Genomic Blueprint of Glycine Betaine Metabolism in Coral Metaorganisms and Their Contribution to Reef Nitrogen Budgets

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HIGHLIGHTS

- Coral tissues contain high concentrations of the osmolyte glycine betaine
- Corals and their microbial symbionts can produce and degrade glycine betaine
- High gene expression patterns signifies role in coral-microbial symbiosis
- Glycine betaine is estimated to encompass 16% of the coral’s nitrogen biomass

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SUMMARY

The osmolyte glycine betaine (GB) ranks among the few widespread biomolecules in all three domains of life. In corals, tissue concentrations of GB are substantially higher than in the ambient seawater. However, the synthetic routes remain unresolved, questioning whether intracellular GB originates from de novo synthesis or heterotrophic input. Here we show that the genomic blueprint of coral metaorganisms encode the biosynthetic and degradation machinery for GB. Member organisms also adopted the prokaryotic high-affinity carrier-mediated uptake of exogenous GB, rendering coral reefs potential sinks of marine dissolved GB. The machinery metabolizing GB is highly expressed in the coral model Aiptasia and its microalgal symbionts, signifying GB’s role in the cnidarian-dinoflagellate symbiosys. We estimate that corals store between \(10^6–10^9\) grams of GB globally, representing about 16% of their nitrogen biomass. Our findings provide a framework for further mechanistic studies addressing GB’s role in coral biology and reef ecosystem nitrogen cycling.

INTRODUCTION

The quaternary ammonium compound glycine betaine (GB) represents one of the most ubiquitous osmolytes on Earth (Roesser and Müller, 2001; Welsh, 2000; Yancey et al., 1982). GB is widespread in the cells and tissues from organisms of all three domains of life (King, 1988; Roesser and Müller, 2001; Welsh, 2000; Yancey et al., 1982), where it generally serves as a compatible solute (osmolyte) to counteract high salinity effects and helps protect membranes and proteins against abiotic stressors (Burg and Ferraris, 2008; Chen and Murata, 2002). Although reliable estimates of dissolved GB in the ocean water are scarce, evidence suggests that dissolved concentrations of betaine analogs (e.g., proline betaine and alanine betaine) are in the picomolar range \([3–482 \text{ pM}; (Muslin, 2017)]\), whereas GB bound in seawater (cellular) particulates are in the nanomolar to lower micromolar range \([0.48 \text{ mM}; (Airs and Archer, 2010; Beale and Airs, 2016; Cree, 2015; Keller et al., 2004)]\). Paradoxically, the extensive knowledge of GB in marine systems stems from studies of cultured phytoplankton species, mostly microalgae/diatoms such as Emiliania huxleyi, Thalassiosira pseudonana, and Amphidinium carterae (Gebser and Pohnert, 2013; Keller et al., 1999a, 1999b; Spielmeyer et al., 2011; Spielmeyer and Pohnert, 2012). In the investigated phytoplankton cultures, maximum intracellular concentrations were higher under nitrogen-replete conditions \([3–172 \text{ mM}; (Keller et al., 1999b, 1999a)]\) than when nitrogen was limiting \([4–15 \text{ mM}; (Keller et al., 1999a)]\). GB concentrations are also sensitive to elevated levels of temperature, CO\(_2\), and salinity (Gebser and Pohnert, 2013; Spielmeyer and Pohnert, 2012) and can be strain-/species-specific and growth-stage dependent (Keller et al., 1999b, 1999a; Spielmeyer et al., 2011). Crucially, GB can comprise up to 20% of the cellular nitrogen pool in phytoplankton (Keller et al., 1999a). Once excreted or released from lysed cells, this organic compound represents a ubiquitous and dynamic constituent of oceanic dissolved organic matter readily taken up by marine heterotrophic bacteria (Kiene et al., 1998; Liburdy et al., 2015; Sun et al., 2011).

Quantitatively significant levels of GB also accumulate in the tissues of tridacnid clams, sponges, and scleractinian corals \([33–215 \text{ mmol kg}^{-1}\text{ wet tissue}]; \text{concentrations of up to 90-fold higher than the alternative sulfur-containing osmolyte dimethylsulfoniopropionate (DMSP; } (Hagedorn et al., 2010; Hill et al., 2017, 2010; Yancey et al., 2010)]\), in turn, suggesting a major role for GB in coral physiology. However, the biosynthesis of GB has not been studied in corals or marine invertebrates in general (see for example [Stephens-Camacho et al., 2015]), limiting our understanding of the synthetic routes for de novo production and...
sequestration of exogenous GB. Hypothetically, the higher intracellular concentrations of GB in reef corals (Hagedorn et al., 2010; Hill et al., 2017, 2010; Yancey et al., 2010), relative to the nanomolar concentrations reported in seawater (Airs and Archer, 2010; Beale and Airs, 2016), suggest an intrinsic capacity of corals to synthesize GB, the retention of extracellular GB obtained via heterotrophic input [e.g., feeding; (Houlbreque and Ferrier-Pages, 2009)], or both.

A two-enzyme oxidation pathway generally synthesizes GB from choline via the unstable intermediate betaine aldehyde in higher plants and bacteria (Figure 1A). The first oxidation step is catalyzed by an NAD⁺-dependent choline dehydrogenase (BetA) in prokaryotes and by a flavin-dependent choline monoxygenase (CMO) in plants, whereas the second reaction is catalyzed by an NAD⁺-dependent betaine aldehyde dehydrogenase (BetB) in all organisms (Chen and Murata, 2002). In some microorganisms, choline (and betaine aldehyde) is directly oxidized by choline oxidase (CodA) to GB with molecular oxygen as the electron acceptor (Lambou et al., 2013). GB can also be generated from glycine via a three-step methylation pathway involving glycine-sarcosine methyltransferase (GSMT) and sarcosine-dimethylglycine methyltransferase (SDMT) (Nyysölä et al., 2000), with S-adenosylmethionine serving as the methyl group donor (Figure 1A). However, in a few organisms, all three successive methylation steps are catalyzed by a single enzyme called glycine sarcosine dimethylglycine N-methyltransferase (GSDMT), essentially a polydonor (Figure 1A). However, only two genomes (Acanthaster planci and Sonneratia caseolaris) possessed a putative codA gene (Figure 1B). Subject to further experimental validation, these findings suggest that the complementary pathway oxidizing choline directly to GB is widespread in marine eukaryotes.

**RESULTS AND DISCUSSION**

**Eukaryotic Genomes Encode Multiple Synthetic Routes for GB Production**

Bioinformatic analyses revealed that the genomes (n = 114) and transcriptomes (n = 16; Table S1) of an overwhelming majority of ecologically important marine invertebrates encode the biosynthetic pathway for GB via choline oxidation (CodA, CMO, BetA, and BADH2), glycine methylation (GSMT/SDMT and GSDMT), or both (Figures 1A and 1B). Genomes of eukaryotes inhabiting terrestrial and freshwater ecosystems exhibited a similar trend (Table S2). Crucially, 85% of the analyzed eukaryotic genomes possessed a putative codA gene (Figure 1B). Putative enzymes of the alternative, dual-enzyme pathways for oxidizing choline to GB using choline monooxygenase (CMO) or choline dehydrogenase (BetA)—for the first reaction—and betaine aldehyde dehydrogenases (BetB or BADH2)—for the second reaction were also present in the genomes/transcriptomes (Figure 1B; Table S2). Potential CMO, betA, and BADH2 homologs with 40%–92% identity to validated proteins (Figure S1) were present in 13%, 23%, and 68% of the studied eukaryotes, respectively (Figure 1B). However, only two genomes (Acanthaster planci and Sonneratia caseolaris) possessed a putative betB homolog (Table S2). In-depth phylogenetic analyses showed that 88% of the predicted BADH2 genes were homologous (>50% amino acid identity; Figure S1) to functionally characterized animal BADHs within aldehyde dehydrogenase (ALDH) family 9 (Julián-Sánchez et al., 2007), whereas the remainder were affiliated to validated plant BADH2s (Figure S2) within the ALDH family 10 proteins (Kirch et al., 2004). To determine the occurrence analysis of predicted CMO, betA, and BADH2 homologs in individual genomes/transcriptomes further showed that 11% and 43% of the analyzed eukaryotes possessed the complete CMO/BADH2- and BetA/BADH2-catalyzed pathways, respectively. However, the complete CMO/BADH2 pathway was predicted absent in Symbiodiniaceae and cnidarian genomes/transcriptomes (Figure 1B; Table S2). Notably, the putative “CMO”-like genes in both groups of organisms showed greater homology to carnitine monooxygenases (30%–40% identity) involved in carnitine metabolism than to ratified eukaryotic CMO proteins (≤25% identity). Crucially, these results indicate the possibility to oxidize choline to GB using the BetA/BADH2-catalyzed pathway in five out of eight cnidarians, which is absent in all Symbiodiniaceae genomes/transcriptomes analyzed (Figure 1B).

Glycine methylation pathways (Figure 1A) were generally underrepresented in the studied genomes/transcriptomes (Figure 1B); the complete GSMT/SDMT- and GSDMT-catalyzed pathways occurred in 18% and
Figure 1. Incidence of Glycine Betaine (GB) Metabolic Pathways in Eukaryotes

(A) Schematic view of pathways for the uptake, biosynthesis (via oxidation and methylation), and catabolism of GB.

(B) Incidence of key genes in eukaryotic genomes (n = 114) and transcriptomes (n = 16) in the upper panel and diverse marine invertebrate lineages (and their close relatives) harboring BCCT carriers (lower panel). Data from transcriptomes are designated with ‘#’. Additional information is provided in Tables S1 and S2.

(C) Copy numbers of predicted genes in each genome (excluding transcriptomes) and the taxonomic distribution of abundant metabolic pathways based on the presence-absence of key genes in the sampled genomes and transcriptomes. Statistically significantly different average copy numbers per genome are denoted by different letters as deduced by one-way ANOVA tests (p < 0.001) with two-stage step-up Benjamin-Krieger-Yekutieli’s multiple comparison test.

Enzymes: ChT, choline transporter; BCCT, betaine-choline-carnitine transporters; BetA, choline dehydrogenase; CMO, choline monooxygenase; BetB/BADH2, betaine aldehyde dehydrogenase; CodA, choline oxidase; GSDMT, glycine-sarcosine methyltransferase; SDMT, sarcosine dimethyltransferase; GSDMT, glycine sarcosine dimethylglycine N-methyltransferase; BHMT, betaine-homocysteine methyltransferase; DMGDH, dimethylglycine dehydrogenase; SDH, eukaryotic sarcosine dehydrogenase; SoxA2, monomeric sarcosine oxidase; SoxABDG, heterotetrameric sarcosine oxidase.

Abbreviations: Na+, sodium ion; H+, proton; NAD+ and NADH, oxidized and reduced nicotinamide adenine dinucleotide; TH4F, tetrahydrofolate; FTH4F, formyl-tetrahydrofolate; MP, Myxozoa and Polyplodiozoa; CPC, Choanoflagellates, Placozoa and Ctenophora; CHP, Cryptophytes, Haptophytes, and Plantae (Algae); MRB, Mollusca, Rotifera, and Brachiopoda; HCCETU, Hemichordates, Cyclosteomes, Cephalochordates, Echinoderms, Tunicates, and Urochordates. Species indicated as “Others” are listed in Table S2.
~8% of the genomes respectively. Interestingly, the GSMT/SDMT-catalyzed pathway was also predicted in five coral genomes but found lacking in Symbiodiniaceae (Figure 1B). Although further biochemical evidence is warranted to establish activities of all these pathways, the possession of the GSMT gene putatively encoding glycine-sarcosine methyltransferase by cnidarians, a demosponge (Amphimedon queenslandica), and many of the investigated mammalian genomes (Figure 1B; Table S2), suggests the potential to turnover glycine to dimethylglycine (Lai et al., 2006; Nyssöla et al., 2001), which also counteracts osmotic and thermal stresses (Bashir et al., 2014). In summary, corals and their dinoflagellate endosymbionts alongside other marine invertebrates are deduced to possess at least one pathway for producing GB de novo, which points to the significance of GB in coral biology and underscores reef-building corals as a potential source of GB in the ocean.

The Ability to Scavenge Exogenous GB Is Taxonomically Constrained but Present in the Coral Host, Microalgae, and Bacteria of Coral Metaorganisms

Intriguingly, genomic profiling of transporters involved in GB metabolism demonstrates the ability to scavenge extracellular GB in higher organisms is constrained to very few eukaryotic lineages, namely cnidarians, ctenophore, diatoms, dinoflagellates, ciliates, and cryptophytes (Figure 1B). The vast majority of eukaryotic genomes/transcriptomes investigated here encode the putative sodium-dependent high-affinity choline carrier ChT (30%–50% identity to validated proteins; Figure S1), which can take up choline for GB biosynthesis (Figures 1B and 1C; Table S2). However, only 25.4% of the eukaryotes harbored a potential GB uptake system (Figures 1B and 1C). Crucially, the eukaryotes that do were predicted to possess numerous copies of genes putatively encoding the betaine-choline-carnitine-transporter (BCCT) family (30%–40% identity to validated carriers; Figure S1). BCCT carriers are sodium- or proton-coupled transporters that are well characterized exclusively in prokaryotes, where they canonicly mediate high-affinity uptake of betaines, choline, or carnitine (Ziegler et al., 2010). Homologs of BCCT carriers were also found in the genomes/transcriptomes of diverse Symbiodiniaceae species (Figures 1B and 2A), the intracellular dinoflagellate symbionts of cnidarians, suggesting the potential capacity to accumulate betaine, choline, or carnitine in hospite. Of evolutionary significance is the fact that BCCT homologs were found only in a few animals hosting microalgal endosymbionts (LaJeunesse et al., 2018) or close photosynthetic relatives, suggesting an important role in the cnidarian-dinoflagellate symbiosis.

It is noteworthy that the alternative ABC-type transport systems commonly found in prokaryotes for GB (OpuA and OpuC) or choline (OpuB, ProU, and ChoABC) (Ziegler et al., 2010) are lacking in the eukaryotes studied here. Also, only the diatom Thalassiosira pseudonana, the cryptomonad alga Guillardia theta and the coccolithophore Emiliana huxleyi encoded putative homologs of the choline and GB exporter EmrE (Tables S3 and S4), which exports naturally occurring quaternary cation compounds (Bay and Turner, 2012). Taken together, the systematic evaluation of potential GB uptake systems across genomes/trancriptomes of model and non-model metazoans depicts a confined distribution of BCCT and EmrE carriers to globally important marine invertebrates such as cnidarians and microalgae.

Cnidarians host a diverse suite of prokaryotes (Rohwer et al., 2002). Therefore, we sought to explore whether their bacterial associates also contribute to GB metabolism. Focusing on the ubiquitous Endozoicomonas species (n = 13), encompassing bacteria that seem to commonly colonize tissues of diverse cnidarians (Neave et al., 2016, 2017a, 2017b), our data show that not all species of these “obligate” endosymbionts have the putative machinery to synthesize and export GB though all possessed the ability to scavenge GB (Figure S3). These findings are also consistent with the incidence of profiled synthetic and uptake routes for GB across S2 metagenome-assembled prokaryotic genomes [MAGs; (Robbins et al., 2019)], representing the most abundant bacterial and archaeal taxa in the environmentally resilient reef-building coral Porites lutea (Table S5).

The majority of the genomes queried (56 out of 64) encoded BHMT proteins that can potentially convert GB to dimethylglycine, but most—all Endozoicomonas species and one-third of MAGs from P. lutea—lack putative enzymes for catabolizing dimethylglycine to sarcosine (Figure S3; Table S5). Thus, we speculate that hosting bacteria with reduced degradative capacity for GB maintains a high intracellular concentration of its precursor dimethylglycine—also an osmolyte, in coral tissues. Considering the ubiquity of Endozoicomonas (and other microbes) in coral microbiomes (Neave et al., 2016, 2017a, 2017b), we infer that the production of osmotically active compounds might be a cooperative mechanism supporting coral holobiont physiology (Ochsenkühn et al., 2017). Although speculative at this point, we propose that the coral host and its associated microbes—that is, Symbiodiniaceae and some bacteria—produce GB from choline or
glycine and facilitate the exchange of GB scavenged from prey or the ambient seawater (by the host) or generated in situ (by the entire metaorganism) within the holobiont. This hypothesized model, however, requires experimental validation.

Figure 2. Evolutionary History of Predicted Eukaryotic BCCT Carriers

(A) An unrooted maximum-likelihood phylogenetic tree showing the affiliation of predicted coral BCCT proteins (red circles) next to their closest cnidarian relatives (blue circles) and the basal prokaryotic BCCT carriers (wedged gray symbol). Colored circles highlight taxonomic affiliation of eukaryotic BCCT proteins. Note that the coral BCCTs are divided into four clusters (≥80% bootstrap support), with some putatively located in the mitochondria.

(B) The average density of predicted BCCT genes in marine invertebrates and marine prokaryotes with fully sequenced genomes (n = 125; Table S6); difference in copy number per mega-base pair (Mbp) of coding genome was only significant (one-way ANOVA, p = 0.0012) for Prokaryotes (gray circles) relative to Alveolata (green circles).

(C) Multiple aminoacid sequence alignment showing the G-x-G-x-G motif (highlighted in green) found in validated prokaryotic BCCT carriers specific for GB (BetP, ButA, OpuD, BetU, and BetL), choline (BetT), and carnitine (CaiT). Glycine residues lacking in choline and carnitine carriers are colored in red and blue (and yellow) respectively, whereas those missing in Symbiodiniaceae proteins are shown in gray. The amino acid positions reflect those in BetP from C. glutamicum. For brevity, only representative eukaryotic proteins are shown; Figure S6 provides the complete data.
Eukaryotic BCCT Homologs Are Distant Relatives of Prokaryotic GB Carriers

The narrow distribution of putative BCCT genes in diverse eukaryotes studied here (Figure 1B) warrants elucidating the evolutionary history of the deduced transport system. Phylogenetic analyses indicate that the predicted eukaryotic BCCT carriers are distantly related to bona fide BCCT transporters from prokaryotes (Figure 2A). Moreover, these carriers group into discrete sequence clusters that reflect the metazoan evolutionary history (Derelle et al., 2015). Cnidarian sequences also form multiple protein clusters (A to D; Figure 2A), which are interspersed by proteins containing signal peptides or mitochondrial-like sequence motifs (Figure 2A), suggesting multiple subcellular localization of the carrier in coral tissues. Collectively, these results imply that the acquisition of BCCT carriers likely occurred independently sensu stricto before the diversification of metazoans.

Further analyses comparing copy numbers of predicted BCCT genes across different species indicate that the average copies (± SD) scaled by the coding genome size (in mega base pairs, Mbp), were only significantly different (one-way ANOVA, p = 0.0012; Figure 2B) between marine prokaryotes (0.59 ± 0.48 copies Mbp⁻¹; n = 125; Table S6) and alveolates (0.03–0.11 copies Mbp⁻¹; n = 8). The genomic density of predicted BCCT genes is also similar among coral species with different life histories (~0.5 ± 0.2 copies Mbp⁻¹; n = 4–5 replicates per species) as deduced from metagenomic coding sequences of ten Red Sea scleractinian coral species (Figure S4; Table S7). However, the clustering of similar sequences at 95% nucleotide identity over 80% of the length of the shorter sequence reveals specificity of predicted BCCT copies to individual coral species (Figure S5). Taken together, this suggests that BCCT copy number and gene isoforms are highly conserved in cnidarians but evolved species specifically.

Coral BCCT Homologs Likely Transport GB as Their Cargo

Additional bioinformatic analyses strongly support the annotation of the predicted eukaryotic BCCT genes as GB transporters. Multiple sequence alignments of the predicted amino acid sequences and functionally characterized BCCT proteins with varied substrate specificity (Figures 2C, S6, and S7) revealed conservation of key active site glycine residues (G-x-G-x-G; Figures 2C and S6) contributing to sodium and substrate binding, as well as tryptophan and tyrosine residues (Figure S7) that catalyze GB binding in sodium-coupled betaine symporters (Perez et al., 2011; Ressl et al., 2009). Interestingly, the central glycine residue in the case of Symbiodiniaceae and some other algal species is substituted by alanine (G-x-A-x-G; Figures 2C and S6), in a region that renders the substrate-binding site accessible (Perez et al., 2011). Although further experimental validation is required, we hypothesize altered kinetics and reduced affinity for GB in Symbiodiniaceae BCCT carriers relative to host BCCT carriers, based on point mutation studies of the GB carrier of Corynebacterium glutamicum (Perez et al., 2011). It is noteworthy that betaine carriers can also transport small quantities of structurally related organic solutes such as proline betaine, dimethylsulfoxoniopropionate (DMSP), and dimethylglycine (Tandon et al., 2020; Ziegler et al., 2010) and that transporters are also fully reversible (Fanwick et al., 1995).

High Expression of GB Metabolism in Aiptasia and Pelagic Symbiodiniaceae

To gain further insight on the relevance of the predicted synthetic and transport pathways for GB in coral physiology, we analyzed previously published transcriptomic data from the coral model system Aiptasia, encompassing different developmental and symbiotic states (Baumgarten et al., 2015). Gene expression analyses indicated that the pathways involved in the metabolism of GB were highly expressed in both symbiotic and aposymbiotic states (i.e., with and without microalgal symbionts) across larval and adult stages (Figure 3). For instance, the choline transporter ChT was highly expressed in aposymbiotic larvae, whereas the BCCT carrier was expressed similarly in larvae and adult stages. Genes putatively involved in the production of GB from choline oxidation (betaA/BADH2 and codA) were significantly highly expressed (one way ANOVA, p > 0.001) in adult anemones, especially in symbiotic colonies (e.g., betaA and codA). In contrast, the glycine methylation pathway catalyzed by GSMT and SDMT was differentially highly expressed in symbiotic and aposymbiotic anemones, respectively (Figure 3). Crucially, transcription was in both cases, four-fold higher than choline oxidation pathways catalyzed by BetA/BADH2 and CodA. The degradative pathway catalyzed by BHMT, DMGDH, SDH, and SHMT, however, followed a trend of higher expression in symbiotic adult anemones (Figure 3), suggesting a role of Symbiodiniaceae in GB catabolism. Our results therefore suggest a greater contribution of GSMT-/SDMT-catalyzed pathway to GB production and the turnover of intracellular GB by Symbiodiniaceae.

We next evaluated whether free-living populations of the microalgal symbionts actively transcribed the metabolic pathway for GB in the global ocean by interrogating a catalog of 116 million expressed...
Figure 3. Transcriptional Activity of GB Metabolism in the Coral Model Aiptasia

Gene expression was determined from published whole tissue transcriptomes (Baumgarten et al., 2015) of aquacultured Aiptasia larvae and adults fed regularly on brine shrimps (n = 2–4 replicates per experiment). Transcriptomes encompass various developmental and symbiotic states. Animals without dinoflagellate endosymbionts are indicated with “−”, whereas adults with intermediate or full endosymbiont levels are denoted with “+” and “++”, respectively. Barplots indicate average (± SD) expression in replicated samples and all predicted gene homologs (denoted by circular symbols). Statistically significantly different expression levels are denoted by different letters as deduced by one-way ANOVA tests (p > 0.05) with two-stage step-up Benjamin-Krieger-Yekutieli’s multiple comparison test. ND, average expression was below ~1 FPKM.
eukaryotic genes of the Tara Oceans Expedition (Carradec et al., 2018) available at http://www.genoscope.cns.fr/tara/. We retrieved several unigenes with significant homology to Symbiodiniaceae genes (≥80% identity and alignment length of ≥60%) involved in GB metabolism, including BCCT (33), codA (1), DMGDH (3), SDMT (1), SHMT (6), and BHMT (31). The genes encoding for BCCT uptake and exchange system for GB were the most highly expressed among the six examined genes (Figure 4). Expression was highest in the mesoplankton size fraction (180–2000 μm)—suggested to correspond to Symbiodiniaceae in symbiosis with their host organisms (Decelle et al., 2018), especially in subsurface samples (Figure 4). In contrast, the codA gene encoding the one-step enzymatic route of GB production was highly expressed in the smaller (0.8–20 μm) size fractions (i.e., free-living Symbiodiniaceae). The expression levels of transport (BCCT) and synthesis (CodA) machineries suggest differences in the metabolism of GB between free-living and symbiotic lifestyles, with higher GB uptake potential inferred in symbiotic Symbiodiniaceae although experimental validation is needed.

A Putative Role for GB in Stress Resilience and Nitrogen Storage of Coral Metaorganisms

The "omics" evidence presented here strongly supports the metabolism of GB as a genetic facet of coral physiology and symbiotic interactions, especially given our estimates that GB comprises about 16% of the total nitrogen biomass of corals [based on the predicted biomass of corals of about 30 g C m⁻² (Crossland et al., 1991), a median GB concentration in coral biomass of 6 μmol cm⁻² (Hill et al., 2010), and a C:N ratio of 6.5 in coral tissues]. Despite its osmolytic properties, GB may have additional properties that supersede other osmo-lytes, such as DMSP. For instance, in plants GB inhibits thermal and photon stress by stabilizing extrinsic proteins vital for the photosynthetic and oxygen-evolving complex of photosystem II (Chen and Murata, 2002). A similar role is hypothesized in microalgae and corals (Hill et al., 2010), because corals located at shallower depths and locations exposed to high irradiance possess elevated pools of GB relative to colonies living in deeper waters and shaded areas (Hill et al., 2010). Moreover, although corals are osmoconformers (i.e., in osmotic equilibrium with their environment), their partnership with Symbiodiniaceae that formed at least 160 million years ago (LaJeunesse et al., 2018) probably generated osmotic constraints, whereby the coral host had to devise a suitable intracellular environment for its endosymbionts (Ochsenkühn et al., 2017).

In addition, we postulate that the thermodynamic constraints imposed by salinity (Oren, 1999) coupled with the inherent biochemical properties of GB rendered this osmolyte particularly suitable for a number of
Corals and Reef Ecosystems Are a Sink for Dissolved GB in the Sea

It is worthwhile to note that the uptake of GB by the high-affinity membrane carriers deduced here might be effective at the ambient dissolved (picomolar range) concentrations reported in the ocean (Muslin, 2017), especially given the high affinity for GB ($K_a$ of up to 8.6 µM in prokaryotic BCCT carriers (Farwick et al., 1995; Peter et al., 1996). However, the ingestion of plankton (Houlbreque and Ferrier-Pages, 2009) most likely comprises the main source of GB in coral tissues since heterotrophic intake (i.e., feeding) can deliver three to four times more nitrogen than dissolved sources in model coral species (Houlbreque and Ferrier-Pages, 2009). The fact that intracellular concentrations of GB make up a significant fraction (7%–23%) of the total organic N in marine phytoplankton (Keller et al., 1999a; King, 1988) implies that incidental GB acquired from feeding might be important in the nitrogen economy of coral reefs. Following this notion, we reasoned that, if heterotrophic input via feeding is vital for a coral’s nutritional ecology, then GB pool sizes in corals have to be placed in the context of their planktonic prey, which likely includes GB producers.

Therefore, as a first step, we profiled the biosynthetic pathways of GB in 189 completely sequenced microbial genomes (mean (±SD) completeness of 99 ± 1%) retrieved from the marine microbial reference (Mar-Ref) genome database (Klemetsen et al., 2018) encompassing diverse species from various oceanic provinces (Table S8). The inventory of genes encoding putative synthetic enzymes for GB showed a high incidence of choline-oxidizing pathways (in 6%–55% of the genomes) and low occurrence of glycine-methylation-based pathways (up to ~17%; Figure S8A), possibly reflecting high energetic costs of methylation reactions, particularly the production of the methyl donor/acceptor S-adenosylmethionine (Luka et al., 2009; Nyysölä et al., 2000). The co-occurrence of choline-oxidizing enzyme complements in individual genomes (Figure S8B) further revealed the predominance of the BetA/BADH2 and BetA/BetB-catalyzed pathways (in ~36% and 13%, respectively) relative to the pathway involving choline monooxygenase (CMO). Also, a low-average copy number per genome (Figure S8C) was observed for betA (1 copy; $n = 71$ genomes), betB (1 copy; $n = 26$), and BADH2 (1 copy; $n = 72$) in comparison to codA (2.3 copies; $n = 104$), where the high coefficient of variation of 67% (versus 11% in betA) around the mean suggested significant species-dependent copy variations. Considering the high incidence of co-occurrence of single-copy genes for the BetA/BADH2-catalyzed pathway in reference marine microbial genomes (Figure S8B; Table S8) implies that betA and BADH2 homologs are feasible proxies for estimating the relative abundance of GB producers from environmental metagenomes, because they have a greater likelihood of co-occurring in individual genomes. It is also noteworthy that two-thirds of the genomes with codA (55% of the genomes) also possessed the complete BetA/BADH2 pathway.

On this basis, we subsequently probed the abundance of betA and BADH2 protein-coding genes by interrogating global ocean metagenomic sequence datasets and normalized counts of retrieved genes using the mean abundance of four conserved single-copy housekeeping genes present in microbial genomes (Table S9). The two respective genes were predicted to be present in 2.4% and 1.2% of prokaryotes sampled in the Tara Oceans marine metagenomes (Sunagawa et al., 2015), which encompass picoplankton-sized microorganisms (0.22–3 µm) from the photic and epipelagic layers (Table S9). The mean of roughly 2% suggests that the environmental abundance of picoplankton producing GB is twice that of...
DMSP-producing microbes (Curson et al., 2017) in the pelagic ocean; however, the value could be higher, if we consider picoplankton encoding the one-step CodA-catalyzed pathway.

Based on the above estimates [i.e., a 2% proportion of GB-producing picoplankton, a cellular concentration of GB between 0.9 and 483 nM in seawater particulates >0.2 μm (Beale and Airs, 2016; Cree, 2015; Keller et al., 2004), ~10^6 cells per ml, and a total biomass of microbial picoplankton in the pelagic ocean of 3.6 × 10^18 cells (Whitman et al., 1998)], we project a global GB stock in pelagic marine picoplankton in the range of ~1 to 410 × 10^12 grams. Scaling these estimates to the areal cover of reefs of ~0.1% of the world’s ocean surface (Spalding et al., 2001), and assuming the global average water depth in areas occupied by reefs of 20 m, suggests that between 0.1 to 41 × 10^9 grams of GB is potentially distributed within picoplanktonic particulates in the waters overlaying corals.

These estimates are similar to the predicted global stock of GB in corals of about 18 × 10^6 to 4.64 × 10^9 grams, based on the GB concentration in coral tissues of 0.05–13.2 μmol cm^-2 (Hill et al., 2010), the molecular weight of GB (117.15 g mol^-1), and a mean areal coral reef cover of 0.3 × 10^6 km^2 (Spalding et al., 2001). However, the predicted global stock of GB present in corals may be significantly underestimated considering that much of the GB is likely catabolized by corals and their symbionts (e.g., via dimethylglycine to serine) or translocated through coral mucus ropes into the adjacent sediments and pore waters. Thus, at the global scale, coral reef ecosystems and their reef-building corals represent a rare, albeit significant and dynamic “hot spot” of microbial-derived GB in the pelagic ocean.

Conclusions
The systematic bioinformatic framework applied here combined with existing biochemical data strongly suggests a role of marine invertebrates in the biogeochemical cycling of GB in the ocean, adding corals to the roster of sinks and organisms producing GB. Because GB is a major reservoir of nitrogen and considering the general scarcity of nitrogen in the pelagic ocean (Bristow et al., 2017), we propose that GB serves both as an osmolyte and as a potent nitrogen reserve in marine environments hosting coral reefs. Moreover, given its photoprotective role in plants including alleviating oxidative stress (Chen and Murata, 2008), we speculate further that GB serves thermo- and photoprotective roles in the coral holobiont. This in turn implicates GB in the integrity of the coral-microalgal symbiosis and the resilience of coral reef ecosystems, further reiterating the notion that coral reefs are biological hotspots, acting as sinks for dissolved organic nitrogen in oligotrophic seas. Our findings open a path for a number of testable hypotheses requiring in-depth biochemical and experimental analysis for (1) the many uncharacterized genes and pathways we elucidated and postulated to play a role in regulating GB uptake, storage, and cycling, as well as (2) the hypothesized protective and nutritional roles of GB in the metabolism of coral holobionts.

Limitations of the Study
In the current study, we examined the genetic potential of corals to metabolize GB, which is present in their tissues at high concentrations. Most of the bioinformatic analyses are based on genomic/transcriptomic data, questioning whether the inference for GB production based on homologs is sufficient in the absence of biochemical validation of the deduced enzymes. A second limiting factor in robust interpretation of our results relates to the localization of the enzyme activities within the coral holobiont and the concentrations of GB in the studied animals species. These limitations call for ratification of the pathways that we deduced to function in GB metabolism in corals and their microbial symbionts and elucidating the contribution of coral holobiont partners in shaping the dynamics of intracellular GB pools.

Resource Availability
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Materials Availability
All data needed to evaluate the results and conclusions are presented in the main text and/or are available in the Supplementary Materials. Additional data or scripts related to this paper are available from the corresponding authors.
Data and Code Availability

The Raw sequence data for the coral metagenomes have been deposited in NCBI under BioProject number PRJNA437202 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA437202).

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101120.

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AUTHOR CONTRIBUTIONS

D.K.N. conceived and designed the study and performed the bioinformatic analyses. M.Z. and C.R.V. determined and provided the metagenomic data. C.M.D. provided the bioinformatic resources for the analyses. D.K.N. wrote the manuscript with contributions from all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental Information

Genomic Blueprint of Glycine Betaine Metabolism in Coral Metaorganisms and Their Contribution to Reef Nitrogen Budgets

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Supplemental Information

Genomic blueprint of glycine betaine metabolism in coral metaorganisms and their contribution to reef nitrogen budgets
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Figure S1. Related to Figure 1. Percentage identity of retrieved gene homologs to validated protein sequences from UniProt. The black dot and horizontal lines show the mean (± SD) identities. The dotted red and blue vertical lines denote identity cutoffs used for interrogating enzymes with no HMM models CMO, codA, SDMT, and GSMT (30%) and BADH2 and SDH (50%) respectively. Identities for genes with HMM models (chT, BCCT, beta, BHMT, DMGDH, and SHMT) are based upon blast search of selected hmmsearch hits for each gene (see methods for details) and subsequent search of the homologs against validated proteins from UniProt.
**Figure S2.** Related to Figure 1. Phylogenetic analysis of betaine aldehyde dehydrogenase-like genes (BADH2-like) predicted in eukaryotic genomes studied here. The tree represents a maximum-likelihood tree, with all nodes having bootstrap values above 50% (out of 1,000 replicates) shown on the tree as proportions. The family designation of the different aldehyde dehydrogenases (ALDHs) follows the work of (Kirch et al., 2004) and are color-highlighted as follows: family 12 (the outgroup); family 6 (methylmalonate semialdehyde dehydrogenase), family 7 (antiquitin-related ALDHs); family 21 and 22 (characterized novel ALDHs); family 11 (non-phosphorylating glyceraldehyde-3-phosphate dehydrogenase); family 5 (succinic semialdehyde dehydrogenase); family 2 and 3 (ALDHs with variable substrates); family 10 (BADH2). The values in parenthesis are the accession numbers, while the values after “#” denote the number of redundant genes (70% global identity over 60% of the protein) belonging to that sequence in the genome(s). The bar denotes 10% amino acid substitutions.
Figure S3. Related to Figure 1. Incidence of metabolic pathways for GB in twelve *Endozoicomonas* species associated with diverse marine invertebrates. The phylogenetic tree shows the relationship based on 20 bacterial single-copy genes inferred using the maximum-likelihood approach. The tree was rooted using *Pseudomonas aeruginosa* PA01 (Accession number NC_002516).
Figure S4. Related to Figure 2. Density of predicted BCCT homologs in diverse coral species based on metagenomics sequence data from the Red Sea. The maximum-likelihood tree on the left panel (a) is based on cytochrome oxidase (Cox2) gene present in representative assemblies of each species. (b) Bars show the average (± SD) counts of coding genes from metagenomes of each species, with values shown in brackets indicating the number of replicates per species. (c) Depicts the overall taxonomic breakdown of the coding genes present in replicates of each species (n = 4–5). (d) Shows the deduced copy number (per mega base pairs) of BCCT homologs present in the different coral species. Statistically significantly different average copy numbers per mega base pairs (Mbp) of coding genome are denoted by different letters as deduced by one-way ANOVA tests (P < 0.001) with two-stage step-up Benjamin-Krieger-Yekutieli’s multiple comparison tests. The dotted grey line shows the overall average copy number per Mbp across all metagenomes.
**Figure S5.** Related to Figure 2. Sequence diversity of BCCT genes predicted in Red Sea corals. The heatmap shows grouping of coral species (n = 4–5 replicates) by BCCT gene clusters. Similarity is defined at 95% global nucleotide identity over 80% of the shorter sequence length. Species and clusters are grouped based on a binary distance matrix and the average-linkage clustering approach. Clusters exclude singletons and sequences homologous to Symbiodiniaceae BCCTs.
Figure S6. Related to Figure 2. Multiple sequence alignments of validated bacterial BCCT carriers and predicted eukaryotic homologs; the third transmembrane helix (TM3) region of the carriers is highlighted. Note the significant homology of predicted eukaryotic BCCT carriers in the region shown by arrows containing the conserved G-x-G-x-G motif present in biochemically defined GB carriers (e.g., BetP from *C. glutamicum*). Amino acid positions are based on BetP.
|   | Corynebacterium glutamicum (CAA63771, BetP) |   |
|---|-----------------------------------------|---|
| 2. | Tetragenococcus halophilus (AAP06751, ButA) |   |
| 3. | Bacillus subtilis (AAC44368, OpuD) |   |
| 4. | Escherichia coli (AAQ10261, BetU) |   |
| 5. | Sinorhizobium meliloti (AA337253, BetS) |   |
| 6. | Listeria monocytogenes (AAD30286, BetL) |   |
| 7. | Escherichia coli (CAQ30790, BetT) |   |
| 8. | Escherichia coli (CAQ30560, CaI) |   |
| 9. | Proteus mirabilis (WP_012368453, CaI) |   |
| 10. | Acropora digitifera (XP_015778301) |   |
| 11. | Stylophora pistillata (Spis13436) |   |
| 12. | Orbicella faveolata (XP_020624688) |   |
| 13. | Discomonas sp. (scaffold_224_10) |   |
| 14. | Amplexidiscus fenestratus (scaffold_164_3) |   |
| 15. | Hydra vulgaris (XP_012562913) |   |
| 16. | Nematostella vectensis (XP_001640076) |   |
| 17. | Aiptasia pallida (AIPGENE15022) |   |
| 18. | Thalassiosira oceanica (EJK50030) |   |
| 19. | Fragilariopsis cylindrus (OEU16937) |   |
| 20. | Ostreococcus tauri (CEF96976) |   |
| 21. | Bathycoccus prasinus (XP_007508379) |   |
| 22. | Mnemiopsis leidyi (ML005533a) |   |
| 23. | Phaeodactylus tricornutus (XP_001827711) |   |
| 24. | Aureococcus anophagefferens (XP_009039779) |   |
| 25. | Symbiodinium minutum (Sminut010127) |   |
| 26. | Symbiodinium kawaguti (Skav202732) |   |
| 27. | Symbiodinium microadriaticum (Smic15475) |   |
| 28. | Linulodinium polyedrum (TRT_00517247) |   |
| 29. | Guillardia theta (XP_005827593) |   |
| 30. | Micromonas commoda (XP_002502671) |   |

**Figure S7.** Related to Figure 2. Multiple sequence alignments of validated bacterial BCCT carriers and predicted eukaryotic BCCT homologs illustrating residues involved in GB binding (orange and green boxes) based on (Ressl et al., 2009).
Figure S8. Related to Figure 2. The density and co-occurrence of synthetic routes for GB in completely sequenced marine prokaryotic genomes (n = 189). (a) Shows the overall number genomes possessing the retrieved gene homolog. (b) Co-occurrence incidence of genes involved choline oxidation pathways (in white) and GB formation via glycine methylation (in grey). (c) Indicates the average (± SD) copy number of predicted homologs per genome. Details of the genomes are provided in Table S8. Values beside each bar graph denote number of genomes (in a) or the proportion of genomes with the pathway (in b) out of 189 genomes; in (c) values indicate the average (± SD) copies across the genomes.
Transparent Methods

Genomic and transcriptomic sequence datasets

Information on GB metabolism is largely restricted to prokaryotes and plants. Hence, our genomic and transcriptomic survey strategy was tailored at capturing sequenced model and non-model organisms, including evolutionary unrelated higher organisms, such as those thriving in terrestrial habitats and marine-freshwater transition zones (e.g., mangrove plants and animals). Eukaryotic genomes \( (n = 114) \) and transcriptomes \( (n = 16) \) representing different metazoan lineages were retrieved from the NCBI/GenBank database or several dedicated databases (downloaded on March 2016; Table S1). Cnidarian and endosymbiotic Symbiodiniaceae sequences were retrieved from the reefgenomics database [http://reefgenomics.org/](http://reefgenomics.org/) (Liew et al., 2016).

Additional reference Symbiodiniaceae transcriptomes \( (n = 8) \) from the Marine Microbial Eukaryote Transcriptome Sequencing Project [MMETSP; (Keeling et al., 2014)] were retrieved from the iMicrobe data server (http://datacommons.cyverse.org/) under project #104. Transcriptomic data for other dinoflagellates (Karenia brevis, Lingulodinium polyedrum, and Prorocentrum minimum) and other marine invertebrates of interest lacking genomes such as sponges (Ephydatia muelleri, Haliclona species, Stylissa carteri, and Xestospongia testudinaria) were retrieved from the comparative genomic platform for early branching metazoans (http://www.compagen.org/) as well as the iMicrobe data server. Three recently published mangrove plant genomes (Avicennia marina, Sonneratia alba, and Sonneratia caseolaris) were also included to cover plants in the marine realm [http://evolution.sysu.edu.cn/Sequences/](http://evolution.sysu.edu.cn/Sequences/) (Xu et al., 2017). Additional genomes such as fungus-cultivating termite (Macrotermes natalensis), the Japanese abalone (Haliotis discus), the estuarine crocodile (Crocodylus porosus), and the Chinese mitten crab (Eriocheir sinensis) were retrieved from the GigaScience Database (http://gigadb.org/site/index).

We also retrieved 189 completely sequenced reference prokaryotic genomes (Table S6) from the marine microbial reference (MarRef) genome database (Klemetsen et al., 2018). These genomes were filtered from a total of 839 genomes (Accessed on June 07th 2019) by selecting the analysis project type “whole genome sequencing” and environmental material “sea water” in the online server and quality assessment with CheckM v1.0.3 (Parks et al., 2015), retaining only “high-quality” genomes (≥90% completeness and <5% contamination) fulfilling the MIMAG standards for single-cell and metagenome-assembled genomes (Bowers et al., 2017).

*Endozoicomonas* genomes, representing a bacterial lineage prevalent in diverse marine invertebrates (Neave et al., 2017a), were retrieved from GenBank (downloaded February 2018). These included the following accession numbers: E. acroporae Acr-14 (GCF_002864045), E. atrinae DSM 22380 (GCF_001647025), E. elysica DSM 22380 (GCF_000710775), E. asciidiicola AVMART05 (GCF_001646945), E. arenosclerae E-MC227 (GCA_001562005), E. namuzuensis DSM 25634 (GCF_000722635), E. montiporae CL-33 (GCF_001583435), and *Endozoicomonas* sp. AB 1-5 (GCA_001729985). Additionally, two single-cell genomes from Stylophora pistillata (“Ca. E. stylophora” type A and B) and
another two metagenome-resolved population genomes from *Acropora humilis* ("Ca. Endozoicomonas humilis") and *Pocillopora verrucosa* ("Ca. Endozoicomonas verrucosa") (Neave et al., 2017b) were retrieved from the RAST server with the following RAST IDs 6666666.127878, 6666666.127879, 305899.6, and 305899.13 respectively.

Another additional set of 52 metagenome-resolved prokaryotic genomes (MAGs), encompassing representative taxa of 12 phyla predominating the microbial microbiome of the Indo-Pacific reef-building coral *Porites lutea* (Robbins et al., 2019) were also retrieved from http://plut.reefgenomics.org/microbiome_download/ (Table S5).

**Prediction of GB metabolism in eukaryotic and prokaryotic genomes**

Profile Hidden Markov Models (profile-HMMs) of protein family alignments were used to query enzymes involved in the transport, biosynthesis, and catabolism of GB and other osmolytes in genomes and transcriptomes. A total of 52 profile-HMMs were retrieved from PFAM (Bateman et al., 2000) and TIGRFAM (Haft et al., 2003) databases (Figure 1; Table S2 and S3). Protein-coding genes and full-length transcriptomic sequences were queried using the program hmmssearch within HMMER v3.2.1 (hmmer.org), applying default settings and an E-value threshold of $1 \times 10^{-6}$ as well as the pre-calculated “gathering” and “noise” cutoffs defined for the 52 model HMMs to search for corresponding protein homologs in the genomes/transcriptomes.

However, eight enzymes lacked profiled HMMs, including those encoding choline oxidase (CodA), choline monooxygenase (CMO), glycine-sarcosine methyltransferase (GSMT), sarcosine-dimethylglycine methyltransferase (SDMT), glycine sarcosine dimethylglycine N-methyltransferase (GSDMT), the GB/choline exporter (EmrE), the eukaryotic sarcosine dehydrogenase (SDH), and betaine aldehyde dehydrogenase (BADH2). For these enzymes, we performed blast searches employing customized protein databases of experimentally characterized proteins available in Swissprot/Uniprot based on the following accession numbers: CodA (*Arthrobacter globiformis*, AAP68832 and Q7X2H8; *Aspergillus fumigatus*, B0YA55); CMO (*Spinacia oleracea*, AAB52509; *Atriplex hortensis*, Q9LKN0; *Amaranthus tricolor*, Q93XE1; *Arabidopsis thaliana*, Q9SZR0; *Beta vulgaris*, BAE07177); GSMT (*Methanohalophilus portucalensis*, 5GWX_A; *Aphanothece halophytica*, Q83WC4; *Synechococcus* sp. WH8102, Q7U4Z8; *Halorhodospira halochloris*, Q9KJ22); SDMT (*H. halochloris*, Q9KJ21; *A. halophytica*, Q83WC3); GSDMT (*Actinopolyspora halophila*, Q9KJ20); EmrE (*Escherichia coli*, P23895); SDH (*Mus musculus*, Q99LB7; *Homo sapiens*, Q9UL12; *Rattus norvegicus*, Q64380); and BALDH2 (*A. thaliana*, Q9STS1 and Q9S795; *Oryza sativa*, Q84L33; *Gadus callarias*, P56533; *Pseudomonas aeruginosa*, Q9HTT1; *Burkholderia mallei*, Q62CH7; *Amaranthus hypochondriacus*, AAB700100; *Gibberella zeae*, EAA74487; *Sinorhizobium meliloti*, P54222; *Bacillus subtilis*, P71016; *Homo sapiens*, P49189). Additional proteins sequences with 90% homology over 80% of the gene length from the UniProt’s (UniRef90) reference database (https://www.uniprot.org) were consolidated into the database of GSDMT (for proteins with 474–564 amino acid sequence length; $n = 140$), EmrE ($n = 100$), and BADH2 ($n = 22$). All customized databases are available from the corresponding author. Protein-coding gene and
transcriptomic sequences were subsequently queried against these customized protein databases using diamond v0.9.24 (Buchfink et al., 2015) with the “sensitive” search mode and an E-value threshold of $10^{-10}$. A minimum amino acid identity of 30% (CMO, codA, GSMT, and SDMT) or 50% (GSMDT, emrE, SDH, and BADH2) and alignment coverage of 60% to the reference sequence was employed for filtering of top hits. The 30% identity cut off was selected because pairwise alignment of ratified enzymes directly in UniProt showed that some of the genes (CMO, codA, GSMT, and SDMT) had pairwise identities as low as 30–40% with alignment coverage minimally around 60%. The 50% cut off was useful in the case of enzymes in which different subfamilies fell into substrate-specific lineages such as BADH2 (s.f. Figure S2) to avoid false positives. We however, contend that the cutoffs my exclude distantly related enzymes catalyzing the target reactions but with different kinetics.

Taxonomic assignment of the retrieved homologs was done based on their blastp matches against the non-redundant NCBI database (downloaded on June 2016) as described by (Ngugi et al., 2017), applying an E-value threshold of $1 \times 10^{-6}$ and a minimum blast score value of 40. Because some eukaryotic genomes likely have microbial contaminants (Artamonova et al., 2015), we employed the lowest-common-ancestor (LCA) approach to flag hits matching prokaryotic genomes (Table S2 and S3). If the taxonomic lineage of the retrieved gene homolog was not consistently supported by at least five blast matches to eukaryotic lineages, then that retrieved gene was completely ignored for downstream analyses. However, only few eukaryotic genomes presented this anomaly; for instance, some genomes (e.g., Avicennia marina and Capitella teleta) and transcriptomes (e.g., Xestospongia testudinaria) were found to have BCCT carriers that were 100% identical to their expected symbionts, indicative of microbial contamination.

Subcellular localization of retrieved protein homologs was predicted using the TargetP server (http://www.cbs.dtu.dk/services/TargetP/) (Emanuelsson et al., 2007) with default “non-plant” settings allowing also for cleavage site prediction.

**Phylogenetic analyses**

All phylogenetic analyses were conducted in Geneious Pro v8.1.9 (https://www.geneious.com) and the best amino acid sequence substitution model for tree inference determined with ProTest3 (Darriba et al., 2011). The deduced eukaryotic BCCT carrier proteins ($n = 72; 749 \pm 185$ amino acid residues) together with validated prokaryotic BCCT proteins ($n = 22; 557 \pm 69$ amino acid residues) were aligned using MUSCLE (Edgar, 2004) with default settings. The alignment was reduced to 92 amino acid sequences by removing shorter proteins and then filtered using Gblocks (Castresana, 2000) to eliminate poorly aligned positions, reducing it to 438 characters based on the liberal settings that retain the maximum number of characters in the alignment. An un-rooted maximum-likelihood phylogenetic tree was reconstructed using RAxML v7.2.830 (Stamatakis, 2014), as implemented in Geneious Pro under the best-fit general time reversible (GTR) + GAMMA model (log-likelihood ratio of $\log likelihood$ ratio of $-33909.16$), with 1,000 bootstraps and a maximum-likelihood search of the best tree topology.
Considering that aldehyde dehydrogenases encompass enzymes with variable substrates (Fitzgerald et al., 2009; Kirch et al., 2004), we were prompted to employ an iterative process to robustly identity genes that were phylogenetically affiliated with *bona fide* betaine aldehyde dehydrogenases (BADH2) among the 100s of matches (in total 1,985) obtained via blast search to our customized blast data (see above). The predicted matches from the genes were clustered into a non-redundant set using cd-hit (Li and Godzik, 2006) by applying global protein identity cutoff of 70% and alignment coverage of 60% for the shorter sequences. Next, the resultant non-redundant proteins (*n* = 927) were aligned together with 22 validated BADH2 amino acid sequences based on the work of Kirch et al. (Kirch et al., 2004) using MUSCLE. The aligned sequences were then used as the basis for a first-run phylogenetic tree with FastTree v.2.1.5 (Price et al., 2010) under the best-fit WAG + GAMMA model (log-likelihood ratio of −34818.25) and using the approximately maximum likelihood method in the fastest mode. This enabled to prune sequences that were distantly affiliated to the validated plant BADH2 sequences, yielding a total of 87 putative BADH genes from the studied eukaryotic genes. To construct the final tree, we gathered 28 additional proteins from NCBI encompassing validated plant aldehyde dehydrogenases with variable substrates and biochemistry (Kirch et al., 2004). The final set of 127 proteins were aligned with MUSCLE resulting in an alignment with 3,727 columns, which were further reduced to 492 column by removing poorly aligned sequence position using Gblocks as stated above. A maximum-likelihood tree was then constructed using RaxML (1000 bootstraps), under the model LG + GAMMA (log-likelihood ratio of −6582.72) selected using Protest3 to accommodate the additional sequences. The reconstructed tree was then manually populated adding the number of redundant sequences in each representative BADH-like gene including the names of organisms harboring that homolog based on the cd-hit information using Adobe Illustrator (available from http://adobe.com/products/illustrator). For the final presence/absence profile only genomes with a gene phylogenetically related to family 10 ALDHs (Figure S2) were scored as having a putative BADH2-like gene (Figure 1b) in addition to genes matching characterized ALDH family 10 proteins (via blast search).

**Coral sampling**

Samples of 10 scleractinian coral species (*Acanthastrea echinata*, *Diploastrea heliopora*, *Fungiia* sp., *Galaxea fascicularis*, *Mycedium elephantotus*, *Pavona varians*, *Plerogyra sinuosa*, *Pocillopora verrucosa*, *Porites lutea*, *Stylophora pistillata*) and one octocoral (*Xenia* sp.) were collected at the ocean-facing side of Al Fahal reef (22.3034, 38.9602) in the central Red Sea, Saudi Arabia (*n* = 4 to 5 replicates per species; Table S7). The sampled species fall across many coral clades and life histories, thus attractive for evaluating metabolism of GB in diverse corals. Five replicate colonies per species were sampled on May 17 and 18 (2016) at depths of 3 to 18 m. One small piece (3–4 cm) was collected per colony using hammer and chisel, transferred into Whirlpack bags, and immediately flash-frozen upon return to the boat.

**DNA extraction and whole-genome sequencing**

DNA was extracted using a DNA All Prep kit (#80204) from Qiagen with modifications. Briefly, corals in Whirlpack bags were first defrosted and a
pressurized air gun was used to remove host tissue and putative Symbiodiniaceae cells away from the skeleton and into the Whirl Pak. Subsequently, 30 mg of air blasted tissue slurry was weighed out in a sterile weigh boat, where 600 µl of buffer RLT was added to the slurry. The mixture was homogenized with a 20-gauge needle on a 1-ml syringe, and the buffer and tissue mixture were placed in a new 2 ml tube. DNA was then extracted as recommended by the manufacturer. The extracted DNA was then quantified on a Qubit prior to library preparation.

100 ng of DNA per sample was sheared to approximately 250–300 bp (Covaris M2). Sheared DNA was end-repaired and A-tailed with a single A dNTP, the NEB adapter (NEB #E7645L) was then ligated to the end-repaired A-tailed DNA fragment. Size selection of the library was performed using Agencourt AMPure XP beads (Beckman Coulter) for size selection of 250 bp fragment inserts. Dual indexes were used in the library enrichment step to add Illumina TruSeq HT indexes (NEB #7600) to the DNA molecules using 6 cycles of PCR. Libraries were quality checked using the Bioanalyzer DNA 1000 Chip (Agilent Technologies), quantified using the Qubit BR DNA system (Invitrogen), and subsequently pooled in equimolar ratios to a final concentration of 10 nM. The pool was re-quantified using qPCR (KAPA Biosystems library quantification) with the ABI HT7900 (Applied Biosystems) and paired-end sequenced (2 × 300 bp) on the Illumina NextSeq 500 at 1.8 pM with 1 % PhiX.

**Metagenomic sequence analyses**

Raw read sequences were quality filtered and trimmed using Trimmomatic v0.32 (Bolger et al., 2014) to remove adapter sequences and leading and trailing bases with a quality score below 20, less than 50 base-pairs (bp) long as well as reads with an average per base quality of 20 over a 4-bp window. Paired-end (PE) reads were mapped onto run quality control sequences (PhiX genome) using BBmap v37.44 (http://jgi.doe.gov/data-and-tools/bbtools/). At each stage, the quality of read sequences was assessed using FASTQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/).

The resulting high-quality PE reads for each dataset (n = 4–5 per coral species) were *de novo* assembled independently with metaSPAdes v3.9.0 (Bankevich et al., 2012) using a kmer range of 21 to 127, while employing the error-correction mode and preset metagenomic options. The assembled contigs were then filtered to a minimum length of 500 bp followed by gene prediction using Prodigal v2.6.2 (Hyatt et al., 2010). Prodigal was run with the options “-p meta -n -m”. Table S7 summarizes the general statistics of read sequences, the assembled contigs, and predicted genes.

Individual metagenomic protein-coding genes were interrogated as described above but focusing on the BCCT carrier family (PF02028) regarding potential high-affinity GB transporters in different coral species from the Red Sea. Here as well, the resultant matches were searched against the nr protein database to clarify the origin of hits, and only those matching to anthozoans were retained for downstream analyses.

Gene density was computed from the counts of retrieved hits per metagenome scaled by the total length of coding sequences (in megabase-pairs) among the
different coral species. All predicted coding genes in each metagenome and the recovered transporters were taxonomically assigned as described above.

**Gene expression analyses using available metatranscriptomes**

Using ten recently published RNA-Seq data covering the larval–adult developmental cycle and two symbiotic states of the laboratory-raised sea anemone Aiptasia (Baumgarten et al., 2015), we tracked the transcriptional activity of GB metabolism in this model dinoflagellate-cnidarian symbiosis. Briefly, raw sequencing reads were downloaded from the short-reads archive under accession number SRX286028 and quality-trimmed, while removing adaptor sequences, using Trimmomatic v0.323 (Bolger et al., 2014) with the following parameters: ILLUMINACLIP::4:30:10 LEADING:20 TRAILING:20 SLIDINGWINDOW:4:20 MINLEN:60. The internal sequencing standard PhiX 174 was subsequently removed by mapping the quality-trimmed reads against the PhiX 174 genome using Bowtie2 v2.2.45 (Langmead and Salzberg, 2012), with default settings. Potential errors in the reads were then corrected using SPAdes’ implementation of the BayesHammer error-correction tool with default parameters (Bankevich et al., 2012). Ribosomal RNA-like reads were removed using SortMeRNA v1.99 (Kopylova et al., 2012). The resulting high-quality PE reads from each sample were mapped against the Aiptasia v1 gene models (http://reefgenomics.org/) using Bowtie2 (Langmead and Salzberg, 2012), with default settings. The mapped read counts per gene (and sample) were subsequently normalized into a common metric of gene expression—that is, fragments per kilobase of exon per million fragments mapped (FPKM) using an in-house perl script.

To evaluate whether genes involved in GB metabolisms are actively expressed in free-living Symbiodiniaceae, we interrogated a catalog of 116 million expressed eukaryotic unigenes of the Tara Ocean Expedition (Carradec et al., 2018) available at http://www.genoscope.cns.fr/tara/.

The available Pfam annotations of the unigene catalog were used to retrieve the expression values of selected four enzymes of GB metabolism, including PF02028 (BCCT), PF02574 (BHMT), PF16350 (DMGDH) and PF00464 (SHMT). In addition, local blast searches were performed as described above in the case of CodA, CMO and SDMT. In both cases, we retained genes taxonomically annotated as “Sueciaceae” at the lowest rank as well as translated blast matches with ≥80% identity and alignment length of ≥60% against reference genes from Symbiodiniaceae genomes and transcriptomes (this study). Expression values of homologs that showed significant similarity to Symbiodiniaceae genes were then plotted using the R package ggplot2 (Wickham, 2009).

**Global estimates of GB stocks bound in marine picoplankton cells**

To estimate the global stock of GB bound in marine picoplankton cells (0.2 to 3.0 µm in size), we interrogated metagenomic datasets from the Tara Ocean global circumnavigation expedition (Sunagawa et al., 2015) to estimate the fraction of marine prokaryotes capable of synthesizing GB using the abundance of betA, betB, BADH2, and codA homologs as marker genes for two different choline oxidation pathways (Figure 1a). Briefly, protein-coding genes from 201 (out of 243) Tara metagenomes covering photic to epipelagic depths (5–70 m) were retrieved from the European Nucleotide Archive.
These metagenomes were probed based on profiled HMMs (BetA and BetB) and the customized BADH2 and CodA protein databases as described in the previous section. These four enzymes were selected because marine reference prokaryotic genomes possess a single copy of each gene homolog on average and also because they are widespread (Figure S8), making them ideal for estimating the abundance of microorganisms capable of producing GB in the ocean.

In parallel, we also retrieved homologs of four conserved single-copy genes (SCGs) in the same metagenomes—that is, recA (recombinase; TIGR02012), ligA (NAD-dependent DNA ligase; TIGR00575), pyG (CTP synthase; TIGR00337), fmt (methyl-tRNA formyltransferase; TIGR00460). Based on the retrieved counts of these SCGs, we then retained metagenomes with at least 15 RecA hits and removed those in which the coefficient of variation (CV) in the counts of all four SCGs was ≥ 25% (mean CV was 16 ± 5%). This yielded a final set of 108 metagenomes encompassing microorganisms with a size range of 0.22 µm to 3 µm (Table S9). Finally, the number of unique hits to betA, betB, BADH2, and codA genes were normalized to the average number of unique hits to the four SCGs in the metagenomes to estimate the environmental abundance of GB-producing picoplankton in the pelagic sea (Table S9). The betA, betB, BADH2, and codA homologs were estimated to be present in 2.4, 0.3, 1.2, and 34.1% (Table S9), respectively of the sampled picoplankton in the Tara metagenomes (n = 108).

To calculate the pool size of GB bound in marine pelagic picoplankton (0.2 to 3 µm in size), we first made four assumptions, namely that (1) a milliliter of seawater contains on average 10^5 pico-sized cells (Whitman et al., 1998); (2) the total biomass of prokaryotic picoplankton in the pelagic ocean is about 3.6 × 10^{28} cells (Whitman et al., 1998); (3) about 2% of marine picoplankton communities produce (and store) GB intracelullarly based on the environmental abundance of betA and BADH2 genes encoding key synthetic routes for GB in marine metagenomes (Table S9)—equivalent to roughly 36 × 10^{26} cells in the upper 200 m globally; and (4) the concentrations of cellular-bound (particulate) GB quantified from 0.2-µm-filtered seawater particulates of about 0.9 to 483 nanomolar (Beale and Airs, 2016; Cree, 2015; Keller et al., 2004)—that is, GB bound in marine picoplankton cells.

**Statistical Analyses**

Significant differences between the average count of recovered genes among different groups and coral species was tested by conducting a one-way analysis of variance (one-way ANOVA) using GraphPad Prism v7.0a (GraphPad Software, Inc.). Multiple comparisons for controlling the false discovery rate (α = 0.05) was conducted with the Benjamini–Hochberg’s two-stage step up method (Benjamini et al., 2006).

**Data and materials availability**

The Raw sequence data for the coral metagenomes have been deposited in NCBI under BioProject number PRJNA437202 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA437202). All data needed to evaluate the results and conclusions are presented in the main text and/or are available in the Supplementary Materials. Additional data or scripts related to
this paper are available from the corresponding authors.

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