



Minimal fermentative metabolism fuels extracellular symbiont in a leaf beetle

Eugen Bauer¹ · Martin Kaltenpoth¹ · Hassan Salem^{2,3}

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Abstract

While genome erosion is extensively studied in intracellular symbionts, the metabolic implications of reductive evolution in microbes subsisting extracellularly remain poorly understood. *Stammera capleta*—an extracellular symbiont in leaf beetles—possesses an extremely reduced genome (0.27 Mb), enabling the study of drastic reductive evolution in the absence of intracellularity. Here, we outline the genomic and transcriptomic profiles of *Stammera* and its host to elucidate host-symbiont metabolic interactions. Given the symbiont's substantial demands for nutrients and membrane components, the host's symbiotic organ shows repurposing of internal resources by upregulating nutrient transporters and cuticle-processing genes targeting epithelial chitin. Facilitated by this supplementation and its localization, *Stammera* exhibits a highly streamlined gene expression profile and a fermentation pathway for energy conversion, sharply contrasting the respiratory metabolism retained by most intracellular symbionts. Our results provide insights into a tightly regulated and metabolically integrated extracellular symbiosis, expanding our understanding of the minimal metabolism required to sustain life outside of a host cell.

The study of reductive genome evolution in bacterial endosymbionts has yielded valuable insights into the minimal gene set necessary to sustain life [1, 2]. Many insects harbor such microbial symbionts with reduced genomes, which complement the metabolism of their host through the biosynthesis of essential amino acids and vitamins [3]. Vertical transmission of these beneficial microbes often inflicts severe population bottlenecks, inducing rapid

genomic changes due to genetic drift [4]. As such, neutral or slightly deleterious mutations can become fixed, leading to the AT-biased genomic composition and a degenerated codon usage [3], as well as severely reduced biosynthetic capabilities. Consequently, intracellular symbionts require extensive nutritional support from the host, blurring the line between organelles and autonomous organisms [1].

While the metabolic implications of reductive genome evolution are well-characterized in intracellular symbionts, our understanding of the corresponding selective pressures in extracellular bacteria is limited. Similar to their intracellular counterparts, some extracellular symbionts are restricted to a defined environment within symbiotic organs, relaxing selection to maintain many biosynthetic pathways [5]. Concordantly, recent studies have described a few extracellular insect symbionts with reduced genomes [5, 6], which represent interesting cases to study the general patterns governing reductive genome evolution. However, the extent of this loss and the corresponding metabolic consequences are generally thought to be less extreme than in intracellular symbionts, as evident by their disparate genome sizes [7, 8].

Recently, we reported on an extracellular symbiont with a drastically reduced genome that encodes for the smallest gene set of any organism subsisting outside of a host cell

These authors contributed equally: Martin Kaltenpoth, Hassan Salem

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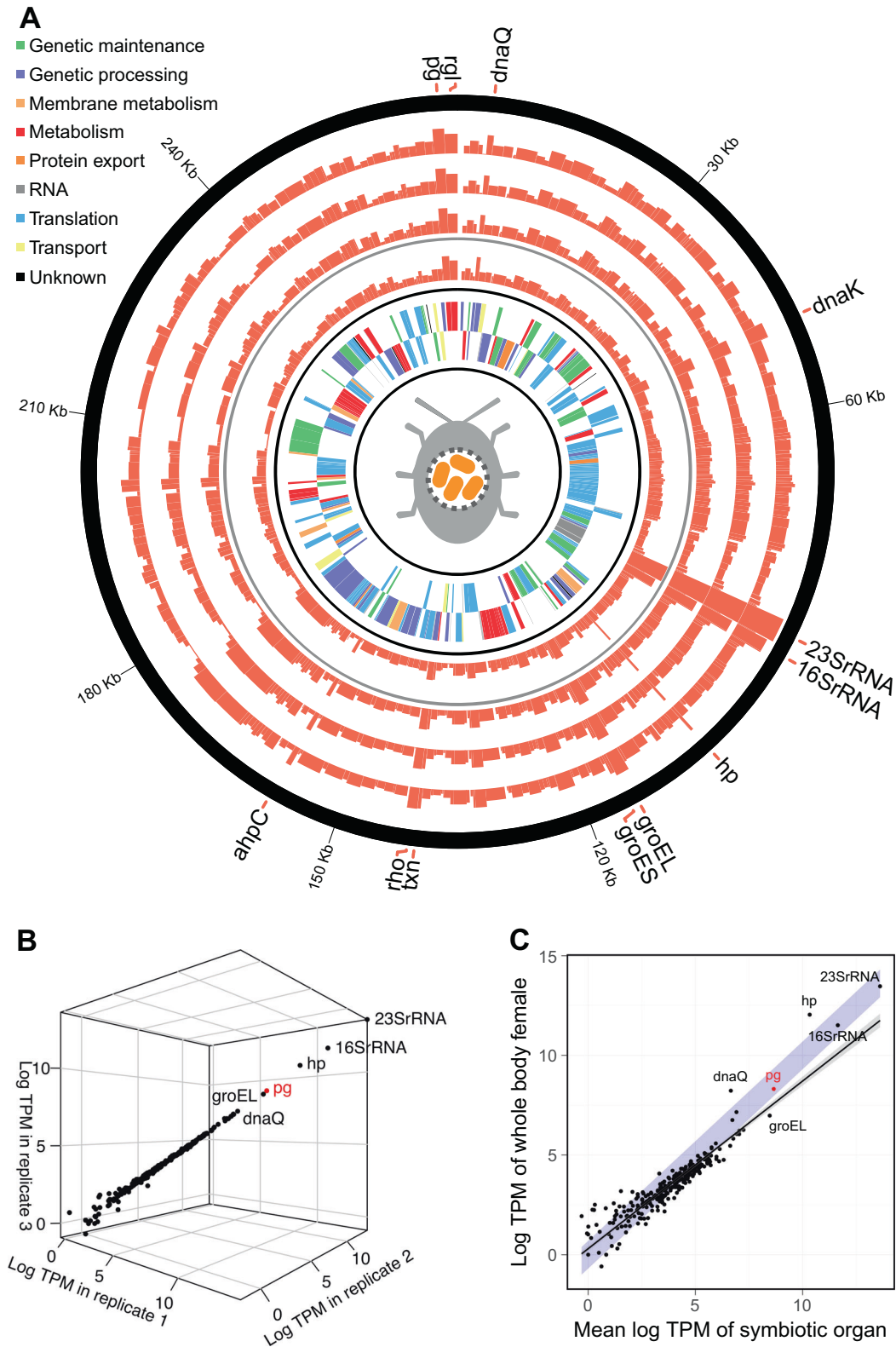
✉ Martin Kaltenpoth
mkaltenpoth@uni-mainz.de

✉ Hassan Salem
hassan.s.salem@gmail.com

¹ Department of Evolutionary Ecology, Johannes Gutenberg University, 55128 Mainz, Germany

² Department of Biology, Emory University, 30320 Atlanta, GA, USA

³ Present address: Mutualisms Research Group, Max Planck Institute for Developmental Biology, Tübingen 72076, Germany



[2]. This gammaproteobacterial microbe, ‘*Candidatus Stammera capleta*’ (henceforth *Stammera*), is vertically transmitted and located in specialized foregut-associated

symbiotic organs of the tortoise leaf beetle (Coleoptera: Chrysomelidae: *Cassida rubiginosa*), an obligate folivore [2]. Through the production and secretion of two pectinases,

◀ **Fig. 1** Expression profile in transcripts per million (TPM) of the annotated *Stammera capleta* genome. The inner two layers in (a) represent the identified genes in the forward and reverse direction, the next layer represents the *Stammera* gene expression profile in the whole body sample of a female *Csassida rubiginosa* (excluding the gut-associated symbiotic organ, but including the symbiont transmission organs associated with the reproductive tract), and the outer three layers represent the three replicates of *Stammera* gene expression profiles in the gut-associated symbiotic organs of two male and one female *C. rubiginosa*, respectively. **b** Correlation of the relative expression of each *Stammera* gene for the three replicates of symbiotic organs (mean pairwise Pearson correlation of gene expression profiles between replicate RNAseq data sets: $R^2 = 0.98$; $p < 0.001$). **c** Correlation of the mean relative *Stammera* gene expression in the gut-associated symbiotic organ replicates and the whole body female sample (Pearson correlation, $R^2 = 0.9$; $p < 0.001$). The line represents a linear regression and the blue area indicates a perfect correlation between both variables. Genes encoding for a hypothetical protein and the polygalacturonase are abbreviated as hp and pg, respectively between both variables. Genes are abbreviated as: hypothetical protein (hp), polygalacturonase (pg), rhamnogalacturonan lyase (rgl), alkyl hydroperoxide reductase C (ahpC), transcription termination factor Rho (rho), and thioredoxine (txn).

Stammera aids its host in the digestion of plant cell wall polysaccharides. Thus, in contrast to most obligate nutritional symbioses, the function for the host is reduced to two isolated enzymatic tasks, instead of pathways with multiple enzymes (e.g., amino acid biosynthesis) [3] (Table S1). In the present study, we investigated the metabolic potential of *Stammera* in the context of the host's as well as symbiont's transcriptomic profiles to better understand how a microbe can afford such a limited metabolism outside the nutritional comforts of a host cell.

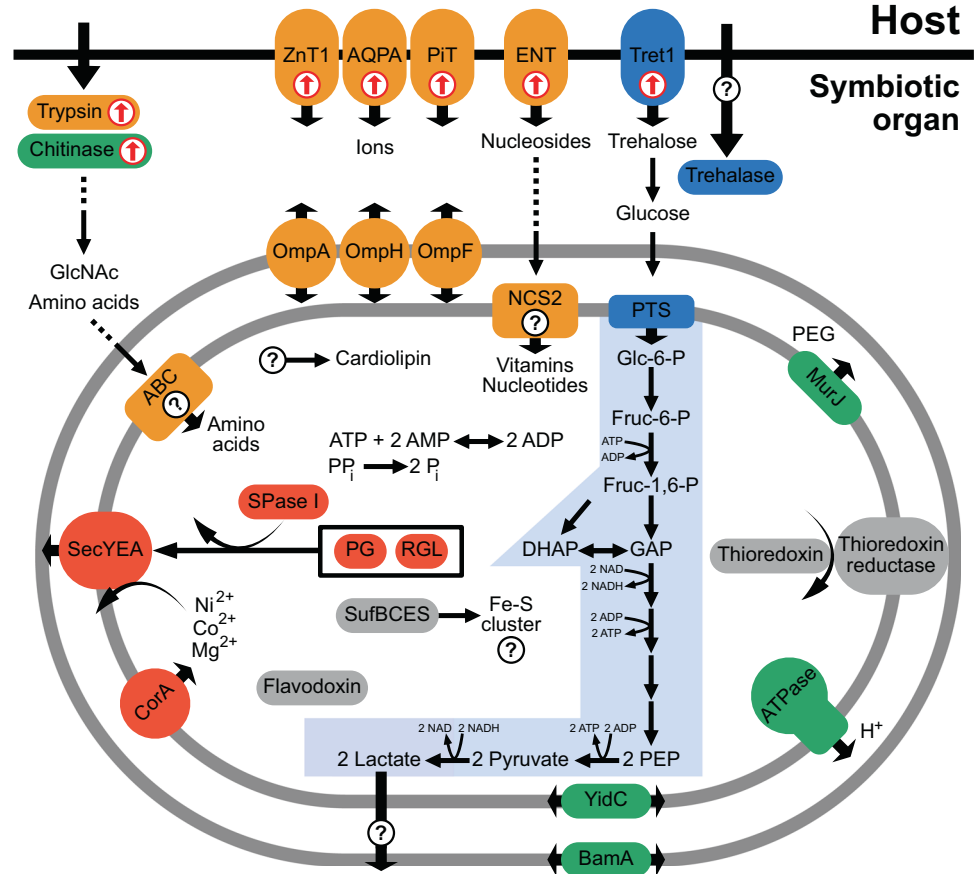
The highly consistent gene expression profile of *Stammera* (mean pairwise Pearson correlation of three replicates: $R^2 = 0.98$; $p < 0.001$) demonstrates a restricted control at the transcriptional level over the reduced genome (Fig. 1). Among the most highly expressed transcripts are chaperone-encoding genes (*groEL*, *groES*, *dnaK*) (Fig. 1; Table S2). Reflecting their capacity to ameliorate protein dysfunction, their preferential expression may indicate the need for protein maintenance and refolding. This is consistent with other symbionts possessing reduced genomes [9, 10], underlining the general challenges in maintaining stability of cellular functions in AT-enriched chromosomes as well as accumulated mutations [11]. Strikingly, the expression of polygalacturonase (pg), a pectinase, exceeded the expression of all other protein-coding genes (Fig. 1), reflecting the mutualistic role of *Stammera* towards its host. Rhamnogalacturonan lyase (rgl), a supplementary pectinase, was also preferentially expressed (Fig. 1A). The elevated expression patterns in *Stammera* are consistent with the transcriptional profiles reported for primary endosymbionts, where genes coding for chaperones and host-beneficial factors (e.g., essential amino acids) are among the most highly expressed [12].

In many insect symbioses, the host is able to control the symbiont population by leveraging its immune system. Among the differentially expressed transcripts in the symbiotic organ were immune signaling genes that were generally downregulated compared with whole body samples (Fig. S1). Interestingly, an isoform of a peptidoglycan recognition protein, as well as a galectin and cathepsin B were strongly upregulated in the symbiotic organ (Fig. S1). In weevils, antimicrobial peptides are known to regulate symbiont populations to avoid overgrowth [13]. Galectins, in particular, are discussed as relevant cell surface-binding proteins that could potentially control microbial colonization [14]. Immunoregulation thus represents an interesting avenue to understand how symbiont populations fluctuate in response to the metabolic demands of their hosts.

Stammera's degraded genome raises the question of how the functional deficiencies of this extracellular bacterium are compensated. Several host genes for managing internal insect resources, such as chitin remodeling and transport of nutrients, are upregulated in the symbiotic organ (Figs. S1 and S2). Given its connection to the foregut [2] the symbiotic organ is likely surrounded by chitin. The turn-over of foregut-lining chitin is a common process in insects and releases several nutrients such as N-acetylglucosamines [15] that serve as precursors for microbial membrane biosynthesis [16], which is eroded in *Stammera* (Fig. 2, Table S1). In addition, the host cells of the symbiotic organ exhibit an upregulated expression of trypsin (Fig. 2). Trypsin-like proteases degrade insect cuticular proteins [17], which could enrich *Stammera*'s environment with amino acids (Fig. 2), since *Stammera* cannot synthesize these nutrients (Fig. 2 and Table S1). The upregulation of nucleoside and ion transporters in the symbiotic organ (Fig. 2) may reflect additional supplementation of essential compounds to *Stammera*, as previously described in nutritional partnerships involving sap-feeding Hemiptera and their bacterial endosymbionts [18, 19]. The most highly upregulated transport protein in the symbiotic organ is a transporter for trehalose (Fig. 2), one of the most abundant sugars in insects [20], potentially supporting the energy metabolism of *Stammera* (Fig. 2). Taken together, we predict that digestive enzymes and transporters of the symbiotic organ could enrich *Stammera* with essential nutrients by remodeling the extracellular host environment.

Although often highly reduced, some intracellular symbionts appear to rely on retained genes in the TCA cycle for the production of amino acids and vitamins [1, 5], tying respiratory energy generation with the host-beneficial factors that underlie the host-symbiont partnership. For *Stammera*, the sole functional contribution to the host lies in the production of pectinases, relaxing selection for the symbiont to maintain vitamin and amino acid biosynthetic pathways and, correspondingly, the TCA cycle alongside the enzymes of the respiratory complex. Instead, the

Fig. 2 *Stammera capleta* reconstructed metabolism and nutritional supplementation by the host, as inferred from genomic and transcriptomic data, respectively. Carbohydrate metabolism is marked in blue, production and export of pectinase enzymes in red, cofactor metabolism in orange, membrane metabolism in green, and individual metabolic enzymes in gray. Cellular processes of which the functions are unknown, or no corresponding gene was found are highlighted with a question mark. Host genes that were found to be significantly upregulated in the symbiotic organ are highlighted with red arrows



reduction equivalents (e.g., NADH) generated during glycolysis, can be oxidized through a lactate dehydrogenase (LDH). This metabolism requires 18 times more glucose than aerobic respiration to generate the same amount of ATP [21], resulting in a high carbohydrate demand, which could be met by the host's carbohydrate-rich diet and the upregulation of a trehalose transporter (Fig. 2). Interestingly, a fermentative LDH is missing in related symbionts, but is present in free-living relatives (Table S3), indicating that a conditional fermentative and respiratory metabolism was restricted to sole respiration in most endosymbionts (Fig. 3). Lactate fermentation has been discussed in the context of parasitic eukaryotes [22], but is unknown in insect symbionts with reduced genomes, likely because lactate accumulation would result in toxicity within host cells [23]. In the case of *Stammera*, however, the extracellular localization supports the erosion to a sole fermentative metabolism by minimizing acid stress. Thus, *Stammera's* central metabolism is reduced considerably beyond that of some intracellular symbionts, maintaining glycolysis and an LDH. Further functional studies are needed to validate the hypothesis this study poses on the evolution of *Stammera's* metabolism (Fig. 3).

In conclusion, despite sharing many metabolic features with intracellular symbionts possessing drastically reduced

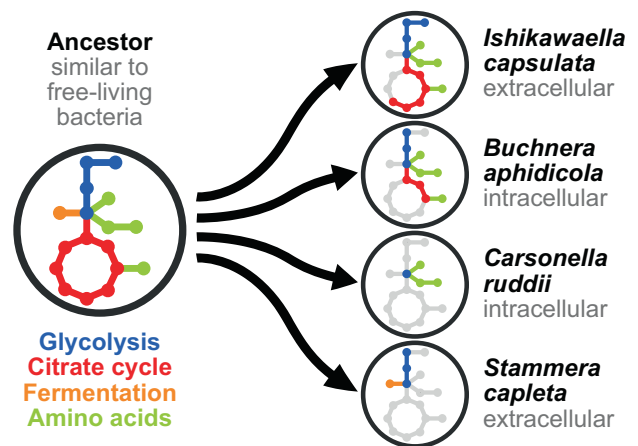


Fig. 3 Proposed reductive evolutionary scenarios leading to variable degrees of metabolic pathway loss in representative Gammaproteobacteria symbionts. The putative relatives are similar to free-living microbes, such as *Escherichia coli*, that possess the complete pathways for glycolysis, aerobic respiration, as well as mixed acid fermentation

genomes, *Stammera* relies on a fermentative metabolism for autonomous energy conversion, permitted by its extracellular location, host supplementation, and pectinolytic function.

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Author contributions EB, MK, and HS conceived the study. EB performed the genome and transcriptome annotation. EB, MK, and HS analyzed data. EB wrote the paper. All authors edited and commented on the paper.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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