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New Sanger sequences for Alviniconcha and Ifremeria hosts and symbionts are available in GenBank under accession numbers MT131487–MT131781 (COI), MT130994–MT131149 (12S), MT146898–MT147385 (H3), MT148092–MT148633 (ATPSa), MT148634–MT149209 (ATPSb), MT147386–MT148091 (EF1a), and MT137388–MT137420 (16S). 16S amplicon sequences have been deposited in the Sequence Read Archive under BioProject numbers PRJNA473256, PRJNA473257, PRJNA610289, and PRJNA610290.

Abstract

Despite significant advances in our understanding of speciation in the marine environment, the mechanisms underlying evolutionary diversification in deep-sea habitats remain poorly investigated. Here, we used multigene molecular clocks and population genetic inferences to examine processes that led to the emergence of the six extant lineages of Alviniconcha snails, a key taxon inhabiting deep-sea hydrothermal vents in the Indo-Pacific Ocean. We show that both allopatric divergence through historical vicariance and ecological isolation due to niche segregation contributed to speciation in this genus. The split between the two major Alviniconcha clades (separating A. boucheti and A. marisindica from A. kojimai, A. hessleri, and A. strummeri) probably resulted from tectonic processes leading to geographic separation, whereas the splits between co-occurring species might have been influenced by ecological factors, such as the availability of specific chemosynthetic symbionts. Phylogenetic origin of the sixth species, Alviniconcha adamantis, remains uncertain, although its sister position to other extant Alviniconcha lineages indicates a possible ancestral relationship. This study lays a foundation for future genomic studies aimed at deciphering the roles of local adaptation, reproductive biology, and host–symbiont compatibility in speciation of these vent-restricted snails.

Key words: Alviniconcha, speciation, allopatric divergence, ecological isolation, deep-sea hydrothermal vents, chemosynthetic symbionts.

Introduction

How novel species evolve in response to environmental pressures remains a fundamental question in evolutionary biology [\(Schluter 1998\)](#page-17-0). New species can originate through a number of mechanisms, including geographic isolation (allopatric speciation) and formation of reproductive barriers due to localized natural selection (sympatric speciation) ([Darwin 1859;](#page-15-0) [Mayr 1942](#page-16-0); [Dobzhansky 1951](#page-15-0); [Coyne and](#page-15-0) [Orr 2004](#page-15-0); [Schuler et al. 2016](#page-17-0); [Yang et al. 2017\)](#page-17-0). The evolution of new species commonly occurs as a result of ecological or mutation-order speciation [\(Schluter 2009](#page-17-0)). Ecological speciation theory predicts that reproductive isolation should evolve between populations adapting to different environments [\(Mayr 1942\)](#page-16-0) and is a consequence of divergent selection on functional traits enabling specialization for distinct ecological niches [\(Rundle and Nosil 2005](#page-17-0); [Schluter 2009](#page-17-0)). Habitat isolation can occur at different spatial scales that range from "macrospatial" (a form of allopatric speciation, where gene flow between populations is absent due geographic separation) to "microspatial" (where distinct populations co-occur geographically but rarely interbreed because of adaptations to different ecological niches) [\(Coyne and Orr 2004\)](#page-15-0). By contrast, mutation-order speciation happens when reproductive isolation evolves due to random fixation of alleles in two populations that face the same selective pressures ([Mani and Clarke 1990](#page-16-0); [Schluter](#page-17-0) [2009](#page-17-0); [Nosil and Flaxman 2011\)](#page-16-0). Evolutionary divergence under this scenario is unlikely (though not impossible) if gene flow is present ([Schluter 2009](#page-17-0); [Nosil and Flaxman 2011](#page-16-0)), whereas ecological speciation is common with or without gene flow [\(Schluter 2009](#page-17-0); [Nosil and Flaxman 2011;](#page-16-0) [Feder](#page-16-0)

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[et al. 2012](#page-16-0); [Campbell et al. 2018\)](#page-15-0). Additional processes including inter-specific hybridization and host–microbe associations have received less attention but are now accepted as significant drivers of speciation [\(Schuler et al. 2016\)](#page-17-0).

Our current understanding of speciation mechanisms in animals stems mostly from evolutionary research on terrestrial and freshwater organisms, whereas less is known about speciation in oceanic environments, in particular deep ocean basins ([Miglietta et al. 2011](#page-16-0)). Processes that lead to evolutionary divergence in marine systems may differ from other environments given that physical barriers are less pronounced and deep-sea habitats might have been less affected by past climatic fluctuations ([Miglietta et al. 2011;](#page-16-0) but see [Vrijenhoek 2013\)](#page-17-0), at least over much of the Cenozoic. Nonetheless, deep-sea hydrothermal vents provide an exception to other deep-sea environments due to their global but island-like distribution and their relative instability on ecological and geological time scales [\(Vrijenhoek 2013](#page-17-0)). The dense animal communities living in these habitats are exposed to highly variable and extreme environmental conditions, and many of the invertebrates are sustained through obligate symbiosis with chemosynthetic bacteria [\(Dubilier et al.](#page-15-0) [2008\)](#page-15-0). The phylogenetically and ecologically closely related snail genera Alviniconcha and Ifremeria (Gastropoda: Abyssochrysoidea) are among the dominant inhabitants of Indo-Pacific hydrothermal vents. The genus Ifremeria is monotypic and apparently restricted to Western Pacific vents in the Manus, Lau, and North Fiji basins. In contrast, the broadly distributed genus Alviniconcha comprises six species ([Johnson et al. 2015](#page-16-0)): Alviniconcha kojimai and Alviniconcha boucheti co-occur in the Manus, Lau, and North Fiji basins; Alviniconcha strummeri is known from the southern Lau Basin; Alviniconcha hessleri and Alviniconcha adamantis appear to be endemic to the Mariana Back-Arc and Volcanic Arc, respectively; and Alviniconcha marisindica occurs along the Central Indian Ridge (fig. 1 and [table 1](#page-4-0)).

Alviniconcha and Ifremeria are nutritionally dependent on horizontally transmitted, chemosynthetic sulfide-oxidizing endosymbionts that are acquired from a diverse community of environmental bacteria and are harbored within vacuoles in the gill filaments ([Windoffer and Giere 1997](#page-17-0); Suzuki, Sasaki, Suzuki, Nogi, et al. 2006; [Trembath-Reichert et al. 2019\)](#page-17-0). Except for A. boucheti and A. marisindica, which are constrained to symbionts of the class Campylobacteria (formerly Epsilonproteobacteria; [Waite et al. 2017\)](#page-17-0), all other Alviniconcha species and Ifremeria nautilei typically host Gammaproteobacteria symbionts [\(Urakawa et al. 2005;](#page-17-0) [Suzuki, Sasaki, Suzuki, Nogi, et al. 2005;](#page-17-0) Suzuki, Sasaki, Suzuki, Tsuchida, et al. [2005](#page-17-0); Suzuki, Kojima, Sasaki, et al. [2006;](#page-17-0) Suzuki, Kojima, Watanabe, et al. [2006;](#page-17-0) [Beinart et al.](#page-15-0) [2012\)](#page-15-0). In zones of sympatry, the two snail genera occupy relatively narrow ranges of chemical and thermal conditions that are well within their physiological tolerance limits and that do not support maximal rates for chemoautotrophic growth ([Sen et al. 2013\)](#page-17-0). This observation suggests that interspecies competition is an important factor determining the realized niche of these snails [\(Sen et al. 2013](#page-17-0)). [Beinart et al.](#page-15-0) [\(2012\)](#page-15-0) proposed that host niche utilization is strongly influenced by the availability and metabolic capacity of locally dominant symbiont phylotypes. In their study of co-occurring Alviniconcha species from the Lau Basin, they showed that the distribution and abundance of host–symbiont combinations (holobionts) followed a latitudinal gradient in vent geochemistry, wherein A. boucheti holobionts dominated at northern localities characterized by high end-member concentrations of sulfide and hydrogen (\sim 3.5–7 mM and \sim 100– 500μ M, respectively), A. kojimai holobionts dominated at midlatitude localities characterized by medium endmember concentrations (\sim 2.5–4 mM and \sim 50–100 µM, respectively), and A. strummeri holobionts dominated at southern localities characterized by low end-member concentrations of these reductants $(\sim 1-3$ mM and 35-135 μ M, respectively). These broad-scale patterns of

NOTE.-Submersibles: H, Hercules; J2, Jason II; S, Shinkai 6500; Su, SuBastian; R, Ropos; ST, ST212; KI, Kiel 6000; MQ, Marum Quest.

microbe-associated habitat segregation suggest that the endosymbionts might have played a significant role in the diversification of these gastropods.

Despite their phylogenetic and ecological affinities, Ifremeria and Alviniconcha have undergone remarkably different degrees of speciation. In the present study, we used fossil-calibrated molecular phylogenies and population genetic analyses based on mitochondrial and nuclear markers to obtain insights into allopatric and sympatric mechanisms that led to the high degree of diversification in the genus Alviniconcha. To assess the role of allopatric isolation, we compared the phylogenetic divergence events among the Abyssochrysoidea with Mesozoic and Cenozoic geological events of the Indo-Pacific Ocean basins and examined genetic subdivision within several broadly distributed species.

To determine evidence for sympatric isolation, we compared the time-calibrated phylogenies with the diversity and phylogenetic relationships of the snails' chemosynthetic endosymbionts and assessed the potential for hybridization among co-occurring Alviniconcha species.

Results

Gene Networks

Haplotype networks for the two mitochondrial genes, COI and 12S, indicated complete sorting among lineages for the six Alviniconcha species and their sister taxon I. nautilei [\(fig. 2\)](#page-5-0), as previously reported [\(Johnson et al. 2015\)](#page-16-0). The COI sequences from Alviniconcha and Ifremeria differed by at least 75 mutations, whereas pairs of Alviniconcha species differed by

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FIG. 2. Haplotype networks for the two mitochondrial (COI and 12S) and four nuclear genes (H3, EF1a, ATPSa, and ATPSb) that were used for population genetic analyses. Circles represent individual haplotypes, where circle size is proportional to haplotype frequency and black circles indicate unsampled haplotypes. Dashes on branches denote the number of mutations between haplotypes.

25–48 mutations. The smallest divergence was observed between A. kojimai and A. hessleri and the largest between A. strummeri and A. adamantis. Sequence divergence was lower for the 12S locus, with a maximum of 19 substitutions between Alviniconcha species pairs (fig. 2). The nuclear markers generally confirmed species identities as seen with the mitochondrial loci, although they were less diagnostic due to varying degrees of incomplete lineage sorting and smaller divergence between sequences (fig. 2). Shared polymorphisms were particularly evident between A. kojimai and A. hessleri, as well as between A. boucheti and A. marisindica, whereas A. strummeri and A. adamantis had higher amounts of private alleles.

Population Structure and Admixture

Structure analyses distinguished the six previously described Alviniconcha species ([Johnson et al. 2015\)](#page-16-0) based on mitochondrial and nuclear markers [\(fig. 3](#page-6-0) and supplementary fig. S1, Supplementary Material online). Runs including the mitochondrial COI gene provided in general higher resolution of population genetic subdivision than runs based on nuclear loci alone (supplementary figs. S1-S6, Supplementary

FIG. 3. Genetic structure of Alviniconcha species (A) and populations (B-F). Each color represents a different genetic cluster (K), whereas each vertical bar represents a distinct individual, where colored line segments are proportional to genetic group memberships. Black lines divide individuals from distinct sampling locations noted along the x-axis. Lau Basin sites: KM, Kilo Moana; TC, Tow Cam; TM, Tu'i Malila. North Fiji Basin sites: WL, White Lady; MH, Mussel Hill; WR, White Rhino. Manus Basin sites: SW8, Solwara 8; SW1, Solwara 1; SSU, South Su; VA, Vanuatu. Mariana Back-Arc sites: FC, Forecast; SN, Snail Site; EDS, E. Diamante Seamount. Central Indian Ridge sites: KA, Kairei; ED, Edmunds.

Material online). Therefore, we report results from the combined marker set in the manuscript, but Structure plots for all K values and marker combinations can be found in the supplementary figures S1–S6, Supplementary Material online. Although the most parsimonious grouping following the method by [Evanno et al. \(2005\)](#page-16-0) was $K = 2$, visual inspections of higher K values [\(Meirmans 2015\)](#page-16-0) showed clear distinctions of up to eight clusters, indicating population structure within the sympatric species A. boucheti, A. kojimai, and A. strummeri (fig. 3 and supplementary figs. S1–S4, Supplementary Material online). For all three species, significant genetic variation among individuals within vent localities was observed (fig. 3).

Alviniconcha boucheti comprised six genetically connected clusters that were broadly partitioned across four geographic regions: 1) Vanuatu and Manus Basin, 2) Kilo Moana (Lau Basin), White Lady, and Mussel Hill (North Fiji Basin), 3) Niua

NOTE. ABBA and BABA sites are parsimony informative sites that result in discordances between gene trees and species trees due to the presence of incomplete lineage sorting or gene flow. ABBA and BABA sites are expected to be equally frequent under incomplete lineage sorting but will have uneven abundances if gene flow is present. $D =$ Patterson's D estimate based on Jackknife resampling. A significant $D > 0$ indicates gene flow between P3 and P2, whereas a significant $D < 0$ indicates gene flow between P3 and P1. P values computed by Fisher's combined probability test were not significant. $Z = Z$ score of Patterson's D.

South and ABE (Lau Basin), and 4) White Rhino (North Fiji Basin). Alviniconcha kojimai was represented by four genetic clusters that were subdivided across the 1) Manus Basin and 2) North Fiji Basin, with Lau Basin populations showing similarities to both of these groups. Alviniconcha strummeri exhibited less genetic partitioning, with the Tu'i Malila locality slightly differentiated from the Tahi Moana locality in the Lau Basin. Populations of A. hessleri and A. marisindica were not structured across the sampled geographic regions, but limited genetic differences were present among individuals [\(fig. 3](#page-6-0) and supplementary figs. S5 and S6, Supplementary Material online). Despite the presence of shared polymorphisms, ABBA– BABA tests provided no evidence for hybridization between sympatric species, indicating that patterns of shared genetic variation were due to incomplete lineage sorting (table 2). Note, however, that D values for the first ABBA–BABA test were close to -1 , and it is possible that gene flow between A. kojimai and A. boucheti might have been detected with a larger genomic marker set.

Fossil-Calibrated Phylogeny and Species Tree

Fossil-calibrated Beast analyses based on ribosomal, mitochondrial, and histone marker genes revealed that the Abyssochrysidae is a relatively old family ([fig. 4\)](#page-8-0), having diverged from the Littorinidae and Buccinidae in the early Jurassic ${\sim}$ 200 Ma (95% Highest Posterior Density (HPD): 172–234 Ma). This range agrees well with the divergence times reported by [Johnson et al. \(2010\)](#page-16-0) (93–228 Ma) but is slightly older than estimates by [Osca et al. \(2014\)](#page-17-0) (144–168 Ma). The split between I. nautilei and Alviniconcha spp. occurred in the mid-Cretaceous \sim 113 Ma (95% HPD: 96–131 Ma). The extant Alviniconcha lineages began to diversify in the Eocene \sim 48 Ma (95% HPD: 39–56 Ma) and comprised two well-supported clades plus A. adamantis, which formed a basal, unresolved branch that split ${\sim}$ 42 Ma (95% HPD: 34–49 Ma). The first well-supported clade included A. strummeri, A. kojimai, and A. hessleri and diverged \sim 25 Ma (95% HPD: 18–31 Ma). The second clade included A. marisindica and A. *boucheti w*ith a divergence time of \sim 38 Ma (95% HPD: 30– 46 Ma). The most recent split occurred between A. kojimai and A. *hessleri* in the later Miocene \sim 10 Ma (95% HPD: 7–13 Ma).

In general, the topology of the species tree for Alviniconcha agreed with that of the concatenated gene tree, although the position of A. adamantis indicated a potential sister relationship to A. boucheti and A. marisindica ([fig. 4\)](#page-8-0). The second species tree including the nuclear markers EF1a, ATPSb, and ATPSa resulted in an unresolved polytomy and we were

unable to obtain robust estimates for phylogenetic placements (supplementary fig. S7, Supplementary Material online).

Phylogenetics of Endosymbionts

Molecular Bayesian analyses of the full-length 16S rRNA revealed geographical partitioning for the endosymbionts of Alviniconcha and Ifremeria, suggesting that symbiont phylotypes associated with I. nautilei, A. boucheti, and A. kojimai were genetically distinct between the Lau, Manus, and North Fiji basins ([fig. 5](#page-9-0)). The endosymbionts of *I. nautilei* were most closely related to free-living Thiolapillus brandeum and the Gammaproteobacteria symbiont of A. hessleri. Together, these bacterial groups formed a neighboring clade to the " γ -1" endosymbionts of A. kojimai ([Beinart et al. 2012](#page-15-0)) and free-living Thiomicrospira and Hydrogenovibrio ([fig. 5](#page-9-0)A). In both the Lau and Manus basins, A. kojimai additionally hosted Campylobacteria lineages that were also found in A. boucheti. The endosymbionts of A. adamantis formed a distinct clade that was most closely related to the " γ -Lau" group present in A. strummeri [\(Beinart et al. 2012](#page-15-0)).

Alviniconcha marisindica and A. boucheti were the only species that harbored predominantly Campylobacteria sym-bionts [\(fig. 5](#page-9-0)B). Although A. marisindica contained a single 16S rRNA phylotype related to free-living Sulfurovum, A. boucheti associated with distinct Campylobacteria lineages that were related to free-living Sulfurimonas, with one exception: A. boucheti from American Samoa harbored a phylotype that was most closely related to the A. marisindica symbiont.

Character traces of symbiont type identified the Gammaproteobacteria endosymbiosis as the ancestral condition given that the evolutionary older host species harbored only Gammaproteobacteria [\(fig. 6](#page-10-0)). By contrast, the Campylobacteria endosymbiosis appears to be a derived trait that evolved multiple times independently.

Symbiont Phylotypic Diversity and Partitioning

Amplicon sequencing of the 16S V4–V5 hypervariable regions generated an average of 21,302 paired-end reads/sample. These were reduced to \sim 9,613 reads/sample after read merging. Sequence denoising resulted in 85 zero-radius operational taxonomic units (ZOTUs), 11 of which were verified as Alviniconcha or Ifremeria endosymbionts. Oligotyping of amplicon reads comprising these symbiont ZOTUs identified a total of 23 symbiont 16S rRNA oligotypes in the data set. For 337 snail samples, we were able to obtain both host species identities and 16S oligotypes. Only these individuals were,

FIG. 4. Fossil-calibrated phylogeny of the Abyssochrysidae. Scale bar is 30 Myr, stars on branches indicate 1.0 support value. Gray bars indicate the highest posterior probability density range for estimates of lineage splitting times. The legend provides symbol codes for the geographic distribution, endosymbiont affinity, and putative sympatric (white circle) and allopatric (gray circle) divergence events of the different Alviniconcha lineages. If both white and gray circles are shown under a branch, both allopatric and sympatric processes might have contributed to the respective divergence event. Inset in the upper right shows the species tree obtained with StarBeast, with gray shading indicating uncertainties in the phylogenetic placements.

therefore, used for further analysis (supplementary table S1, Supplementary Material online).

Each host individual contained one majority symbiont phylotype together with up to 12 minority variants, with A. hessleri exhibiting the highest and A. boucheti, A. kojimai, and A. strummeri exhibiting the lowest symbiont richness ([fig. 7\)](#page-10-0). In general, symbiont phylotypes corresponded to host species, except in the case of A. kojimai and A. strummeri, where symbiont compositions largely overlapped [\(figs. 7 and 8\)](#page-10-0). All host species, with the exception of A. boucheti, comprised mostly a combination of different Gammaproteobacteria lineages. Alviniconcha boucheti was dominated by Campylobacteria symbiont phylotypes, although in the Lau Basin this species also harbored Gammaproteobacteria symbionts that were typically associated with A. kojimai and A. strummeri. However, due to presence of the A. kojimai symbiont (Oligo1) in negative controls on the same well plate where these A. boucheti samples were processed, we cannot exclude the possibility that some of these observations resulted from well-to-well contamination. In accordance with our phylogenetic analyses, the symbionts associated with A. boucheti were partitioned across basins. In most cases, we did not have sufficient sample sizes to reveal potential within-basin structuring of the Gammaproteobacteria or Campylobacteria symbiont phylotypes, with the exception of the A. hessleri symbionts, which seemed to form two latitudinal clusters [\(fig. 8\)](#page-11-0).

Partial Mantel tests supported the described patterns, indicating significant correlations between symbiont and host genetic distances ($r = 0.76$; P = 1e-04) as well as between symbiont genetic distance and geographic distance $(r = 0.55; P = 1e-04)$. By contrast, no association between host genetic distance and geographic distance was found.

Discussion

In the present study, we performed fossil-calibrated phylogenetic reconstructions and population genetic analyses to illuminate mechanisms of speciation in provannid snails of the genus Alviniconcha, which are dominant members of deepsea hydrothermal vent ecosystems in the Western Pacific and Indian Ocean.

Major tectonic and climatic changes in the Indo-Pacific during the Mesozoic and Cenozoic probably played a central

Campylobacteria **B**

FIG. 5. Phylogenies of the (A) Gammaproteobacteria and (B) Campylobacteria symbiont lineages. Node labels indicate node support probabilities, whereas scale bars show the expected number of substitutions per nucleotide site. Color codes are based on host species identities as shown in [figures 1](#page-3-0) and [2](#page-5-0).

role for geographic isolation during the early evolution of Alviniconcha snails. Our fossil-calibrated phylogeny indicates that Ifremeria and Alviniconcha split 96–131 Ma close to the

peak of the Aptian oceanic anoxic event [\(Li et al. 2008\)](#page-16-0). Although anoxic events led to widespread declines of marine biodiversity, they also created novel opportunities for habitat specialization and allopatric speciation within oxygendepleted deep-sea environments, such as hydrothermal vents ([Rogers 2000\)](#page-17-0). Similarly, expansion of oxygen minimum zones might have contributed to the evolution of the most recent common ancestor of extant Alviniconcha, which originated during the upper Eocene (\sim 50 Ma) shortly after the Paleocene–Eocene Thermal Maximum, another period that was characterized by increasing anoxia in the deep sea. Around the same time, the Philippine Sea Plate developed above the Manus hotspot, and a wide deep-water passage connected the Indian and Pacific Oceans ([Hall et al. 1995;](#page-16-0) [Hall](#page-16-0) [2002;](#page-16-0) see figs. 14–25 in [Hall 2002\)](#page-16-0). This passage started to narrow around 30 Ma and eventually closed 25 Ma, when the Australian Plate subducted under the Philippine Sea Plate and created physical barriers to dispersal (Gaina and Müller 2007).

Gradual vanishing of the Indo-Pacific deep-water connection is consistent with the timing of the most recent common

FIG. 6. Analysis of endosymbiont trait evolution in Alviniconcha. Associations with Gammaproteobacteria lineages represent the ancestral condition, whereas the Campylobacteria endosymbiosis is a derived state.

ancestors of the two major Alviniconcha clades and likely promoted their diversification: 1) A. marisindica and A. boucheti (\sim 30–46 Ma) and 2) A. hessleri, A. kojimai, and A. strummeri (\sim 18–31 Ma). A variety of along-arc splitting and rifting events in the Miocene to Pliocene subsequently led to the formation of the North Fiji Basin (\sim 12 Ma), Mariana Back-Arc Basin (\sim 7 Ma), and Lau Basin (\sim 5 Ma) [\(Hall 2002](#page-16-0); [Crawford et al. 2003;](#page-15-0) [Straub et al. 2015](#page-17-0)). Evolution of the North Fiji Basin and Mariana Back-Arc Basin corresponds well with the divergence of A. hessleri and A. kojimai (\sim 10 Ma), suggesting a potentially allopatric origin of these lineages that might have allowed A. hessleri to invade deep-water vent sites of the Mariana Back-Arc, where it is nowadays an endemic species. Alviniconcha adamantis is the only other Alviniconcha species that occurs in the Mariana region, but it is restricted to shallow vents on volcanoes of the Izu-Bonin-Mariana Arc, which have been active in some form for 40 My ([Stern et al. 2003\)](#page-17-0). Interestingly, the phylogenetic position of A. adamantis was not well supported and formed a basal branch to the two other clades of Alviniconcha. Although we do not have enough data to determine its placement, the A. adamantis lineage might be ancestral to other Alviniconcha lineages. If true, this genus might have followed a gradual transition from shallow to deep-water vent sites, as has been proposed for bathymodiolin mussels (e.g., [Jones](#page-16-0) [et al. 2006;](#page-16-0) [Miyazaki et al. 2010;](#page-16-0) but see [Thubaut et al. 2013\)](#page-17-0).

Although our results suggest that splits between some Alviniconcha species involved geographic subdivision, allopatric divergence alone might be insufficient to explain all instances of speciation. For example, A. boucheti, A. kojimai, and A. strummeri live sympatrically in the Lau Basin, and A. boucheti and A. kojimai also coinhabit vents in the North Fiji and Manus basins. For both of these species, genetic structure analyses indicated varying degrees of genetic

FIG. 7. (A) Fractional abundances and (B) phylogenetic relationships of symbiont 16S V4-V5 oligotypes in Alviniconcha (163 specimens) and Ifremeria (173 specimens).

FIG. 8. (A) Nonmetric dimensional scaling plot of the symbiont 16S phylotypes based on weighted Unifrac distances. The plot shows that the symbiont composition correlates with the type of host species and is partitioned across back-arc basins for Alviniconcha boucheti. (B) Detailed nonmetric dimensional scaling plots for symbiont phylotypes associated with Alviniconcha hessleri. Symbiont populations form two latitudinal clusters corresponding to \sim 16 $^{\circ}$ N (Hafa Adai, Perseverance) and \sim 18 $^{\circ}$ N (Burke, Illium).

subdivision not only among but also within disparate backarc basins. However, based on recent biophysical models for the Western Pacific region [\(Mitarai et al. 2016](#page-16-0)), ocean circulation should support within-basin connectivity, especially for species that are assumed to have planktotrophic larvae with high dispersal potentials, such as Alviniconcha [\(War](#page-17-0)é[n and](#page-17-0) [Bouchet 1993](#page-17-0)). These contrasting findings suggest that antagonistic ecological factors might exert strong selection against immigrants with maladapted phenotypes, thereby counterbalancing the homogenizing effect of dispersal and allowing speciation in the presence of gene flow. The present genetic data do not allow us to confidently distinguish whether the observed speciation patterns occurred during primary sympatry or resulted from secondary colonization following allopatry. At least the ancestors of A. kojimai and A. strummeri likely co-occurred in the predecessor of the modern Manus Basin and, thus, may have evolved sympatrically. Nevertheless, opening of the younger Lau and North Fiji basins could have allowed for brief geographic separation (<5–10 Ma), if species invaded these regions asynchronously.

Hydrothermal vents are environmentally dynamic habitats that offer ample opportunities for niche segregation along steep thermal and geochemical gradients. Geochemical patchiness and the availability and/or fitness of symbiont phylotypes (rather than functional host anatomy) are believed to affect spatial partitioning among Alviniconcha species in the Eastern Lau Spreading Center [\(Beinart et al. 2012;](#page-15-0) [Laming et al. 2020\)](#page-16-0). Alviniconcha boucheti, which typically hosts Campylobacteria symbionts, was most abundant at northern vent localities characterized by high sulfide and hydrogen concentrations. On the other hand, A. kojimai and

bionts, were more frequent at southern vent localities with lower sulfide and hydrogen concentrations. Functional differences among the symbiont types combined with genetically based host–symbiont specificities could explain the broadscale segregated distribution of these species and might have provided a mechanism for their diversification. Free-living chemosynthetic Campylobacteria are known to encode sulfide-oxidizing enzymes that are most effective at high $H₂S$ concentrations [\(Han and Perner 2015](#page-16-0)), providing a selective advantage over Gammaproteobacteria at sulfide-rich vents [\(Nakagawa and Takai 2008;](#page-16-0) [Yamamoto and Takai](#page-17-0) [2011;](#page-17-0) [Patwardhan et al. 2018\)](#page-17-0). Recent physiological experiments and transcriptomic analyses imply that this might also be the case for the Lau Basin Alviniconcha symbionts [\(Breusing et al. 2020](#page-15-0)). Possible relationships between vent geochemistry, symbiont phylotypes, and host species have not been similarly explored in other back-arc basins. Nevertheless, the PACMANUS and Vienna Woods vent fields in the Manus Basin are known to exhibit physicochemical gradients similar to those found in the Lau Basin ([Reeves et al.](#page-17-0) [2011\)](#page-17-0), providing comparable opportunities for symbiontrelated niche differentiation. Our partial Mantel tests and NMDS analyses involving Alviniconcha species from the Lau and Manus basins, and the Mariana Back-Arc revealed a significant association of symbiont genotypes with host species and vent localities, suggesting that symbiont-mediated habitat segregation might be common in chemosynthetic vent snail symbioses. Together, our host phylogenetic and bacterial phylotype

A. strummeri, which usually host Gammaproteobacteria sym-

analyses imply a possible role for endosymbionts in

Alviniconcha speciation. For instance, A. boucheti and A. marisindica comprise a monophyletic clade associated predominantly with Campylobacteria symbionts. A trace of character trait evolution indicated that Campylobacteria endosymbionts were acquired in the most recent common ancestor of these two host species, possibly prior to geological events that contributed to splitting of the A. boucheti–A. marisindica clade. It is possible that a switch to a new symbiont type offered novel ecological opportunities, thereby promoting adaptive divergence from the Gammaproteobacteria-dominated Alviniconcha lineages. Both A. boucheti and A. marisindica associate with symbionts related to free-living Sulfurovum, whereas A. boucheti also associates with symbionts of the genus Sulfurimonas, indicating that the Sulfurovum endosymbiosis might be the evolutionarily older condition in these two host species. Speciation due to symbiont-driven niche expansion has been proposed for deep-sea mytilid mussels, with similar time intervals of diversification as observed for provannid snails [\(Lorion et al. 2013](#page-16-0)). Interestingly, geographic partitioning of the A. boucheti symbiont phylotypes corresponded broadly with the genetic structure of the host populations. Although we cannot exclude the hypothesis that these covarying patterns were mediated by geographic separation, gene flow between basins is expected on evolutionary time scales ([Mitarai et al. 2016](#page-16-0)); therefore, genetically based host–symbiont specificities might also shape the diversification of both partners.

Presently, no substantive evidence exists for past hybridization between broadly co-occurring A. boucheti and A. kojimai (although future genomic studies may provide other insights into this question). Host–symbiont incompatibilities might contribute to hybrid dysgenesis, a scenario that has been described in insect–microbe symbioses (e.g., [Feder](#page-16-0) [et al. 1994;](#page-16-0) [Brucker and Bordenstein 2012,](#page-15-0) [2013\)](#page-15-0). Although A. kojimai occurs in association with Campylobacteria symbionts, these observations were relatively rare, indicating a bias toward a Gammaproteobacteria endosymbiosis ([Beinart et al. 2012\)](#page-15-0). Microbial symbionts are currently considered the most disregarded element in animal and plant evolution [\(Shropshire and Bordenstein 2016](#page-17-0)). A recent study has shown that both transmission mode and symbiont function influence the degree of host dependence that can ultimately lead to major evolutionary transitions and innovations ([Fisher et al. 2017\)](#page-16-0). Although strong host dependence is usually correlated with vertical transmission mode, the same effect can be achieved in horizontally transmitted symbioses if symbionts provide an essential nutritional benefit [\(Fisher](#page-16-0) [et al. 2017](#page-16-0)). Consequently, although further studies are needed to characterize the role of symbionts in host speciation, their impact is likely of particular significance in chemoand photosynthetic ecosystems where host fitness is tightly linked to symbiont metabolism, such as hydrothermal vents, cold seeps, and coral reefs.

An alternative, though not mutually exclusive, scenario for sympatric speciation in Alviniconcha could be rapid reproductive isolation and sexual selection. In many organisms, including marine snails of the genera Haliotis and Tegula, egg and sperm proteins are under strong positive selection and evolve at an accelerated rate that could ultimately split populations into reproductively isolated species [\(Galindo](#page-16-0) [et al. 2003;](#page-16-0) [Panhuis et al. 2006;](#page-17-0) [Hellberg et al. 2012\)](#page-16-0). Subsequent inter-specific gene flow could be reduced further due to assortative mating, which is often observed in gastropods with internal fertilization ([Johannesson et al. 2008;](#page-16-0) [Zahradnik et al. 2008](#page-17-0)). Although little is known about the reproductive biology of Provannidae, internal fertilization is the implied reproductive mode in this family [\(Gustafson and](#page-16-0) [Lutz 1994](#page-16-0)).

Our study suggests that both geographical and ecological divergence (followed by putative reproductive isolation) played significant roles in the evolutionary diversification of the six extant Alviniconcha lineages at Indo-Pacific hydrothermal vents. It remains to be determined why the same phenomena did not similarly affect speciation within the genus Ifremeria, which is represented by only a single species. The bounded hypothesis on diversification proposes that the evolution of new species is constrained by competition for available niche space (e.g., [Rabosky 2009](#page-17-0); [Price et al. 2014;](#page-17-0) [Larcombe et al. 2018\)](#page-16-0) and that an increase in the number of species within one clade may be compensated by a decline in another [\(Ricklefs 2010\)](#page-17-0). Given that Ifremeria has similar ecological requirements as Alviniconcha ([Podowski et al.](#page-17-0) [2010\)](#page-17-0), it is possible that competitive exclusion and niche filling might have limited diversification within this genus. Future genomic and ecological studies may be helpful to illuminate some of these questions and to assess the relative importance of host–symbiont incompatibilities, local adaptation, and reproductive barriers in the evolutionary history of these gastropods.

Materials and Methods

Sample Collection and DNA Methods

Snail samples were collected from 32 localities in the Western Pacific and Indian Oceans [\(table 1\)](#page-4-0). DNA extraction, general polymerase chain reaction (PCR) conditions, amplicon purification, and DNA sequencing used methods that were reported for provannid snails ([Johnson et al. 2010,](#page-16-0) [2015\)](#page-16-0). Twelve primer sets were used to amplify three mitochondrial (COI, mt-12S, and mt-16S) and seven nuclear (H3, 28S-D1, 28S-D6, 18S, ATPSa, ATPSb, and EF1a) DNA targets from vent snail hosts as well as the 16S rRNA locus from the snails' chemo-synthetic symbionts [\(table 3\)](#page-13-0). The 16S rRNA gene of gill endosymbionts of two A. *adamantis* and five *I. nautilei* individuals was cloned prior to Sanger sequencing [\(Goffredi et al. 2007](#page-16-0)) with 24 colonies per individual. All PCR products were diluted in 40–50 μ of sterile water and purified with a Multiscreen HTS PCR 96 vacuum manifold system (Millipore Corp., Billerica, MA). PCR products were sequenced bidirectionally on an ABI 3130XL sequencer with BigDye Terminator v3.1 chemistry (Life Technologies Corp., Carlsbad, CA) and primers used in PCR. Samples from American Samoa were added late to the study and were only included in phylogenetic analyses of the symbionts. Forward and reverse sequences were trimmed, merged, and manually curated in Geneious Prime [\(http://www.geneious.com; last accessed July 24, 2020\)](http://www.geneious.com).

Table 3. PCR Primers and Methods for Gene Loci Used in Alviniconcha spp. and Ifremeria nautilei.

^aPCR program: 95 °C/10 min; 35 \times [94 °C/1 min, 55 °C/1 min; 72 °C/1 min], extension at 72 °C/7 min.
^bTouchdown and East PCP: Amplitag Gold East PCP Master Miv. LIP (Life Technologies Corp.) and the r

 b Touchdown and Fast PCR: Amplitaq Gold Fast PCR Master Mix, UP (Life Technologies Corp.) and the protocol for the Taq supplied by the manufacturer of the Veriti thermal cycler with an annealing temperature at 50 \degree C (Life Technologies Corp.).

Wodified after [Jarman et al. \(2002\),](#page-16-0) ATPSb-AlviniF: 5'-TCAGCCTCACCGATGACACCT-3' and ATPSb-AlviniR: 5'-CRGGGGGYTCRTTCATCT-3'.
^dPCP program: 95 °C/10 min: 25 × 102 °C/1 min: 53 °C/1 min: 72 °C/1 min), extension at 72 °C/7

^dPCR program: 95 °C/10 min; 25 \times [92 °C/1 min, 53 °C/1 min; 72 °C/1 min], extension at 72 °C/7 min.

Heterozygous sites at nuclear loci were phased with Phase v2.1.1 ([Stephens and Donnelly 2003](#page-17-0)) as in [Breusing et al.](#page-15-0) [\(2015\)](#page-15-0).

Haplotype Networks, Genetic Structure, and Admixture

Haplotype networks for two mitochondrial genes (COI, 12S) and four nuclear genes (H3, EF1a, ATPSa, and ATPSb) were constructed in Popart v1.7 [\(http://popart.otago.ac.nz/; last](http://popart.otago.ac.nz/) [accessed July 24, 2020](http://popart.otago.ac.nz/)) with the median-joining algorithm as in [Breusing et al. \(2019\)](#page-15-0).

Structure v.3.5 [\(Pritchard et al. 2000](#page-17-0)) was applied to investigate the number of genetic clusters K in the total Alviniconcha sample set and to test for contemporary admixture in the co-occurring species A. boucheti, A. kojimai, and A. strummeri. Analyses were run for all six species together as well as for each species separately using 1) mitochondrial and nuclear data and 2) nuclear data only. Allelic information was provided as a number code. Inter-specific Structure analyses were based on five (COI, H3, EF1a, ATPSa, and ATPSb) and four genetic loci (H3, EF1a, ATPSa, and ATPSb), respectively. The same genetic markers were used in intra-specific analyses, except that we excluded the H3 locus given that it was essentially monomorphic within Alviniconcha populations. Model runs and settings for Structure generally followed [Breusing et al. \(2016\)](#page-15-0) with ten repetitions for each K between 1 and 8 for inter-specific analyses and between 1 and 6 for intra-specific analyses. We subsequently used Clumpak v.1.1 ([Kopelman et al. 2015\)](#page-16-0) to integrate results across independent runs for each K value and determine the optimal clustering solution. We initially used the ΔK method [\(Evanno](#page-16-0) [et al. 2005](#page-16-0)) to objectively infer the number of genetic clusters in the data set. However, as this method is biased toward identifying the uppermost hierarchical level of population structure (i.e., lowest parsimonious K value) [\(Cullingham](#page-15-0) [et al. 2020](#page-15-0)), we visually inspected all K plots for biologically

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meaningful structuring and chose the highest K value where genetic subdivision could be observed ([Meirmans 2015](#page-16-0)).

To distinguish hybridization from incomplete lineage sorting in the sympatric species A. boucheti, A. kojimai, and A. strummeri, we performed two ABBA–BABA tests on the concatenated nuclear gene alignments of 90 snail individuals using HybridCheck [\(Ward and van Oosterhout 2016\)](#page-17-0). Gene flow was assessed between 1) A. boucheti (P3) and A. kojimai/ A. strummeri $(P1 + P2)$ and 2) A. strummeri (P3) and A. kojimai/A. hessleri ($P1 + P2$), with A. adamantis and A. boucheti as outgroups, respectively. Sequences from I. nautilei were not used because of missing data. Jackknife sampling was performed on four blocks of 305 bp (i.e., the approximate size of each nonlinked locus).

Fossil-Calibrated Phylogeny of the Abyssochrysidae

All Alviniconcha species were included in a fossil-calibrated phylogenetic reconstruction of the Abyssochrysidae with the program Beast v.2.5.7 [\(Bouckaert et al. 2014\)](#page-15-0). Fossil calibrations included the divergence of Neptunea amianta and N. antiqua, which have probably been separated since the earliest appearance of the genus in the Late Eocene, 33–37 Ma [\(Amano 1997](#page-15-0)), as well as Provanna and Desbruyeresia, which are known from Cenomanian (Cretaceous) seep deposits in Japan, 93–100 Ma [\(Kaim et al. 2008\)](#page-16-0). These estimates can be considered a minimum age of the split between the genera. As sequences for the nuclear DNA markers EF1a, ATPSa, and ATPSb could not be obtained for fossil samples, only ribosomal and mitochondrial markers plus the nuclear H3 locus were used for analysis (table 3). New sequences for all species of Alviniconcha from several localities were added to existing alignments from [Johnson et al. \(2010,](#page-16-0) [2015\)](#page-16-0) with default parameters using Muscle executed in Geneious Prime [\(http://www.geneious.com; last accessed July 24, 2020\)](http://www.geneious.com). Fragments of the 16S mtRNA and 12S mtRNA loci that had been previously excluded due to alignment ambiguities with more distantly related taxa were informative and therefore included within the genera Alviniconcha and Ifremeria. For the rest of the Abyssochrysidae, these regions are represented as missing data. For each locus and species, the most common allele was chosen as representative sequence for phylogenetic reconstructions. JModelTest v2 ([Darriba et al. 2012](#page-15-0)) was used to identify the best fitting evolutionary model for each gene and to partition the data for phylogenetic analyses. Concatenated gene trees were estimated based on a relaxed, lognormal clock with a Yule tree prior, a GTR $+$ I $+$ Γ substitution model as well as unlinked base frequencies and rates. The Markov chain Monte Carlo chains were run for 200 million generations, with parameters sampled every 5,000 generations. We used Tracer v1.7.1 [\(Rambaut et al. 2018\)](#page-17-0) to determine if effective sample sizes were adequate for all parameters and if analyses had reached convergence. Node dates were calibrated as a normal distribution around the timing of genera splits.

A StarBeast v2.5.7 [\(Bouckaert et al. 2014\)](#page-15-0) species tree was estimated including only the genera Alviniconcha and Ifremeria as replicate individuals for more distantly related taxa were missing. These estimated trees included at least one individual from each population per species or replicates from a single population. Species trees were calculated based on 1) the same loci used for the concatenated gene tree as well as 2) all available genetic markers (i.e., including the nuclear genes EF1a, ATPSa, and ATPSb). Phylogenetic reconstructions were performed with loci unlinked by partition, a $GTR + I + \Gamma$ site model with empirical frequencies, an uncorrelated lognormal clock, an analytical population size integration model, and a Yule tree prior. Markov chain Monte Carlo analyses were run for 200 million generations, with a tree sampled every 10,000 generations, a preburnin of 1,000 and 10 initialization steps. All analyses were run multiple times and results assessed in Tracer, FigTree v1.4.4 [\(http://tree.bio.](http://tree.bio.ed.ac.uk/software/figtree/) [ed.ac.uk/software/figtree/; last accessed July 24, 2020\)](http://tree.bio.ed.ac.uk/software/figtree/), and Densitree v2.2.1 ([Bouckaert 2010](#page-15-0)).

Phylogenetic Analyses of Snail Endosymbionts

New 16S rRNA endosymbiont sequences from the gills of A. adamantis and I. nautilei were compared against the NCBI nonredundant database. The new sequences were then aligned with Sina v1.2.11 ([Pruesse et al. 2012\)](#page-17-0) against the global Silva 16S rRNA alignment with available 16S rRNA endosymbiont sequences from GenBank for Alviniconcha species, I. nautilei, and other representative chemosynthetic taxa [\(Beinart et al. 2012](#page-15-0), [2015\)](#page-15-0). Phylogenetic trees for the Gammaproteobacteria and Campylobacteria symbionts were constructed with MrBayes v.3.2.7a [\(Altekar et al. 2004](#page-15-0)) under a GTR $+$ I + Γ substitution model in the Cipres Science Gateway v3.3 [\(Miller et al. 2010\)](#page-16-0). Two sets of three heated chains and one cold chain were run for 1,100,000 generations and sampled every 100 generations. The first 100,000 generations were discarded as burnin. Reference sequences of Nautilia lithotrophica (NR_025497) and Beggiatoa alba (NR_041726) were used as outgroups for the Campylobacteria and Gammaproteobacteria trees, respectively. Ancestral endosymbiosis traits were identified with Mesquite v.2.75 [\(Maddison WP and Maddison DR 2011\)](#page-16-0).

High-Throughput 16S V4–V5 Amplicon Sequencing and Analysis

The 16S V4–V5 hypervariable region was investigated to assess symbiont phylotypic variation in relation to host species and geographic location. After overnight lysis in proteinase K, DNA from RNALater (Thermo Fisher Scientific, Inc.) preserved gill samples was extracted with the Zymo Quick DNA 96 Plus kit (Zymo Research, Inc.) and further cleaned with the ZR-96 Clean-up kit (Zymo Research, Inc.) following the manufacturer's protocols. Barcoded amplicon libraries (515 samples and 62 negative controls) targeting a \sim 375bp fragment were prepared using the primer pairs 515F/ 926R [\(Walters et al. 2015\)](#page-17-0) and sequenced on an Illumina MiSeq platform with a 2×250 -bp protocol at the Argonne National Laboratory (Lemont, IL). Host species identity was determined based on morphology (Ifremeria) or molecular barcoding of the mitochondrial COI gene (Alviniconcha).

After quality-checking the demultiplexed reads with FastQC v0.11.5 [\(Andrews 2010;](#page-15-0) [http://www.bioinformatics.](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) [babraham.ac.uk/projects/fastqc/; last accessed July 24, 2020\)](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/), we clipped adapters with Trimmomatic v0.36 [\(Bolger et al.](#page-15-0) [2014\)](#page-15-0) and followed the Usearch v11 [\(Edgar 2010](#page-15-0)) pipeline for analysis. Reads were merged allowing lengths >300 bp, and then filtered applying a maximum error rate of 0.001, a quality threshold of 20, and a minimum length of 300 bp. The fastx_uniques and unoise3 commands were used to dereplicate, denoise, and cluster the reads into ZOTUs, whereas the otutab command was used to generate the OTU table. Taxonomy was assigned to each ZOTU in Qiime2 ([https://](https://qiime2.org) [qiime2.org; last accessed July 24, 2020](https://qiime2.org)) with a Naïve Bayes classifier that we trained against the target 16S V4–V5 region using the Silva 132 99% reference database. Ambiguous anno-tations were resolved using BlastN searches ([Altschul et al.](#page-15-0) [1990\)](#page-15-0). Recent studies have shown that well-to-well contamination accounts on average for 2.37% of reads in high biomass samples that were processed at the Argonne National Laboratory ([Minich et al. 2019\)](#page-16-0). Consequently, for further analysis, we removed ZOTUs that had <2.37% abundance in a sample, ignored all ZOTUs that were not annotated as vent snail endosymbionts and we excluded all samples with <1,000 symbiont reads. To identify any unrecovered phylotype variation, we analyzed the filtered symbiont read set with the Oligotyping v2.0 method [\(Eren et al. 2013\)](#page-15-0) based on 24 informative nucleotide positions. Only oligotypes with at least 2.37% abundance in a sample (–a), 100 supporting reads (–M), and occurrence in more than one individual (–s) were considered. We further excluded reads that had base quality scores $\langle 20$ in the nucleotide positions of interest $(-q)$.

Nonmetric multidimensional scaling and diversity analyses for the final symbiont oligotype data set were performed with the PhyloSeq package in R v3.5.2 ([McMurdie and Holmes](#page-16-0) [2013;](#page-16-0) [R Core Team 2018\)](#page-17-0) based on weighted Unifrac distances on log-transformed read counts. We also performed partial Mantel tests using the Vegan package [\(Oksanen et al.](#page-16-0) [2019\)](#page-16-0) to identify relationships between Alviniconcha host and symbiont genetic distances and geography based on Pearson's product–moment correlations. Pairwise host genetic distances were calculated for mitochondrial COI sequences based on Kimura's two-parameter model ([Kimura 1980](#page-16-0)). For geographic distances, we determined the geodesics between vent sites using the Geosphere package ([Hijmans 2019\)](#page-16-0).

Supplementary Material

Supplementary data are available at Molecular Biology and Evolution online.

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Author Contributions

S.B.J. designed the study with advice from R.C.V., conducted laboratory work, performed the fossil-calibrated phylogenetic analyses, and edited the manuscript. C.B. analyzed host population genetic structure, symbiont diversity, and phylogenetic relationships and wrote the paper. V.T., R.A.B., and R.C.V. contributed study specimens and manuscript edits. R.A.B. generated the 16S rRNA gene amplicon data set. D.A.C wrote sections on tectonics of the region and edited several drafts of the manuscript.

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