001–ICQ01000008, ICQ01000001–ICQ01000004, and ICQ01000001–ICQ01000024) (35). All other study data are included in the article and/or supporting information.

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