Horizontal transmission enables flexible associations with locally adapted symbiont strains in deep-sea hydrothermal vent symbioses

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Symbiont specificity, both at the phylotype and strain level, can have profound consequences for host ecology and evolution. However, except for insights from a few model symbiosis systems, the degree of partner fidelity and the influence of host versus environmental factors on symbiont composition are still poorly understood. Nutritional symbioses between invertebrate animals and chemosynthetic bacteria at deep-sea hydrothermal vents are examples of relatively selective associations, where hosts affiliate only with particular, environmentally acquired phylotypes of gammaproteobacterial or campylobacterial symbionts. In hydrothermal vent snails of the sister genera Alviniconcha and Ifremeria, this phylotype specificity has been shown to play a role in habitat distribution and partitioning among different holobiont species. However, it is currently unknown if fidelity goes beyond species-level associations and influences genetic structuring, connectivity, and habitat adaptation of holobiont populations. We used metagenomic analyses to assess sequence variation in hosts and symbionts and identify correlations with geographic and environmental factors. Our analyses indicate that host populations are not differentiated across an ~800-km gradient, while symbiont populations are clearly structured between vent locations due to a combination of neutral and selective processes. Overall, these results suggest that host individuals flexibly associate with locally adapted strains of their specific symbiont phylotypes, which supports a long-standing but untested paradigm of the benefits of horizontal transmission. Symbiont strain flexibility in these snails likely enables host populations to exploit a range of habitat conditions, which might favor widespread genetic connectivity and ecological resilience unless physical dispersal barriers are present.

chelomysynthetic symbiosis | hydrothermal vents | host-symbiont population genomics | habitat adaptation | symbiont transmission and specificity

Mutualistic relationships between eukaryotes and bacterial microbes are ubiquitous in nature. Symbionts enable hosts to gain access to novel resources and habitats, provide protection against pathogens and predators, and can be essential for the hosts’ diets (1, 2). For symbiotic associations to persist over evolutionary time, hosts must successfully transmit their symbionts from one generation to the next, either through symbiont acquisition from the environment (horizontal transmission), direct inheritance of symbiont lineages (vertical transmission), or through the host germline, or a combination of both mechanisms (mixed transmission) (3, 4). The mode of transmission has significant implications for the composition and variation of symbionts within and between host individuals. Vertical transmission necessarily results in strong genetic coupling between host and symbiont lineages and an accompanied reduction in intrahost symbiont diversity (3). By contrast, horizontal transmission exposes asymbiotic hosts to a potentially heterogeneous environmental pool of symbiont lineages (3). This can promote the formation of generalist partnerships, where multiple hosts and symbionts associate with each other, to more specialized associations between only one or a few potential partners (1, 5). This range is often referred to as host–symbiont specificity, which can vary in its taxonomic level for both partners (i.e., partner fidelity) depending on the symbiotic system. The degree of partner fidelity and its effect on symbiont composition can have dramatic impacts on holobiont functioning (6). For example, host–symbiont specificity, both at the species and genotype level, is crucial for light production in bioluminescent squid (7), while shifts in symbiont communities have been shown to affect phytoplankton growth rates (8) and the efficiency of nitrogen fixation in nodule-forming legumes (9–11).

Environmental transmission of obligate symbionts is particularly common in marine ecosystems (2). Apart from host–symbiont genetic interactions, isolation by distance through geographically limited dispersal, selection by environment, or interactive

Significance

In marine ecosystems, transmission of microbial symbionts between host generations occurs predominantly through the environment. Yet, it remains largely unknown how host genetics, symbiont competition, environmental conditions, and geography shape the composition of symbionts acquired by individual hosts. To address this question, we applied population genomic approaches to four species of deep-sea hydrothermal vent snails that live in association with chemosynthetic bacteria. Our analyses show that environment is more important to strain-level symbiont composition than host genetics and that symbiont strains show genetic variation indicative of adaptation to the distinct geochemical conditions at each vent site. This corroborates a long-standing hypothesis that hydrothermal vent invertebrates affiliate with locally adapted symbiont strains to cope with the variable conditions characterizing their habitats.

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effects of these processes can shape the composition, diversity, and structure of horizontally transmitted symbionts across host populations (12). However, despite their importance for holobiont biology, the relative contributions of these factors to symbiont composition within and among hosts remain understudied in most environmentally acquired marine symbioses. It has long been hypothesized that horizontal transmission enables host organisms to associate with locally adapted symbiont strains (i.e., subtypes within a bacterial species), conferring fitness advantages in spatially and temporally variable marine habitats (3, 13–15). Especially for long-dispersing aposymbiotic larvae that are likely to encounter new habitat conditions when they settle, association with a locally adapted symbiont strain may be advantageous compared to carrying a vertically transmitted symbiont that might be maladapted at a nonnative site. This hypothesis has been indirectly supported by evidence that marine animals with horizontally transmitted obligate microbial symbionts often host location-specific strains (15–20). However, the influence of local adaptation relative to neutral evolutionary processes on symbiont geographic structure and genomic traits has not been formally evaluated.

Chemosynthetic animal–microbe symbioses are globally significant phenomena that dominate hydrothermal vent and hydrocarbon seep ecosystems in the deep sea. In these associations, the bacterial partner uses chemical energy from the oxidation of reduced compounds, such as hydrogen, sulfide, or methane, to synthesize organic matter, which serves as primary nutrition for the host (21). Vent animals harboring chemosynthetic symbionts are typically very specific in their partnerships: In the predominant number of cases, host individuals associate with only one or two phylotypes of gammaproteobacterial or campylobacterial symbionts (i.e., species- or genera-level phylogenetic clades based on 3 and 5% 16S ribosomal RNA [rRNA] sequence divergence, respectively) (21), whereas symbionts can exhibit a comparatively broad host range. Chemosynthetic endosymbioses in deep-sea snails of the Indo-Pacific sister genera *Alviniconcha* and *Ifremeria* are examples of reciprocally relatively specific partnerships, where host species harbor only particular symbiont species or genera across their geographic distribution. Given the absence of host–symbiont phylogenetic concordance, the symbionts are assumed to be environmentally acquired in both genera (22) despite contrasts in reproductive mode. Although reproductive strategy does not always predict symbiont transmission mode (23), a vertical transmission component is more likely in *Ifremeria*, as this genus is known to brood its larvae (24), giving symbionts more developmental time to transfer.

In the Eastern Lau Spreading Center (ELSC), previous work suggested that specificity to functionally distinct symbiont phylotypes drives local- and regional-scale habitat partitioning among four co-occurring *Alviniconcha* and *Ifremeria* species (25–28). *Alviniconcha boucheti* from the ELSC contains a campylobacterial phylotype (Epsilon) and is usually found at northern vent sites with high concentrations of sulfide and hydrogen, while *Alviniconcha kojima* and *Alviniconcha strummeri* associate with different gammaproteobacterial phylotypes (Gamma1 and Gamma1/GammaLau, respectively) and usually occupy midlatitude to southern vent sites, where the concentrations of these chemical reductants are lower (25, 27–29). *Ifremeria nautili* establishes dual symbioses with thiophoric and methanotrophic gammaproteobacterial endosymbionts (29) and is codistributed with *Alviniconcha* across their geographical range, although it typically segregates into habitat patches with reduced fluid flow relative to its sister genus (30). While it has been well-established that niche differentiation across hydrothermal vents is likely linked to symbiont phylotype specificity in *Alviniconcha* and *Ifremeria* host species, nothing is known about the fidelity of these associations at the population level, how strain-level specificity might influence host population structure and connectivity, and how regional adaptation is conferred functionally.

In this study, we applied population genomic methods to assess symbiont strain-level genetic variation and patterns of host–symbiont genetic subdivision in *Alviniconcha* and *Ifremeria* species from the ELSC and Tonga Volcanic Arc (Fig. 1A and Dataset S1). We first assembled transcriptomes and pangenomes for each host species and symbiont phylotype, respectively, and then determined sequence and functional variation between populations by mapping reads from multiple individuals against the reference assemblies. Using multivariate statistical analyses, we subsequently evaluated the impact of host genetics, environment, and geography on symbiont composition in populations of all four snail species and assessed the effect of local adaptation on symbiont geographic structure. To test if vertical transmission has linked symbiont and host genealogies or if horizontal transmission has randomized these relationships, we further assessed the topological concordance among symbiont and host mitochondrial phylogenies.

**Results**

**Host Populations Are Not Structured across Geographic Locations.** Host transcriptomes were coassembled from 11 to 25 individual gill samples per species and used as references for variant calling to assess host population genomic structure. Transcriptome assemblies consisted of 24,176 to 35,654 transcripts (totaling 20.68 to 28.68 Mb) and were ∼30.30 to 55.40% complete (Dataset S2). Mapping of 1,138,186 to 541,418,710 (mean 20,984,011) host reads against the transcriptome references (Dataset S3) and subsequent filtering of variant sites yielded 1,655 to 9,185 single-nucleotide polymorphisms (SNPs) per species for population genetic analyses. Irrespective of host taxon, *F*~ST~ and ordination analyses revealed low genetic differentiation among host populations sampled from different vent localities (Fig. 1B and Dataset S4; *F*~ST~ = 0.0400 to 0.1579). With the exception of four SNP sites in *A. kojima* (24), no *F*~ST~ outliers that could be indicative of adaptive variation were detected in any host species at a *q* value ≤ 0.05 (Dataset S5). The effect of geography on host population genetic structuring was not significant in Mantel correlations (Dataset S6).

**Snail Symbionts Belong to Six Bacterial Species That Are Frequently Horizontally Transmitted.** Pangenomes of each symbiont phylotype were reconstructed from 4 to 59 high-quality symbiont metagenome assembled genomes for population genomic and phylogenomic analyses (Datasets S7 and S8). Based on average nucleotide identities and taxonomic assessments, symbiont phylotypes recovered in our dataset comprised six species from five genera of chemosynthetic bacteria: *Sulfurimonas* “Epsilon” (*A. boucheti* from the ELSC), a second *Sulfurimonas* species (*A. boucheti* from Niua South), unclassified Gammaproteobacteria “Gamma1” (*A. kojima* and *A. strummeri*), *Thiolapillus* “Ifr-SOX”, and *Methylomonadaceae* “Ifr-MOX” (L. *nautili*) (Dataset S7). Because of the low coverage of the *GamaLau* and *Ifr-MOX* species that resulted in poor variant recovery (SI Appendix, Fig. S1 and Dataset S7), these symbionts were excluded from further analyses. In addition, the *Sulfurimonas*
species from Niua South was not included in population genomic analyses due to taxonomic divergence from the Epsilon symbiont associated with *A. boucheti* from the ELSC. The dominant symbiont species within the ELSC (Epsilon, Gamma1, and Ifr-SOX) were considered for all downstream analyses.

To assess if symbionts exhibit evidence of vertical transmission, we determined the congruence between tree topologies of the host mitochondrial and symbiont core genomes. For all host species, symbiont and mitochondrial phylogenies were clearly discordant (Fig. 1C). This indicates that symbionts are primarily environmentally acquired or that enough horizontal transmission has occurred to erode the signal of vertical transmission in both *Alviniconcha* and *Ifremeria* despite differences in reproductive strategy.

**Fig. 1.** (A) Sampling map for *Alviniconcha* and *Ifremeria* species at vent fields along the ELSC (Kilo Moana, Tow Cam, Tahi Moana, ABE, and Tu’i Malila) and the Tonga Volcanic Arc (Niua South). Vent field locations are shown as colored diamonds and the symbols below each site label indicate the species sampled from the respective vent field. Map data ©2021 Google. (B) Principal-component plots for *Alviniconcha* and *Ifremeria* host species based on genetic covariance matrices. Axes show percent of variation explained by the first and second principal components. Each symbol represents a single snail sample colored by host species, symbiont and mitochondrial phylogenies were considered for all downstream analyses.

Symbiont Populations Are Segregated by Geography, with Little Effect of Host Genetics. Populations of the dominant ELSC symbiont species, Epsilon, Gamma1 and Ifr-SOX, were structured by geography based on 1,716 to 7,741 variants (Fig. 2, SI Appendix, Fig. S2, and Dataset S9). Genetic variation between symbiont populations typically increased with geographic distance between vent fields and could be broadly partitioned into four regions: Tu’i Malila, ABE + Tahi Moana, Tow Cam + Kilo Moana, and Niua South (Dataset S9; $F_{ST} = 0.1012$ to 0.8682). In general, differentiation within the ELSC was larger for the *Alviniconcha* symbionts than for the *Ifremeria* symbionts across the same geographic scale. Genotype–site associations were significant even when corrected for host genetic variation (Dataset S6; $P \leq 0.0285$, $r = 0.2861$ to 0.74).
0.8512), with the strongest geographic effect being observed in the Epsilon and Ifr-SOX symbionts of *A. boucheti* and *I. nautilei*, respectively.

The Gamma1 symbiont was associated with both *A. kojimai* and *A. strummeri* and we therefore investigated whether populations of this phylotype varied between host species. Ordination analyses and Mantel tests indicated a clear affiliation to host taxon that superseded the effect of geography by about a factor of 3, suggesting that different Gamma1 strains associate with host taxon that superseded the effect of geography by about a factor of 3, suggesting that different Gamma1 strains associate with host species, while 3.27% were correlated with other environmental factors (Fig. 2 and Dataset S10; $P = 0.001$, $r^2_{adj} = 38.54%$). Likewise, the overall marginal effects of each explanatory variable that was included in the RDA models for the different host–symbiont pairs were significant, except for year in the case of the Gamma1 symbiont (Dataset S11).

Candidate adaptive loci in each symbiont species encompassed a wide range of functional categories, from metabolism of macronutrients, cofactors, and cell wall/membrane to energy production, maintenance of cellular homeostasis, motility, mobile elements, and response to environmental stressors and phages (Dataset S10). In concordance with the known environmental differences among vent sites in the ELSC and previous observations of habitat partitioning between *Alviniconcha* holobionts linked to vent geochemistry (25), a subset of adaptive variants in the Epsilon and Gamma1 symbionts was located in
genes related to sulfur and hydrogen metabolism (e.g., sulfide-quinone oxidoreductase, sulfate permeases, carbamoyltransferase HypF). Conversely, variants associated with these functions appeared to not be under positive selection in the Ifr-SOX symbiont. Notably, this symbiont encoded loci involved in conjugation that showed evidence for adaptive evolution and that were not detected as outliers in the Alviniconcha symbionts. Interestingly, all symbionts contained variants in genes related to vitamin B12 metabolism that were linked to local adaptation. In the Gamma1 symbiont, adaptive loci related to host species comprised several genes involved in immunity (e.g., production of cell-wall/membrane compounds, virulence, response to oxidative stress), nutrient provisioning (carbon fixation, cofactor and macronutrient metabolism), and cellular respiration. In general, candidate variants in all symbiont species were characterized by elevated \( F_{ST} \) values and were located in genes with increased or reduced \( pN/pS \) ratios (Dataset S12), which further supports their potential role in habitat adaptation and highlights the essentiality of the respective genes for symbiont functioning.

**Symbiont Populations Differ in Gene Content.** We used the symbiont pan-genome references to further assess variation in gene content between symbiont populations from different vent localities or broader geographic regions based on groupings observed in the ordination and \( F_{ST} \) analyses (Fig. 3, SI Appendix, Fig. S4, and Dataset S13). For the Gamma1 symbiont, we additionally determined gene content variation between populations from different host species, given that this symbiont species occurs in both *A. strummeri* and *A. kojimai* (SI Appendix, Fig. S3 and Dataset S13). Differentially preserved genes between symbiont populations comprised about 1.33 to 3.84% of protein-coding regions.

Independent of symbiont species, the most common differences among geographic or host-specific populations concerned genes of unknown or miscellaneous function as well as a smaller number of genes related to mobilome (e.g., transposases, integrases), antiviral defense (e.g., restriction–modification systems, CRISPR-Cas systems), and DNA metabolism (e.g., DNA polymerases, helicases, repair systems) (Fig. 3, SI Appendix, Figs. S3 and S4, and Dataset S13). Populations of the *A. bouchei* Epsilon symbiont from Kilo Moana/Tow Cam and ABE further differed in several cell wall/membrane-related genes, a protein phosphatase, a 3-dehydro-bile acid delta(4, 6)-reductase, and an inosine-5‘-monophosphate dehydrogenase (Fig. 3, SI Appendix, Fig. S4, and Dataset S13). Within the Gamma1 symbiont of *A. kojimai*, populations from Tu’i Malila contained multiple private genes that were related to cell signaling, iron–sulfur cluster biosynthesis (*sofS*), iron uptake, and other membrane transport processes, whereas populations from ABE encoded genes for succinate dehydrogenase assembly and hydrogenase maturation. Differences in hydrogenase-related genes were also evident in the Gamma1 symbiont of *A. strummeri*, where genes for uptake and sensory hydrogenases (*hoxKL, hydL*) as well as hydrogenase maturation and regulation (*hypB, hypDEF, hupR1, hupU*) were largely missing in populations from Tu’i Malila (Fig. 3, SI Appendix, Fig. S4, and Dataset S13). In addition, these populations lacked several transcription factors and RNA helicases that were present in lineages from Tahi Moana, while containing private genes involved in cell signaling, membrane transport, as well as carbon, vitamin B12, and secondary metabolism. Similar contrasts in metabolic functions were observed between host-specific strains of the Gamma1 symbiont. Most notably, host-specific populations differed in the presence of genes for a sulfite dehydrogenase complex (*soeABC*), which was present in strains from *A. kojimai* but not *A. strummeri* (SI Appendix, Fig. S3 and Dataset S13). The *I. nautilis* Ifr-SOX populations from Niua South and the ELSC showed contrasting conservation patterns in genes that spanned a variety of metabolic and cellular functions, including asparagine biosynthesis, macro-nutrient, cofactor, nitrogen, nucleotide, and RNA metabolism, transport of iron and other inorganic compounds, cell regulation, conjugation, and cellular respiration (Fig. 3, SI Appendix, Fig. S4, and Dataset S13).

**Discussion**

Strain-level variation within microbial symbionts is increasingly recognized as an important driver of host ecology and evolution (6), yet its patterns, determining factors, and functional implications remain poorly investigated in many nonmodel symbioses. In this study, we used metagenomic analyses to assess sequence and gene content differences in chemosynthetic symbionts associated with four co-occurring species of deep-sea hydrothermal vent snails and determined the impact of host genetic, geographical, and environmental factors on symbiont strain composition and variation.

Despite fidelity between hosts and symbionts at the host species level (25, 27), our results indicate that specificity between host genotypes and symbiont strains in *Alviniconcha* and *Ifremeria* is weak: Host populations were not partitioned across an ~800-km gradient, whereas symbiont populations were clearly structured between vent locations or broader geographic regions. Correlations between symbiont and host genetic distances were either insignificant or low, suggesting that host genetics has a minor impact compared to environment or geography on strain composition within both *Alviniconcha* and *Ifremeria*.

These findings qualitatively agree with observations in deep-sea mussel hybrids that appear to associate with locally available symbiont strains (19), but contrast markedly with patterns in other horizontally transmitted partnerships, such as the squid–*Vibrio* symbiosis and some legume–rhizobia associations, where hosts exhibit strong strain specificity (6). Environmental uptake of locally adapted symbiont strains provides host organisms with the opportunity to optimally exploit novel habitats, but carries the risk of unsuccessful symbiont acquisition and infection by cheaters (3). Depending on the amount of partner reliance, chances of symbiont encounter, and fitness variation among symbiont strains, holobionts likely find different trade-offs between these opposing factors. Associations with chemosynthetic bacteria are obligate for hydrothermal vent animals, but are restricted to relatively ephemeral habitats that are characterized by large temporal and spatial fluctuations in environmental conditions and associated shifts in microbial communities (31–33). Strong nutritional dependency in vent symbioses combined with the uncertainty of habitat (and thus symbiont) encounter might promote specificity toward a mutualistic symbiont phylotype, while enabling enough flexibility toward different local strains of that phylotype to maximize recruitment success of host larvae at a new vent site. Although vent tubeworms, for example, are known to release symbionts into their environment upon death (34), it is not guaranteed that larvae self-recruit and take up the same symbionts as their population of origin, given the transient nature of hydrothermal vents. By contrast, juvenile bobtail squids affiliate with symbiotic bacteria that are expelled by adult hosts into the native environment (35), which might favor increased strain selectivity in this association. Similarly, some legume species
appear to preferentially associate with certain rhizobial strains that are abundant in the native host range (36). Although the factors underlying strain specificity are not fully understood, it is possible that the relative fitness advantages provided by these locally available strains (36) coupled with the benefit of decreased cheater invasion (11) might have selected for strong partner fidelity in these symbioses.

\textit{A. kojimai} and \textit{A. strummeri} represent notable exceptions to the observed patterns of symbiont specificity, as they share one species of gammaproteobacterial symbionts (Gamma1), but appear to take up functionally distinct, site-specific strains even in habitats where these host species co-occur. Strains of both host species showed evidence for positive selection in genes encoding outer-membrane proteins and factors for the biosynthesis of cell-wall lipopolysaccharides, which likely play a role in host–symbiont recognition and specificity (37).

The phylogenetic divergence between \textit{A. kojimai} and \textit{A. strummeri} (~25 million years ago; MYA) is more recent than that between either species and \textit{A. boucheti} (~38 MYA) or \textit{I. nautili} (~113 MYA) (27). Perhaps the closer evolutionary relationship between these two species favors associations with the same symbiont phylotype but distinct symbiont strains. A similar scenario has been suggested for \textit{Bathymodiolus azoricus} and \textit{Bathymodiolus puteonserpentis} on the Mid-Atlantic Ridge (~8.4 MYA) (38) as well as \textit{Bathymodiolus thermophilus} and \textit{Bathymodiolus antarcticus} on the East Pacific Rise (~2.5 to 5.3 MYA) (39), which do not show phylotype specificity in contrast to their more divergent relatives from Atlantic cold seeps (~15.4 MYA) (19, 38). However, compared with the bathymodiolin mussel system, the split between \textit{A. kojimai} and \textit{A. strummeri} is significantly older, which could indicate that the timing of phylotype specificity evolution varies among taxonomic groups, possibly as a result of contrasting ecological or evolutionary contexts. The fact that \textit{A. kojimai} and \textit{A. strummeri} nevertheless associate with distinct symbiont strains in sympatry is potentially a mechanism to avoid competition for niche space. Gamma1 strains of these two species notably differed in the presence of a molybdenum-containing sulfite dehydrogenase (SoeABC), which was preserved in strains of \textit{A. kojimai} but not \textit{A. strummeri}. SoeABC is the predominant enzyme for sulfite oxidation in many purple sulfur bacteria (40) and is further involved in taurine and dimethylsulfoniopropionate degradation in \textit{Roseobacter} clade bacteria (41–43). Although the physiological role of SoeABC in \textit{A. kojimai}’s Gamma1 strain

![Fig. 3. Dotplot showing differences in gene content for each functional category between habitats and symbiont species. Dots are scaled by the number of genes in each category and colored by geographic region. ELSC includes all vent sites in the Lau Basin sampled for \textit{Ifremeria} (Tow Cam, Tahi Moana, ABE, Tu’u Malila). Presence/absence of genes was determined in PanPhAna based on gene coverage profiles.](https://www.pnas.org/doi/10.1073/pnas.2115608119)
is unknown, it is possible that it contributes to partitioning of sulfur resources among co-occurring snail holobionts.

Our data support a model of horizontal symbiont transmission in both *Alviniconcha* and *Ifremeria*. While these results are expected for *Alviniconcha* that produce free-swimming planktrophic larvae and do not invest in their young (44), they are rather surprising for *Ifremeria* that brood their offspring in a modified pouch in the female’s foot (24) and would thus have more opportunities to vertically transmit their symbionts (independent of possible transovarial inheritance). Although topological discordance between host mitochondrial and symbiont phylogenomic trees suggests frequent horizontal transmission, it is possible that *Ifremeria* symbionts exhibit a mixed transmission mode and maternally acquired symbionts get replaced or complemented by more competitive strains in the habitat where the snail larvae settle. Our current dataset cannot distinguish this possibility from a strict horizontal transmission mode, though future studies assessing symbiont composition in different developmental stages of *Ifremeria* would be helpful to address this hypothesis.

While the majority of genetic variants in the symbionts did not deviate from neutral expectations, a subset of loci showed evidence for natural selection, implying a role of both genetic drift and local adaptation in shaping symbiont population structure. Candidate adaptive loci spanned a surprisingly broad range of metabolic functions, many of which were probably not causally linked to the investigated environmental predictors but other correlated variables that we could not account for in this study. For example, in all investigated symbiont species, we observed adaptive variation in genes that were related to antiviral defense (e.g., CRISPR-associated proteins, restriction-modification systems) or mobile elements. Although differences in depth or geochemistry might contribute to these patterns, it is more likely that they reflect exposure of symbionts strains to distinct viral assemblages that might covary with local habitat conditions, as has been suggested for vestimentiferan tubeworm symbionts (45). Similarly, all symbionts showed evidence for local adaptation associated with vitamin B12 metabolism. This could be due to habitat-specific differences in the availability of cobalt, which is an essential trace element for the formation of vitamin B12. Though cobalt concentrations are unknown for hydrothermal fluids in the Lau Basin, they are typically correlated with those of iron and manganese and might thus show similar variation between vent fields, given the solubility dependence of all of these metals on temperature and chlorinity (46).

Polymorphisms in other genes, by contrast, are likely directly explained by variation in the analyzed environmental factors. Geographic strains of *A. bouchei*’s Epsilon symbiont, for instance, contained several variants under positive selection that were located in genes involved in sulfur oxidation and respiration. Likewise, the Gamma1 strains of *A. kojimai* and *A. strummeri* showed adaptive differences in variants related to hydrogen and sulfur metabolism. Niche-specific differences in hydrogen metabolic genes were also observed in comparisons of gene content among symbiont strains. In both *A. kojimai* and *A. strummeri*, Gamma1 strains from *Tu‘i Malila* lacked subunits of uptake and hydrogen-sensing hydrogenases as well as various genes for hydrogenase maturation and regulation. Given that H2 concentrations at *Tu‘i Malila* can drop to 35 μM in endmember fluids and are probably lower in diffuse-flow habitats (25), it is possible that hydrogen does not constitute a major energy source for Gamma1 strains from this locality. This hypothesis is in agreement with previous physiological experiments that revealed strikingly low hydrogen oxidation and associated carbon fixation rates in Gamma1 symbionts from *Tu‘i Malila* (28). An alternative though mutually nonexclusive explanation for the loss of hydrogenase-related genes at least in the *A. strummeri* strain from *Tu‘i Malila* could be avoidance of intrahost competition with co-occurring GammaLau strains, which sometimes codominate in *A. strummeri* individuals at this vent site (25). Such functional diversity is predicted to enable symbiont coexistence in a variety of hydrothermal vent symbioses, including bathymodiolin mussels (47, 48), alvinocarid shrimp (49), and vestimentiferan tubeworms (50). In contrast to the *Alviniconcha* symbionts, the *Ifremeria* Ifr-SOX symbiont did not show evidence for adaptive variation related to hydrogen and sulfur metabolism. This finding is not surprising, given that *Ifremeria* holobionts typically inhabit patches with less variable sulfide concentrations than *Alviniconcha* (51) and do not show the same patterns of regional-scale niche partitioning. Interestingly, we observed a higher number of adaptive variants linked to conjugal transfer in this symbiont. Perhaps site-specific selection in the respective genes promotes increased exchange of genetic material in the Ifr-SOX symbiont, which could improve adaptation to diverse environmental conditions and might thus enable this symbiont to occupy a broader range of ecological niches.

Although geography and environment appeared to be the dominant drivers of symbiont strain composition, some differences were linked to year of collection. In particular, strain pools of the Ifr-SOX symbiont from *Tu‘i Malila* showed significant temporal variation. Given that the Kilo Moana vent field expired between 2009 and 2016, it is likely that the ELSC experienced changes in hydrothermal activity and associated geochemical conditions during this time period, which could have induced shifts in biological communities. If individuals only acquire their symbionts during a short developmental window, the observed strain pools might reflect distinct host cohorts, or, alternatively, it is possible that long-lived host individuals replace their symbionts over time, resulting in compositional differences between years. Under both scenarios, however, potential symbiont subdivision within vent fields might have exacerbated temporal variation in intraspecific strain composition, given that different snail patches (41 to 87 m apart) were sampled between years.

Compared with their symbionts, host populations were markedly less structured across the same spatial scales. Although a proportion of genetic markers was differentiated across vent sites (0.41 to 12.70% with FST > 0.15), we did not find strong evidence for local adaptation in any host species, suggesting that these patterns likely reflect random variation among localities. The limited genetic subdivision between host populations agrees with predictions from biophysical models that indicate absence of physical dispersal barriers for vent larvae within the Lau Back-Arc Basin (52). While symbiont population structure, by contrast, appears to be at least partly driven by natural selection, it is possible that symbionts also experience stronger dispersal limitations than their hosts, as has been hypothesized in some coral–algae symbioses (18). The environmental distributions and life cycles of the free-living stages of *Alviniconcha* and *Ifremeria* symbionts are currently unknown, and a better understanding of these aspects will be necessary to evaluate the relative importance of dispersal barriers in symbiont biogeography.

Overall, our findings reveal that a lack of strain-level specificity in *Alviniconcha* and *Ifremeria* symbioses is common and possibly reflects an evolutionary strategy to cope with the transient and dynamic nature of hydrothermal vent habitats. However, a deviation from this pattern appears to occur in relatively closely related sympatric host taxa, which can evolve strong fidelity to local strains of the same symbiont species. In general,
strain flexibility might contribute to the widespread genetic connectivity observed among host populations, since the ability to partner with local strains should facilitate success across broad geographic and environmental scales and decrease ecological barriers to gene flow. In turn, this could favor resilience to natural but also anthropogenic environmental disturbances, a relevant consideration given the increasing human pressures on hydrothermal ecosystems worldwide (53). Our observations further support the fundamental hypothesis that horizontal transmission enables host organisms to associate with locally adapted symbiont strains (3, 12–14). Though the genomic basis of local adaptation can be detected in natural populations using population genomics methods, as we did here, evaluation of the phenotypic consequences of the observed strain-level genomic trait variation will be necessary to confirm local adaptation in these symbiont strains. Future work using organism-based manipulative experiments will be helpful to compare the fitness of hydrothermal vent animals hosting site-specific symbiont strains when exposed to native and foreign conditions (54). These assessments will be critical to understand the commonality of horizontal symbiont transmission in the marine environment (2), given the hypothesis that local adaptation is less common in marine than terrestrial systems due to higher levels of gene flow (55).

Materials and Methods

Sample Collection, Nucleic Acid Extraction, and Sequencing. Samples of Alviniconcha and Ithymera were collected from six vent sites (1,164 to 2,722 m) of the Lau Basin and Tonga Volcanic Arc in 2009 and 2016 using remotely operated vehicles (Fig. 1 and Dataset S1). Upon recovery, animal samples were dissected, placed in RNALater Stabilization Solution (Thermo Fisher Scientific), and frozen at −80 °C until further analysis. DNA was extracted with the Quick-DNA 96 Plus Extraction Kit (Zymo Research) and further purified with the MO BIO PowerClean DNA Pro Clean-Up Kit (Qiagen).

Host Transcriptome Sequencing and Assembly. Illumina RNA sequencing reads for host transcriptome assemblies were obtained from sequencing experiments performed in refs. 28 and 56. Adapter clipping, quality trimming, and removal of RNA and symbiont reads were performed as in ref. 28. Cleaned reads were error-corrected with RECOVERER (57) and filtered for uncorrectable and overrepresented sequences with the TRIMMOMATIC ASSEMBLYTOOLS package (https://github.com/harvardinformatics/TranscriptomeAssemblyTools). Host transcriptome coassemblies for each Alviniconcha and Ithymera species were performed with TRINITY (58) with the PasaFilt algorithm using 12,487,748 to 559,864,144 total reads per species. For each Alviniconcha species, additional transcripts were assembled from 6,434 to 46,731 454 reads obtained from ref. S9. Assembled contigs for each species were clustered with CD-HIT-EST (60) at a 95% identity threshold to reduce transcript redundancies. Open reading frames (ORFs) were predicted with TRANSDECODER (https://github.com/TransDecoder/TransDecoder) using a gene-wise best-hit criteria. Transcripts that did not contain any ORF or had a non-canonical translation initiation site were excluded. We further filtered reads based on strand bias (sb_pval < 0.05), heterozygote bias (hetbias_pval = 0.05), and probability of being variable (SNP_pval = 1e-6). In addition, we removed spurious and improperly paired reads, adjusted mapping qualities for excessive mismatches (C = 50), and computed per-base alignment qualities (BAQ = 1) to disregard variants close to indel regions. Putative paralogous variants were excluded by discarding reads with multiple mappings and by considering only sites that had a maximum depth of 40 to 80 (based on distributions of average coverage). Genetic distances between individuals were inferred by calculating pairwise covariance matrices.

Host population genomic variation was assessed in ANGSD (85) by inferring genotype likelihoods based on Hardy-Weinberg equilibrium considering individual inbreeding coefficients. To increase accuracy of the analyses, variant sites with mapping qualities <30 (minMapQ = 30), base qualities <20 (minQ = 20), and minimum minor-allele frequencies <0.01 (minMaf = 0.01) were excluded. We further filtered sites based on strand bias (sb_pval = 0.05), heterozygote bias (hetbias_pval = 0.05), and probability of being variable (SNP_pval = 1e-6). In addition, we removed spurious and improperly paired reads, adjusted mapping qualities for excessive mismatches (C = 50), and computed per-base alignment qualities (BAQ = 1) to disregard variants close to indel regions. Putative paralogous variants were excluded by discarding reads with multiple mappings and by considering only sites that had a maximum depth of 40 to 80 (based on distributions of average coverage). Genetic distances between individuals were inferred by calculating pairwise covariance matrices.

Symbiont Metagenomic Sequencing and Pangenome Generation. One hundred and ninety-two barcoded high-throughput DNA-sequencing libraries were prepared with a TN5 transposase-based protocol after ref. 63 at the University of California, Santa Cruz and then sent for 150-bp paired-end sequencing on a NovaSeq 6000 instrument at the University of California, Davis. However, due to low read allocation, 66 of the libraries were excluded from further analysis. Raw reads were trimmed with TRIMROMATIC (64) and filtered for sequence contaminants through mapping against the human (GRCh38) and Phix reference genomes. An average of 31,747,773 (2,015,718 to 882,674,692) filtered Illumina reads for each host individual were then assembled with minSPades (65), using k-mers from 21 to 121 in 10-step increments, and independently binned with MetaBAT 2 (66), MaxBin 2 (67), and GneissBin (68). Binning results were evaluated with DdxTox (69). Taxonomic identity of the top-scoring symbiont metagenome-assembled genomes (MAGs) was assessed with GTDB-Tk (70) and assembly quality was checked with CheckM (71) based on Gammaproteobacteria- or Campylobacteria-specific marker genes. High- to moderate-quality MAGs with >85% completeness and <10% contamination were chosen for further analysis (Dataset S4). Additional reassamplings of previously published draft symbiont genomes (29) were constructed from a combination of Illumina and Nanopore reads (SI Appendix, Methods). Average nucleotide identities between MAGs were calculated with fastANI (72) to determine species boundaries at the 95% threshold (73). All MAGs were annotated with Proka (74) and combined into species-specific symbiont pan-genomes with Panaroo (75), using default parameters for gene clustering and a gene-refining step to locate genes that were missed by gene prediction software within a radius of 1,000 nt. Genes without functional annotation were characterized through comparisons against the UniRef90 database with BlastP (76) and the Shimodaira-Hasegawa-like approximate-likelihood ratio test with 5,000 replicates by resampling both partitions and then sites within resampled partitions. To reduce the impact of model violations, each bootstrap tree was optimized using a hill-climbing nearest-neighbor interchange search (option --bnni). The best maximum-likelihood trees were midpoint-rooted with Pagcon (81) and cladograms were plotted with GGTREE (82) in R.

Phylogenomic Analyses of Host Mitochondria and Symbionts. To evaluate the presence of vertical transmission in all host species, we reconstructed phylogenies for host mitochondrial and symbiont core genomes and determined the concordance between tree topologies. Host mitochondrial genomes were assembled with MitoOmm (77) and annotated with GtSeq (78), using the A. boucheti (NC_049893.1) and l. nautilis (NC_024642.1) reference mitogenomes as bait for mitochondrial reconstruction and functional characterization. Concatenated alignments of 15 mitochondrial (13 protein-coding and 2 rRNA genes) and 222 to 1,679 symbiont-core specific genes were produced with Maft (79) in Geneious Prime (https://www.geneious.com) and Panaroo, respectively. Phylogenomic trees were calculated with IQTree 2 (80) in 10 independent runs using a gene-wise best-fit partition model selected with ModelFinder based on a greedy or relaxed hierarchical clustering strategy. Branch support was assessed through the ultrafast bootstrap approach and the Shimodaira-Hasegawa-like approximate-likelihood ratio test with 5,000 replicates by resampling both partitions and then sites within resampled partitions. To reduce the impact of model violations, each bootstrap tree was optimized using a hclimbing-nearest-neighbor interchange search (option --bnni). The best maximum-likelihood trees were midpoint-rooted with Pagcon (81) and cladograms were plotted with GGTREE (82) in R.

Population-Level Variant Identification in Hosts and Symbionts. Filtered metagenomic reads from each host individual were mapped against the corresponding host transcriptome and symbiont pan-genome with Bowtie 2 (83) in very sensitive mode. Optical duplicates were removed with Picard's MarkDuplicates tool (https://github.com/broadinstitute/picard). To resolve common alignment errors and improve base call accuracy, we locally realigned reads around indels and recalibrated base quality scores with LoFreq (84).

Host population genomic variation was assessed in Acesu (85) by inferring genotype likelihoods based on Hardy-Weinberg equilibrium considering individual inbreeding coefficients. To increase accuracy of the analyses, variant sites with mapping qualities <30 (minMapQ = 30), base qualities <20 (minQ = 20), and minimum minor-allele frequencies <0.01 (minMaf = 0.01) were excluded. We further filtered sites based on strand bias (sb_pval = 0.05), heterozygote bias (hetbias_pval = 0.05), and probability of being variable (SNP_pval = 1e-6). In addition, we removed spurious and improperly paired reads, adjusted mapping qualities for excessive mismatches (C = 50), and computed per-base alignment qualities (BAQ = 1) to disregard variants close to indel regions. Putative paralogous variants were excluded by discarding reads with multiple mappings and by considering only sites that had a maximum depth of 40 to 80 (based on distributions of average coverage). Genetic distances between individuals were inferred by calculating pairwise covariance matrices.

Symbiont population genomic variation was determined with FireBates (86) using input parameters adjusted for the analysis of metagenomic data (F 0.01 -C 1 -p 1 -pooled-continuous -haplotype-length 0 -report-monomorphic). Variant
calls were restricted to sites with a minimum base quality of 20, a minimum mapping quality of 30, and a minimum coverage of 10. To eliminate bias in variant identification and other downstream analyses due to uneven read depth between samples, we normalized all samples to the lowest amount of coverage found in a sample for a particular symbiont phylotype (>10x coverage). Variants were further filtered based on strand bias (SRP > 5 & SAP > 5 & EPP > 5), proximity to indels (5 bp), and maximum mean depth with Bcftools (87) and VcTools (88). In addition, sites and individuals with more than 25% missing data were excluded from the analysis. Allele counts (=symbiont strain abundances) and consensus haplotypes (=dominant symbiont strains) were extracted with GATK’s VariantTo Allele tool (89).

Population Genomic Structure and Differentiation. We performed ordination analyses with the Ape and Stats packages in R (90, 91) to assess genetic variation between symbiont and host populations. Host genetic structure was determined through principal-component analyses based on genetic covariance matrices, while symbiont genetic structure was inferred through principal-coordinate analyses based on both consensus haplotype and allele count data transformed into Euclidean distances and Bray-Curtis dissimilarities, respectively. For each sample, absolute allele counts were normalized to relative counts prior to analysis. Negative eigenvalues were corrected using the method by Cailliez (92) and final ordination plots were produced with GgAncor (93). FST values between host and symbiont populations were calculated in A构成了和 scikit-allel (https://github.com/cgh/scikit-allen), respectively, following the procedure in ref. 94. For the host species, potential outlier loci under selective pressures were inferred with OutFlank (95) based on neutral FST distributions that were obtained from quasi-independent SNP subsets determined with Punk (96).

Assessment of Symbiont Variation Based on Environment and Host Genetics. We conducted traditional and partial Mantel tests with the NCF package in R (97) to assess relationships between host identity-by-state distances, symbiont genetic distances, and geography based on Spearman rank correlations. Geographic distances were determined by calculating the geodesics between vent sites with the GeoSphere package (98). We used redundancy analyses following the approach in ref. 99 to evaluate the influence of hydrothermal fluid composition and depth on the genetic structure of each symbiont species. We further included the effect of sampling year, which encompasses changes in hydrothermal circulation within the ELSC as indicated by the cessation in fluid flow at the Kilo Moana vent field between 2009 and 2016. Endmember concentrations for geochemical compounds were obtained from the literature (25, 100, 101) or data provided by A. Diehl, MARUM, Bremen, Germany, and J. Seewald, Woods Hole Oceanographic Institution, Falmouth, MA (Dataset S1). Due to multicolinearity among the chemical species, we used the first eigenvector from principal-coordinate analyses (corresponding to the eigenvalue with the largest explanatory power) as the composite value. For each symbiont species, we further assessed the strength of correlation with other predictors to exclude variables that were highly collinear. Unless geographic location was strongly linked to environmental predictors, we used latitude as a conditioning factor in the analyses to correct for isolation by distance, given that vent sites are separated along a latitudinal range. Based on these corellarity evaluations, we tested the effect of all factors on the IberiaSea SOX symbiont, the effect of fluid composition on the A. bouchei Epsilon symbiont, and the effect of fluid composition and year (uncorrelated for geography) on the A. kojimai and A. strummeri Gamma1 symbionts. As the Gamma1 symbiont was present in two host species, we ran an additional model including host as explanatory variable. SNPs were considered candidates for local adaptation if their loadings on significant structured RDA axes deviated more than 2.5 SDs from the mean of the distribution. For each symbiont species, we further calculated the ratio of nonsynonymous polymorphisms per nonsynonymous site to synonymous polymorphisms per synonymous site (pN/pS) to infer candidate genes evolving under natural selection. The number of observed nonsynonymous and synonymous polymorphisms was determined with SnpEff (102), while the number of expected mutations was assessed with a modified Python script from ref. 103.

Assessment of Symbiont Gene Content Variation. We used PanPAN (104) to investigate potential differences in gene composition between symbiont populations from different vent localities. Downsized symbiont reads for each species were mapped against the corresponding symbiont pangeneome using custom annotations for functional categorizations. As exact zeros are not an appropriate threshold for considering a gene absent due to the possibility of mapping artifacts, PanPAN calculates the expected strain abundance by assessing the plateau in the coverage curves. Genes that are called absent through this method therefore include alignment errors but also genes that might be present in strains with very low abundance. Samples were profiled for gene presence/absence based on the following sensitive parameter thresholds: -min_coverage 1 -left_max 1.70 -right_min 0.30. Only samples that passed the retention thresholds in the variant-calling analyses were included. Gene content variation between symbiont strains was assessed by identifying genes that were present/absent in about 90% of samples from one or the other geographic region. For populations with small sample sizes (<5), we reduced this threshold down to 80% to avoid being overly stringent in determining differential gene preservation between locations. Presence/absence heatmaps were produced with the ComplexHeatmap package in R (105).

Data Availability. The raw Nanopore and Illumina reads, host transcriptomes, and symbiont reference genomes reported in this article have been deposited in the National Center for Biotechnology Information under BioProjects PRJNA523619, PRJNA562263, and PRJNA741492. Scripts for bioinformatic analyses are available on GitHub, https://github.com/cbreusing/Panvandid_host-symbiont_popgen. All other study data are included in the article and/or supporting information.

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PNAS 2022 Vol. 119 No. 14 e2115608119 https://doi.org/10.1073/pnas.2115608119 9 of 11
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