



The Osmolyte Ties That Bind: Genomic Insights Into Synthesis and Breakdown of Organic Osmolytes in Marine Microbes

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The production and consumption of organic matter by marine organisms plays a central role in the marine carbon cycle. Labile organic compounds (metabolites) are the major currency of energetic demands and organismal interaction, but these compounds remain elusive because of their rapid turnover and concomitant minuscule concentrations in the dissolved organic matter pool. Organic osmolytes are a group of small metabolites synthesized at high intracellular concentrations (mM) to regulate cellular osmolarity and have the potential to be released as abundant dissolved substrates. Osmolytes may represent an essential currency of exchange among heterotrophic prokaryotes and primary and secondary producers in marine food webs. For example, the well-known metabolite dimethylsulfoniopropionate (DMS) is used as an osmolyte by some phytoplankton and can be subsequently metabolized by 60% of the marine bacterial community, supplying up to 13% of the bacterial carbon demand and 100% of the bacterial sulfur demand. While marine osmolytes have been studied for decades, our understanding of their cycling and significance within microbial communities is still far from comprehensive. Here, we surveyed the genes responsible for synthesis, breakdown, and transport of 14 key osmolytes. We systematically searched for these genes across marine bacterial genomes ($n = 897$) and protistan transcriptomes ($n = 652$) using homologous protein profiles to investigate the potential for osmolyte metabolisms. Using the pattern of gene presence and absence, we infer the metabolic potential of surveyed microbes to interact with each osmolyte. Specifically, we identify: (1) complete pathways for osmolyte synthesis in both prokaryotic and eukaryotic marine microbes, (2) microbes capable of transporting osmolytes but lacking complete synthesis and/or breakdown pathways, and (3) osmolytes whose synthesis and/or breakdown appears to be specialized and is limited to a subset of organisms. The analysis clearly demonstrates that the marine microbial loop has the genetic potential to actively recycle osmolytes and that this abundant group of small metabolites may function as a significant source of nutrients through exchange among diverse microbial groups that significantly contribute to the cycling of labile carbon.

Keywords: osmolytes, glycine betaine, mannitol, transporters, biosynthesis, metatranscriptomics

1. INTRODUCTION

Marine microbes live in a high salinity environment and must maintain cellular mechanisms for rapid response to any small changes in external water potential in order to maintain cellular homeostasis (Bisson and Kirst, 1995). Ions are most readily transported by microbes, and therefore their concentrations typically respond first to external salinity changes. However, the charged nature of ions can only be tolerated by cellular physiology at specific concentrations, and ions are most often stored in solution within vacuoles in phytoplankton. Thus, organic osmolytes represent a critical addition to cellular osmotic balance. Throughout the rest of this paper when we refer to osmolytes, we will be exclusively discussing organic osmolytes. Osmolytes are often referred to as compatible solutes as they are small, organic molecules that are highly soluble and can accumulate at high concentrations within the cytoplasm without interfering with cellular function (Stefels, 2000; Roeßler and Muller, 2001). Although organic osmolytes are energetically more costly to use than inorganic ions, because they must be either synthesized or actively transported into the cell, many serve multiple functions in addition to their osmotic contributions. Osmolytes have been shown to contribute to protein stabilization and to have other cytoprotective properties under salinity, temperature, and pressure stress (Kirst, 1989; Burg and Ferraris, 2008; Yancey and Siebenaller, 2015; Ma et al., 2017). Some osmolytes have been shown to be efficient scavengers of reactive oxygen species (Smirnov and Cumbes, 1989; Sunda et al., 2002; Brands et al., 2019). Osmolytes have also been predicted to function in a ballasting mechanism, where the synthesis of osmolytes with different densities would alter cellular buoyancy (Boyd and Gradmann, 2002; Lavoie et al., 2015). Osmolytes have been of interest in the marine context for decades (Challenger, 1951; Yancey et al., 1982; Kiene and Hoffmann Williams, 1998; Keller et al., 1999a). However, with the exception of a few extensively studied osmolytes (e.g., dimethylsulfoniopropionate, DMSP), we know relatively little about osmolyte distributions in the ocean, the role of environmental conditions in controlling intracellular osmolyte abundances and subsequent release into the dissolved phase, or the significance of osmolytes as substrates in the microbial loop.

In marine systems, a range of amino acids and their derivatives, carbohydrates, methylsulfonium compounds, and methylammonium compounds have been identified as osmolytes used by marine organisms ranging from bacteria to tube worms (Yancey, 2005) (Table 1). All macro- and microalgae exhibit the ability to accumulate both inorganic ions and organic osmolytes (Bisson and Kirst, 1995), though their salinity tolerances have been shown to be a function of the isolation location's salinity conditions (open ocean vs. coastal ocean) (Brand, 1984; Kirst, 1989). In general, phytoplankton contain one or two major organic osmolytes with concentrations >50 mM, and many minor organic osmolytes with concentrations <5 mM (Gebser and Pohnert, 2013). However, monoculture studies have demonstrated that the composition of organic osmolytes is variable and strongly taxa dependent (Dickson and Kirst, 1987a,b). Marine phytoplankton use many well-known plant

osmolytes (Yancey, 2005), but also osmolytes that are primarily found in the marine environment. For example, DMSP is found in a diverse array of marine phytoplankton, and was originally described as an osmolyte due to its similar structure to glycine betaine (Andreae, 1986). DMSP is hypothesized to be multi-functional, but likely functions predominantly as an osmolyte in dinoflagellate and haptophyte groups which typically contain >50 mM intracellular DMSP (Keller et al., 1989; Stefels, 2000; McParland et al., 2020). 2,3-dihydroxypropane-1-sulfonate (DHPS) was very recently reported to be present at osmolyte-like concentrations in some diatoms (15 mM) and coccolithophores (18 mM) (Durham et al., 2019), and its intracellular concentrations in the Antarctic sea-ice diatom *Nitzschia lecontei* (~ 85 mM) correlate with salinity and temperature changes (Dawson et al., 2020a). Some marine diatoms have also been shown to be capable of transporting ectoine or DMSP for osmotic functions (Spielmeyer et al., 2011; Lavoie et al., 2018; Fenizia et al., 2020). Across prokaryotes, halophilic archaea and bacteria have uniquely adapted cellular machinery that can withstand molar concentrations of potassium and chloride ions (Roeßler and Muller, 2001), but most marine prokaryotes also utilize organic osmolytes. Some prokaryotes maintain *de novo* synthesis pathways, while many others are capable of transporting osmolytes available in the dissolved pool (Poretsky et al., 2010; Poli et al., 2017). Two well-known marine nitrogen-fixing Cyanobacteria use different organic osmolytes from each other: *Crocospaera* synthesizes trehalose, while *Trichodesmium* synthesizes homoserine betaine (Pade et al., 2012, 2016). The chemoautotroph *Sulfurimonas* was recently shown to synthesize proline with increasing salinity (Götz et al., 2018), while *Vibrio parahaemolyticus* synthesizes ectoine as an osmolyte (Ongagna-Yhombi and Boyd, 2013). Rather than synthesizing osmolytes *de novo*, prokaryotes can also transport and retain them under salinity stress. For example, uptake of radiolabeled DMSP and glycine betaine by natural communities of bacterioplankton has been shown to increase with salinity, and a significant proportion of the spiked osmolytes remained untransformed after uptake (Kiene and Hoffmann Williams, 1998; Motard-Côté and Kiene, 2015).

Osmolytes are some of the most concentrated (mM) small molecules in marine microbes, and therefore have the potential to be released as abundant carbon substrates to the dissolved organic matter (DOM) pool. The small size of osmolytes make them inherently labile substrates compared to larger biopolymers, such as proteins, polysaccharides, and lipids, which require additional extracellular processing. Uptake studies of DMSP, glycine betaine, and taurine have estimated that these molecules provide up to 13% (Kiene and Linn, 2000), 38% (Kiene and Hoffmann Williams, 1998), or 4% (Clifford et al., 2020) of bacterial carbon demand, respectively. When we consider the scale of the global carbon flux that is used by bacterioplankton (50% of net primary productivity, Ducklow, 2000), these substrates could contribute monumentally to this flux of energy and matter and dictate relationships amongst organisms. Catabolic pathways for other osmolytes, such as those identified for ectoine and 5-hydroxyectoine in the marine bacterium, *Ruegeria*

TABLE 1 | Osmolytes previously described in marine microbes.

Common name	g/mol	Category	N atoms	S atoms	Cellular	Dissolved (nM)
Trigonelline	137	Alkaloid	0	1	+	-
TMAO	75	Amine oxide	1	0	+	2–77 (Hatton and Gibb, 1999; Gibb and Hatton, 2004)
ectoine	142	Amino acid	2	0	+	0.1–0.3 (Widner et al., 2021)
GABA	103	Amino acid	1	0	+	0.1 (Widner et al., 2021)
Glutamate	147	Amino acid	1	0	+	0.7–5 (Mopper and Lindroth, 1982; Widner et al., 2021)
Glutamine	146	Amino acid	2	0	+	0.2–3 (Mopper and Lindroth, 1982; Widner et al., 2021)
Glycine betaine	117	Amino acid	1	0	+	-
Homarine	137	Amino acid	1	0	+	-
Homoserine betaine	161	Amino acid	1	0	+	-
Hydroxyectoine	158	Amino acid	2	0	+	-
Proline	115	Amino acid	1	0	+	0.4 (Widner et al., 2021)
Sarcosine	89	Amino acid	1	0	+	0.1–0.2 (Widner et al., 2021)
Taurine	125	Amino acid	1	1	+	0.1–320 (Clifford et al., 2017; Widner et al., 2021)
Glucosylglycerol	254	Sugar	0	0	+	-
Isofloridoside	254	Sugar	0	0	+	-
Sucrose	342	Sugar	0	0	+	BD (Sakugawa and Handa, 1985)
Trehalose	342	Sugar	0	0	+	BD (Sakugawa and Handa, 1985)
Glycerol	92	Sugar alcohol	0	0	+	-
Mannitol	182	Sugar alcohol	0	0	+	-
Sorbitol	182	Sugar alcohol	0	0	+	-
DHPS	155	Sulfonate	0	1	+	0.26–0.44 (Widner et al., 2021)
DMSP	134	Sulfonium	0	1	+	1–2 (Kiene and Slezak, 2006)
Gonyol	178	Sulfonium	0	1	+	-

The table includes: Common name of osmolyte, molecular weight (g/mol), nominal category, number of nitrogen (N) atoms, number of sulfur (S) atoms, cellular concentrations reported (+) in monocultures, and previously measured dissolved concentrations (nM) from coastal or open ocean environments. Names in bold were used in the analyses presented here. As osmolyte cellular concentrations are extremely variable, we only report if an osmolyte has been previously reported in monocultures (Brown and Hellebust, 1978; Reed et al., 1986; Dickson and Kirst, 1987a,b; Keller et al., 1989; Mountfort and Pybus, 1992; Yancey, 2005; Pade et al., 2012, 2016; Gebser and Pohnert, 2013; Durham et al., 2019; Fenizia et al., 2020; Lin et al., 2020). To our knowledge, dissolved measurements of most osmolytes do not exist. GABA = 4-aminobutanoate, TMAO = trimethylamine *n*-oxide, DMSP = dimethylsulfoniopropionate, DHPS = dihydroxypropane-sulfonate, BD = below detection.

pomeroyi, indicate that other osmolytes could also be valuable substrates (Schulz et al., 2017), but lack relevant environmental measurements to quantify their abundance and role within the microbial loop.

Previous measurements of osmolytes within the DOM pool range from \sim pM to nM concentrations (Table 1), which contrast with the mM intracellular concentrations of osmolytes, suggesting that osmolytes are rapidly consumed upon release to the dissolved pool. As osmolyte concentrations can approach limits of detection with current analytical techniques, the down-regulation of transcripts associated with osmolyte consumption have been quantified as a proxy for utilization (Vorobev et al., 2018). When applying this technique to natural communities, the osmolytes glycine betaine, mannitol, taurine, proline, and sorbitol were all predicted to be rapidly recycled in <24 h (Vorobev et al., 2018). Osmolyte concentrations and availability have been hypothesized to be a function of both community composition and other environmental drivers. For example, DMSP concentrations were predicted to be determined by phytoplankton community composition (McParland and Levine, 2019). Recent analytical advances have expanded our knowledge to a wide array of osmolytes and their environmental variability (including trehalose, sucrose, glucosylglycerol, DHPS, DMSP,

glycine betaine, proline, homarine). Particulate abundances of some osmolytes are linked to diel cycles (Boysen et al., 2020), and different osmolytes dominate on sinking particles (glycine betaine, proline) compared to suspended particles (i.e., planktonic cells; DMSP) (Johnson et al., 2020). Although extensive measurements of DMSP concentrations and turnover rates exist (Kiene and Linn, 2000), the dissolved concentrations and turnover rates of most marine osmolytes are unknown (Table 1) and require further experimental work to fully understand the factors that drive their distributions.

In this study, we explored the role of osmolytes within the marine microbial loop by identifying microbes with the potential to synthesize, catabolize, assimilate, or transport osmolytes. We hypothesized that different patterns of osmolyte metabolisms would reveal the role of osmolytes as valuable nutrients (i.e., potential carbon, nitrogen, and/or sulfur substrates) in the microbial loop. We first explored the taxonomic trends of potential producers and consumers by surveying osmolyte synthesis and utilization in genomes and transcriptomes from monocultures. We then compared these results with *in situ* transcription of mannitol and glycine betaine transport, synthesis, and breakdown in the Tara metatranscriptomes from the surface ocean.

TABLE 2 | Osmolytes included in prokaryote and eukaryote surveys.

Common name	KEGG ID	Breakdown	Synthesis	Transport
Ectoine	C06231	2 (1–3)	2 (4–5)	0
GABA	C00334	4 (1–2)	11 (1–4)	0
Glutamate	C00025	61 (1–5)	78 (1–3)	0
Glutamine	C00064	30 (1)	2 (1)	0
Glycine betaine	C00719	1 (3)	4 (1–2)	2 (3)
Proline	C00148	12 (1)	6 (1–2)	0
Sarcosine	C00213	2 (1)	7 (1–4)	0
Taurine	C00245	6 (1)	5 (1)	1 (3)
Glycerol	C00116	7 (1–3)	9 (1–4)	1 (3)
Mannitol	C00392	2 (1–2)	1 (2)	1 (3)
Sorbitol	C00794	4 (1)	4 (1)	1 (3)
Sucrose	C00089	12 (1)	3 (1–3)	0
Trehalose	C01083	6 (1)	5 (1–3)	0
TMAO	C01104	1 (1)	1 (1)	0

The table includes: Common name of osmolyte, the KEGG ID, the total number of curated pathways with associated KOs used in the searches for breakdown, synthesis, and transport, and the range of steps in each pathway in parentheses. Some pathways required the presence of multiple steps (i.e., KOs) to be considered complete (**Supplementary Material 1**). A single step value indicates all pathways had the same number of steps. For osmolyte pathways with differing number of steps, a range is reported which reflects the minimum and maximum number of steps for all pathways. GABA = 4-aminobutanoate, TMAO = trimethylamine *n*-oxide.

2. MATERIALS AND METHODS

2.1. Manual Curation of Osmolyte-Related KEGG Orthologs

Osmolytes ($n = 23$) were initially selected for investigation based on their ubiquity within microorganisms or their previously identified importance in marine systems (**Table 1**). We chose to use the Kyoto Encyclopedia of Genes and Genomes (KEGG) as the database from which to identify metabolic pathways associated with the osmolytes and the corresponding gene orthologs, referred to as KEGG orthologs (KOs) throughout this paper (Kanehisa and Goto, 2000; Kanehisa et al., 2012). Only 18 of these osmolytes of interest were metabolites mapped in KEGG. In addition, glucosylglycerol had to be excluded because there were no genes associated with this compound in KEGG and trigonelline was also excluded due to missing KOs in the pathway available on KEGG. Dimethylsulfoniopropionate (DMSP) was also removed from the list of osmolytes considered because the pathways involved in its biosynthesis and breakdown are incomplete in KEGG. Finally, hydroxyectoine was not analyzed in detail as it is a closely linked derivative of ectoine. Thus, our analysis was performed on 14 osmolytes (**Table 2**).

The KOs associated with the synthesis, breakdown, and transport of these metabolites were manually compiled (**Supplementary Material 1**). We sought to identify synthesis pathways that began with a common metabolite, such as a sugar or amino acid, rather than an intermediate. Similarly, we mapped breakdown pathways to endpoints that could be incorporated back into metabolism for either catabolic or anabolic purposes. Transporters are not comprehensively represented in KEGG

and thus we chose to focus on the available ABC transporters. This limited us to transporters for glycerol, glycine betaine, mannitol, sorbitol, and taurine. By mapping the pathways in this way, we sought to identify organisms that could use ubiquitous building blocks to synthesize an osmolyte and those who could breakdown an osmolyte into biologically relevant molecules that could be used for another metabolic purpose.

To facilitate identification of synthesis and breakdown capabilities in organisms, we identified each of the genes required for a selected pathway by numerically labeling each gene as a step in a pathway as well as indicating alternative complete pathways (**Supplementary Material 1**). Finally, we created a comprehensive list of the KOs associated with synthesis, breakdown, and transport for each osmolyte to use in downstream homology searches. This comprehensive list resulted in the definition of 482 possible reaction steps or transporter components, which represented 379 unique reactions with assigned KEGG reaction numbers. Ultimately, 486 unique KOs were identified (**Supplementary Material 2**). To confirm that our manual curation had comprehensively captured the genes associated with each osmolyte according to KEGG, we computationally searched a list of KOs directly associated with each osmolyte and checked this list against our manually curated pathways.

2.2. Homolog Searching of KEGG Orthologs Against Marine Prokaryotic Genomes and Eukaryotic Protistan Transcriptomes

Genomes from representative marine bacteria and archaea were collected from MarRef (v5) ($n = 897$) (Klemetsen et al., 2017; ten Hoopen et al., 2017), and transcriptomes from representative marine protists were collected from the Marine Microbial Eukaryotic Transcriptome Sequencing Project (MMETSP) ($n = 652$) (Keeling et al., 2014; Johnson et al., 2018) (**Supplementary Material 2**). The predicted proteins from both the prokaryotic genomes and MMETSP transcriptomes were used in the subsequent analyses. All 486 KOs that were identified through manual curation as associated with the metabolic processing or transport of key osmolytes (**Table 2**) were searched against the predicted proteins from the MMETSP and MarRef datasets. A snakemake pipeline (v. 5.24.2) (Mölder et al., 2021) based off of kofamscan (Mistry et al., 2013; Aramaki et al., 2019) was built to search for the likely presence of the osmolyte-related KOs in our protein sets of interest. KO hmm profiles were downloaded from KEGG (<https://www.genome.jp/tools/kofamkoala/>) on October 6, 2020. Both the MarRef and MMETSP predicted protein sets were searched for a given KO with hmmsearch (HMMER v. 3.3.1). True hits for KOs were parsed from the resulting hmm output with a custom script (parse_hmmsearch.py) run with python v. 3.9.1 and biopython v. 1.78 (Cock et al., 2009), which parses hits based on the adaptive bitscore cutoff values used by kofamscan as published in the ko_list file (Aramaki et al., 2019). A final table with presence or absence of all osmolyte-related KOs for each organism was generated (**Supplementary Material 2**). This pipeline and

all associated scripts and analysis notebooks is available as a GitHub repo (<https://github.com/AlexanderLabWHOI/2021-marine-osmolytes>).

2.3. Metabolic Potential Prediction

Using the manually curated pathway listings (**Supplementary Material 1**), a python based script was used to identify the presence or absence of a given metabolic pathway associated with the synthesis, breakdown, or transport for an organism (**Supplementary Material 3**). To be considered capable of osmolyte synthesis or breakdown, an organism was required to have at least one complete pathway between the identified common metabolite (as described above). In cases where a pathway had more than one step (**Table 2**), the organism had to have all steps present for it to be annotated as capable of synthesis or breakdown. In cases where a step was identified on a pathway map, but was not annotated with a KO, the pathway was not considered. For the annotation of transporters, genomes were required to contain all KOs (subunits) of a given transporter to be annotated as capable of transport. In the case of glycine betaine, which has two different transporters, only one transporter system was required to be present to confer the ability to transport.

2.4. Prokaryotic Phylogeny Comparison

Phylogenetic trees from the MarRef genomes were constructed using GToTree v1.4.7 (Price et al., 2010; Lee, 2019). Trees were constructed with the ‘Universal’ dataset of 15 genes as defined by Hug et al. (2016) and default parameters. Trees were visualized using iTOL (Letunic and Bork, 2016) and the presence or absence of synthesis, breakdown, and transport capabilities were visualized across all genomes and publication-quality figures were generated.

2.5. Identifying Orthologs in Tara Oceans

To assess the potential importance of osmolyte metabolic processes across the global ocean, KOs from genes associated with glycine betaine and mannitol synthesis, breakdown, and transport were searched against the surface data from the Tara Oceans dataset using the Tara Oceans Gene Atlas (Villar et al., 2018). A selenium-based web bot was used to automate the search submission and result download process (<https://github.com/AlexanderLabWHOI/2021-marine-osmolytes/tree/master/genome-searching/tara-scraping>). KO file hmm profiles from kofamscan were uploaded to Tara Ocean gene atlas and searched against the metagenome and metatranscriptome datasets for both the “eukaryotic” dataset, Marine Atlas of Tara Oceans Unigenes (MATOU v1) (Carradec et al., 2018), and Marine Atlas of Tara Oceans Unigenes and the “prokaryotic” dataset, Ocean Microbial Reference Gene Catalog (OM-RGC v2, including the Arctic dataset) (Salazar et al., 2019). All searches were initially run with a generous *e*-value cutoff of $1e-10$ and all count-based data were returned as “percent of mapped reads.” We filtered the returned alignment files to retain only orthologs which passed the identified bitscore cutoff from kofamscan (**Supplementary Material 4**), such as we used in our genomic and transcriptomic analyses above. Notably, these cutoff values

are quite stringent and significantly reduced the number of hits we reported (**Supplementary Material 4**). It is likely that assembled metagenomic or metatranscriptomic orthologs that are incomplete were not included in the analysis, but we believe this conservative cutoff best reflects true hits in the Tara dataset. Taxonomic annotation of orthologs recovered was assessed using the reported taxonomy from Ocean Gene Atlas.

2.6. Plotting and Statistical Analysis

Data was analyzed and figures were generated with matplotlib v. 3.3.1 (Hunter, 2007), pandas v. 1.1.2 (McKinney, 2010), and seaborn v. 0.11.0 (Waskom, 2021). Spearman correlations were calculated with the stats.spearmanr function within scipy v. 1.5.2 (Virtanen et al., 2020). Multi-testing *p*-value correction was done with the multipletests function within statsmodels 0.12.1 using the Bonferroni method. Global map distributions were plotted using cartopy 0.18.0 (Met Office, 2010–2015).

3. RESULTS AND DISCUSSION

3.1. Osmolyte Synthesis, Breakdown, and Transport in Marine Prokaryote and Eukaryote Monocultures

In this study, we manually curated the genes responsible for the synthesis, breakdown, and transport of marine osmolytes ($n = 14$) (**Tables 1, 2**) that are classified as either amino acids and their derivatives ($n = 8$), sugars ($n = 5$), or an amine oxide ($n = 1$). We then assessed the presence of these osmolyte-associated genes across marine prokaryotic genomes (MarRef) and marine eukaryotic transcriptomes from protists (MMETSP). The MarRef genomes ($n = 897$) contain representatives from 23 bacterial and 5 archaeal phyla. Notably, the genomes are biased toward copiotrophic lifestyles (and consequently more easily cultured and maintained in a laboratory setting) (Pachiadaki et al., 2019). The eukaryotic data targeted here constitute transcriptomes from protists, drawn from the MMETSP ($n = 652$). These transcriptome datasets contain representatives from across eight of the major supergroups within the eukaryotic tree of life.

Specifically, we looked at patterns of breakdown, synthesis, and transport across major prokaryotic and eukaryotic groups (**Figures 2, 3**). The total number of pathways searched ranged from 1–75 for synthesis and 1–50 for breakdown (**Table 2**). The majority of pathways were comprised of a single step, though one breakdown (glutamate) and one synthesis pathway (ectoine) had a maximum 5 steps (**Table 2**). Relative to osmolyte synthesis and breakdown, we looked at only a limited set of osmolyte transporters, focusing only on ABC transporters ($n = 5$ pathways total). Glycine betaine has two different transport systems annotated in KEGG, and mannitol and sorbitol have the same transporter (**Supplementary Material 1**). More than half of all prokaryotic genomes surveyed contained at least one of the transporter systems ($n = 469$), demonstrating the prevalence of transport in marine prokaryotes. The presence or absence of osmolyte synthesis, breakdown, and

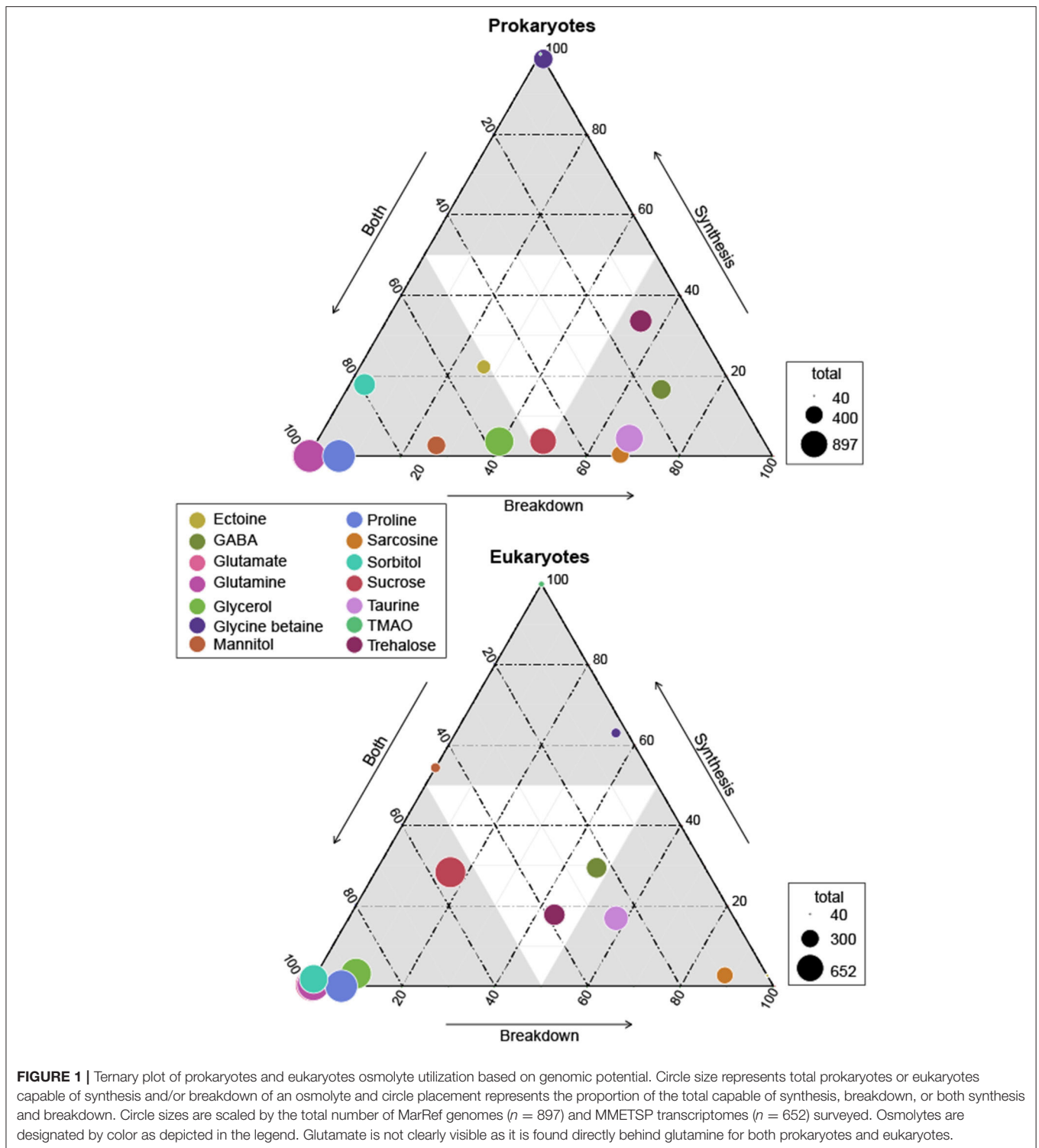


FIGURE 1 | Ternary plot of prokaryotes and eukaryotes osmolyte utilization based on genomic potential. Circle size represents total prokaryotes or eukaryotes capable of synthesis and/or breakdown of an osmolyte and circle placement represents the proportion of the total capable of synthesis, breakdown, or both synthesis and breakdown. Circle sizes are scaled by the total number of MarRef genomes ($n = 897$) and MMETSP transcriptomes ($n = 652$) surveyed. Osmolytes are designated by color as depicted in the legend. Glutamate is not clearly visible as it is found directly behind glutamine for both prokaryotes and eukaryotes.

transport appeared to be phylogenetically linked in prokaryotes (Supplementary Figures 1–3).

We found that either: (1) breakdown and synthesis were tightly linked, suggesting core metabolism/internal recycling, or (2) breakdown and synthesis ability were unequal suggesting

that the utilization of these osmolytes is a more specialized metabolism potentially only present in a smaller portion of the community. Additionally, most prokaryotic genomes that contained a transport system, also harbored the ability to synthesize and/or breakdown the respective osmolyte, except

for glycine betaine for which many genomes were capable of transport without synthesis or breakdown.

3.2. Amino Acids and Their Derivatives

3.2.1. Glutamate and Glutamine

The ability to both synthesize and breakdown glutamate and glutamine was found in almost all prokaryotes ($n = 897$, $n = 895$, respectively) and eukaryotes ($n = 647$, $n = 645$, respectively) surveyed (Figure 1). This widespread function was expected as these amino acids play a fundamental role in the creation and structure of proteins, and also in nitrogen recycling in cells (Reitzer, 2003). Additionally, glutamate and glutamine abundance has been documented to rapidly change in response to altered osmotic conditions (Saum and Müller, 2008). Only two prokaryotic genomes lacked the ability to synthesize glutamine, which was likely due to either an incomplete genome assembly or an issue of homology in our searches. The synthesis and breakdown of glutamate and glutamine was also missing in just 3 and 4 eukaryotic transcriptomes, respectively. Again, this was likely due to lack of coverage in the transcriptome, and potentially related to the physiological condition of the organism at the time of sequencing.

3.2.2. Proline

Both breakdown and synthesis of proline was also common and widespread across prokaryotes ($n = 838$) and eukaryotes ($n = 349$), though slightly less so than glutamate and glutamine (Figure 1). In particular, Archaea (52%) were less likely to have an identified proline synthesis pathway than bacterial groups (75–100%) (Figure 2 and Supplementary Figure 4). Proline synthesis was much more variable across eukaryotic supergroups, ranging from 14 to 92% (Figure 3). The observed trends for proline synthesis were similar to a previous study, which found that around 50% of bacterial genomes had a complete proline synthesis pathway, and only 40 and 30% of archaeal and eukaryotic genomes, respectively, were capable of synthesis (Mee and Wang, 2012).

As with glutamate and glutamine, proline fulfills an important role as an amino acid in proteins, but is also an osmolyte for bacteria (Burg and Ferraris, 2008; Brill et al., 2011) and some marine diatoms (Dawson et al., 2020a,b), and can protect membranes from freezing (Yancey, 2005). More recently, proline was identified as the major organic osmolyte in the chemoautotroph *Sulfurimonas* found around hydrothermal vents (Götz et al., 2018). Interestingly, proline has been found to be abundant in deep sea particulate samples (Takasu and Nagata, 2015; Johnson et al., 2021) and in sinking marine particles (Johnson et al., 2020). One possible explanation for this could be increased use of proline as an osmolyte by organisms adapted to the colder temperatures of the deep ocean.

3.2.3. Taurine

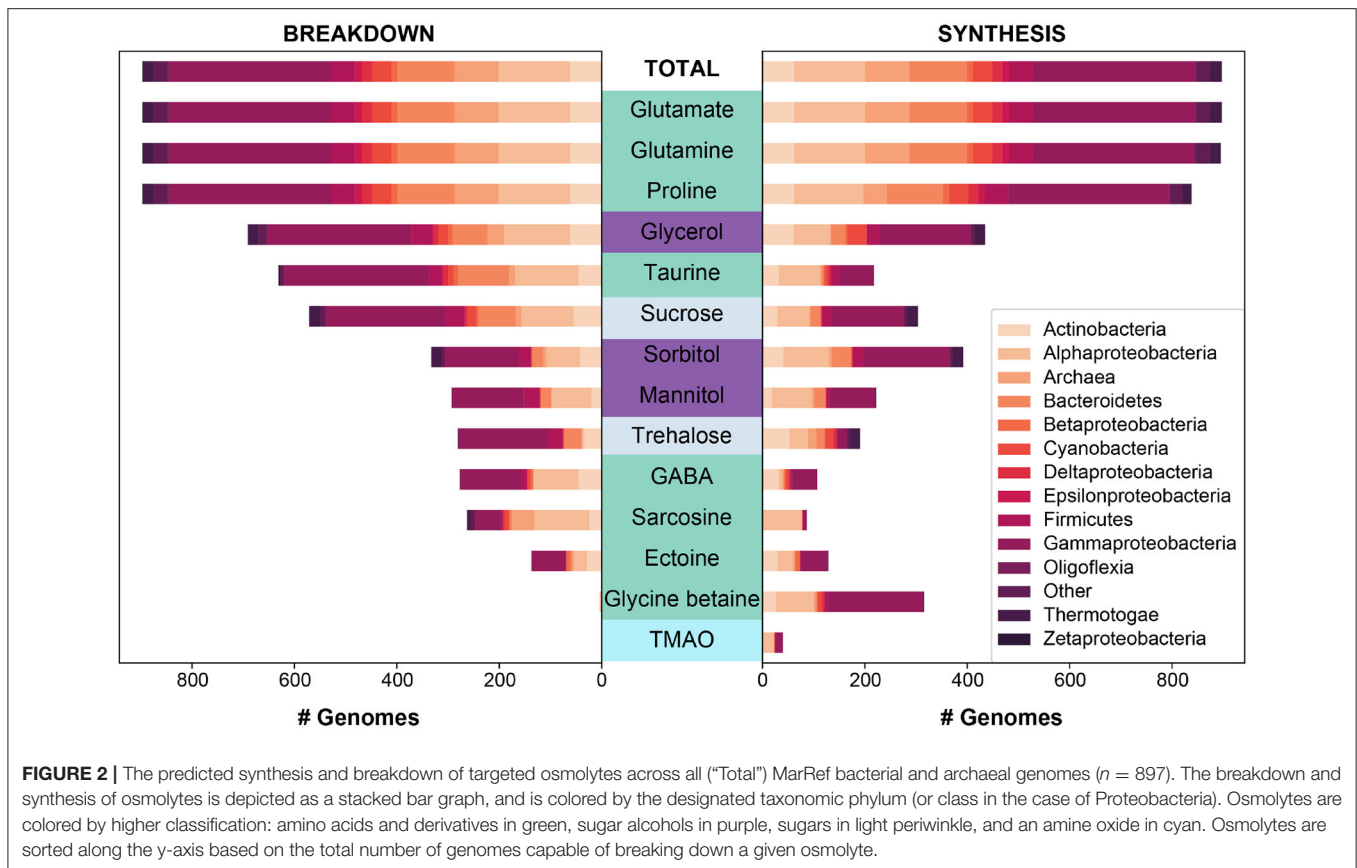
Broadly, the ability to synthesize taurine was less common than the ability to breakdown taurine across both prokaryotes and eukaryotes (Figure 1). A majority of prokaryotes were capable only of taurine breakdown ($n = 442$), and a smaller number were capable of both taurine breakdown and synthesis

($n = 189$) (Figure 1). Taurine synthesis was most common within Alphaproteobacteria (58%) and Actinobacteria (52%) (Figure 2 and Supplementary Figure 4). The breakdown of taurine in typically heterotrophic bacteria (i.e., Proteobacteria, Firmicutes, Bacteroidetes) ranged from 21 to 100% (Figure 2 and Supplementary Figure 4). Breakdown and synthesis of taurine was limited in Cyanobacteria (29% breakdown and 16% synthesis). These findings are consistent with previous work that identified bacterial uptake of taurine by three Proteobacteria groups: SAR11, Roseobacter, and *Alteromonas*, and also Thaumarchaeota and Euryarchaeota (Clifford et al., 2019, 2020). Metaproteomic data from the Ross Sea also indicated that SAR11 was a significant taurine sink via uptake and degradation (Williams et al., 2012). Prokaryotes capable of taurine transport ($n = 53$) were almost exclusively limited to Proteobacteria and were all also capable of taurine synthesis and/or breakdown (Figure 4).

Taurine synthesis and breakdown was overall less common across eukaryotes (Figures 1, 3). As observed for prokaryotes, the ability to only breakdown taurine was also most common in eukaryotes surveyed ($n = 226$) (Figure 1). Taurine synthesis was most common in Archaeplastida (61%) (Figure 3). Interestingly, Archaeplastida were far less likely to be capable of taurine breakdown (29%). This contrasts with the other eukaryotic supergroups which had limited synthesis [e.g., Stramenopiles (6%), Hacrobia, (10%), and Alveolata (41%)] and greater potential for breakdown (38, 49, and 68%, respectively) (Figure 3). Dissolved taurine concentrations in the Gulf of Alaska and the North Atlantic indicate that concentrations are typically in the low nanomolar range, and that release by amphipod-copepod assemblages in the Pacific occurs at rates of around $0.8 \mu\text{mol g}^{-1} \text{C-biomass hr}^{-1}$ and from Atlantic copepods at rates ranging from 1.3 to $9.5 \mu\text{mol g}^{-1} \text{C-biomass hr}^{-1}$ (Clifford et al., 2017). In the Adriatic Sea, even higher rates of release were found in mixed mesozooplankton communities, reaching the highest rates in the fall of an average of $59 \mu\text{mol g}^{-1} \text{C-biomass hr}^{-1}$ (Clifford et al., 2020). While our results indicate that protists are potential taurine sinks, zooplankton, specifically copepods, are clearly important taurine sources in the microbial loop. Multicellular eukaryotes (not represented in MMETSP) should be considered with respect to taurine recycling in the oceans, and possibly in addition to some other marine eukaryotes (mussels and tubeworms) that use taurine (Yin et al., 2000; Hosoi et al., 2005).

3.2.4. Glycine Betaine

Glycine betaine synthesis was more common than breakdown in both prokaryotes and eukaryotes (Figure 1). Most prokaryotes were only capable of glycine betaine synthesis ($n = 316$) (Figure 1). In particular, a majority of Alphaproteobacteria (54%) and Gammaproteobacteria (59%) were capable of synthesis. The ability to breakdown glycine betaine was extremely rare across prokaryotes (Figure 2 and Supplementary Figure 4). The small number of prokaryotes capable of breakdown ($n = 4$) were not able to synthesize glycine betaine (Figure 1) and were exclusively observed in Alphaproteobacteria strains: Candidatus *Pelagibacter ubique*, Alphaproteobacterium HIMB5, Alphaproteobacterium



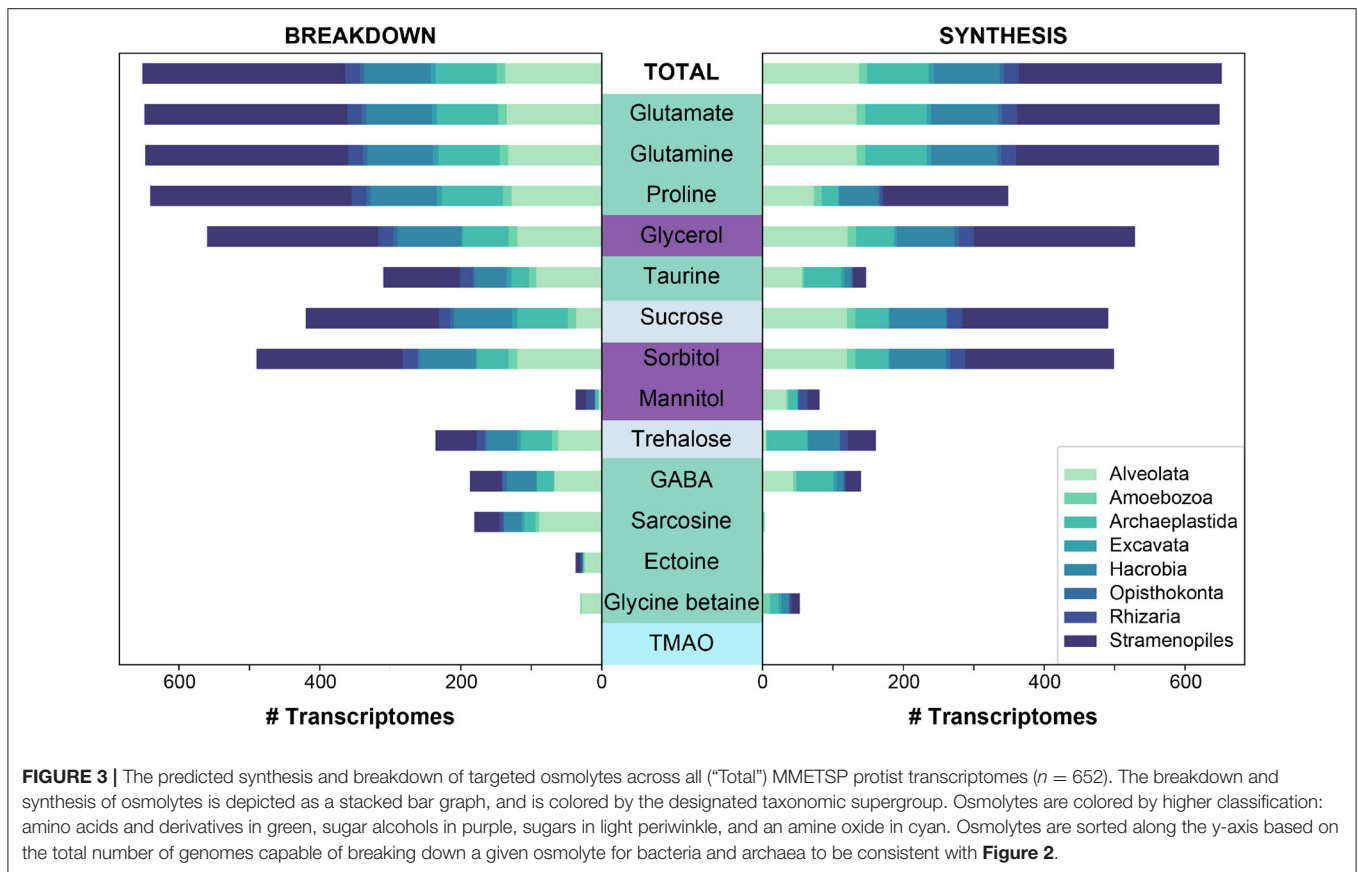
HIMB59, and *Rhodovulum sulfidophilum*. Glycine betaine synthesis and breakdown was also rare in Archaea, where only 4% were capable of synthesis and no Archaea were capable of breakdown. The ability to breakdown glycine betaine by bacteria is clearly a highly specialized metabolic ability. Interestingly, the majority of prokaryotes capable of glycine betaine transport cannot breakdown or synthesize ($n = 190$) or can only synthesize glycine betaine ($n = 234$) (Figure 4). This stands in stark contrast to our observations of taurine transporter co-occurrence with synthesis and/or breakdown, where no genomes contained only a taurine transporter (Figure 4). In addition, transport of glycine betaine was found more frequently than for other osmolytes, and the presence of the two different glycine betaine transporters was taxa dependent. Proteobacteria primarily have the osmoprotectant ABC transporter, whereas Actinobacteria primarily have the glycine betaine/proline ABC transporter. A majority of Firmicutes contained both the osmoprotectant and the glycine betaine/proline ABC transporters, and notably did not contain any other osmolyte transporters (Figure 4). The widespread ability for glycine betaine transport without breakdown suggests that this osmolyte is likely targeted by bacteria for cellular retention to function as an osmolyte, which was observed previously by direct measurements of radiolabeled glycine betaine uptake (Kiene and Hoffmann Williams, 1998). Glycine betaine was the only osmolyte in our

survey that appeared to be targeted for osmotic function by bacteria, rather than as a carbon and/or nitrogen substrate to be metabolized.

Glycine betaine synthesis and breakdown was also limited in eukaryotes (Figure 3), and the majority were only capable of synthesis ($n = 53$) (Figure 1). Notably, glycine betaine synthesis was most common in Amoebozoa (75%) and Excavata (57%), while in other groups, synthesis was < 14% prevalent (Figure 3). The breakdown of glycine betaine was absent from all groups except for Alveolata (21%) and Amoebozoa (8%) (Figure 3). Glycine betaine was previously measured in monocultures of Hacrobia (*Chrysochromulina sp.*, *Emiliania huxleyi*) and Alveolata (*Amphidinium carterae*, *Prorocentrum minimum*), and some diatoms (*Thalassiosira pseudonana*) (Keller et al., 1999a,b; Gebser and Pohnert, 2013). These direct measurements of cellular osmolytes suggest we under-predicted glycine betaine synthesis in our analysis. It is possible however, that glycine betaine synthesis was not upregulated under the conditions the reference transcriptomes were collected.

3.2.5. Sarcosine

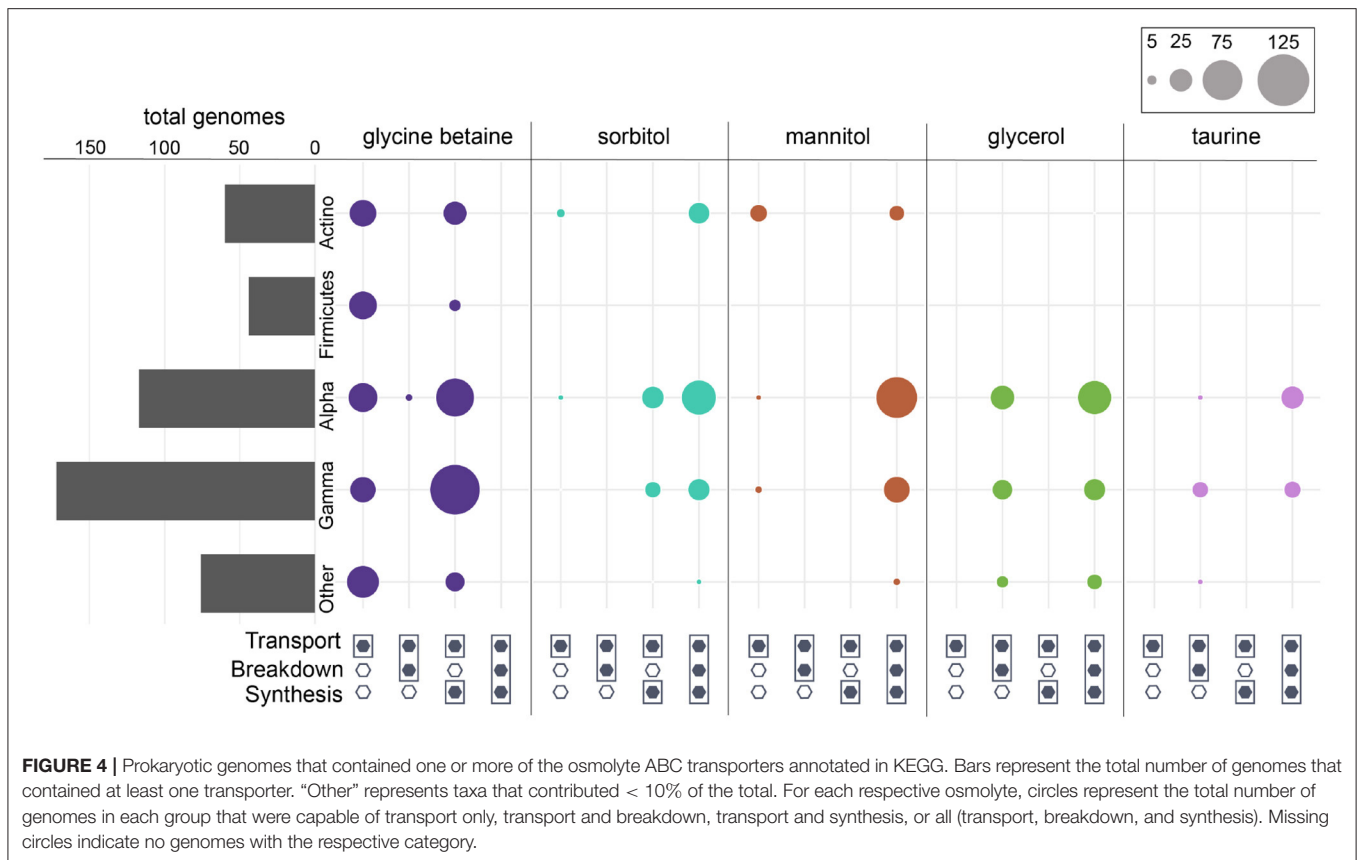
Sarcosine is structurally very similar to glycine betaine, but it lacks two methyl groups relative to glycine betaine. Sarcosine breakdown was much more common in both prokaryotes and eukaryotes (Figure 1). Sarcosine synthesis was not present in



prokaryotes, except Alphaproteobacteria of which 55% contained a synthesis pathway (**Figure 2** and **Supplementary Figure 4**). Relative to synthesis, sarcosine breakdown was far more common across prokaryotic groups, particularly in Actinobacteria, Archaea, and Alphaproteobacteria (39, 51, 77%, respectively). Sarcosine synthesis was almost completely absent in all eukaryotes (**Figure 3**). As observed for prokaryotes, more eukaryotes were capable of sarcosine breakdown, including most Amoebozoa, Excavata, and Alveolata (42, 57, 65%, respectively). While sarcosine as an osmolyte in marine organisms does not seem to have been extensively studied, elasmobranchs have been shown to use sarcosine to counteract increased levels of urea (Treberg et al., 2006), and, generally, for decades, sarcosine has been considered to act as an osmolyte (Arakawa and Timasheff, 1985). Sarcosine can be synthesized from creatine, choline, or glycine (**Supplementary Material 1**). As the ability to synthesize sarcosine was mostly absent in prokaryotic and eukaryotic groups, except in Alphaproteobacteria, it suggests that the source of sarcosine in the ocean is relatively unknown. However, sarcosine is an intermediary product formed during the breakdown of glycine betaine (glycine betaine \rightarrow dimethylglycine \rightarrow sarcosine \rightarrow glycine) (**Supplementary Material 1**), and therefore the incomplete breakdown of glycine betaine could be a possible source of sarcosine in the ocean.

3.2.6. Ectoine

Ectoine metabolism was less widely distributed across prokaryotes and eukaryotes compared to the other osmolytes (**Figures 2, 3** and **Supplementary Figure 4**). Within the small number of prokaryotes with ectoine metabolism, most were capable of both synthesis and breakdown ($n = 90$), while a smaller number were only capable of breakdown ($n = 47$) or only capable of synthesis ($n = 39$) (**Figure 1**). Ectoine synthesis and breakdown had a fairly similar taxonomic distribution within prokaryotes. For example, $\sim 48\%$ of Actinobacteria and $\sim 20\%$ of Alphaproteobacteria were capable of both synthesis and breakdown (**Figure 2** and **Supplementary Figure 4**). However, the Betaproteobacteria ($n = 12$ genomes in MarRef) had the highest proportion of synthesis (75%) and breakdown (58%). Notably, ectoine synthesis and breakdown pathways were completely absent in Cyanobacteria, and were very rare in Archaea (5% synthesis, 2% breakdown). Ectoine is known to be an important osmolyte for *Halomonas* sp. (Ono et al., 1999) and has also been found to be used across species of *Vibrio*, both those that are associated with other organisms, like fish or shellfish, or planktonic species (Pflughoeft et al., 2003; Ongagna-Yhombi and Boyd, 2013; Ma et al., 2017). Ectoine synthesis was completely absent in eukaryotes, except for one Hacrobia strain, which was likely a result of prokaryotic contamination of the transcriptome (**Figure 3**). A very small number of eukaryotes were capable



of ectoine breakdown (< 18% in all supergroups). Ectoine is typically considered to be a prokaryotic osmolyte, but was recently observed in Stramenopiles ($n = 2$ diatoms), Hacrobia ($n = 2$ haptophytes, $n = 1$ coccolithophore), and Alveolata ($n = 1$ dinoflagellate) (Fenizia et al., 2020). Specifically, the diatoms were shown to take-up ectoine in xenic cultures, and also synthesize ectoine in axenic monocultures, but only one of the three genes used by bacteria for ectoine synthesis had a putative homolog in the genome of the diatom *Phaeodactylum tricorutum* (Fenizia et al., 2020). This low homology with bacterial ectoine synthesis genes may explain why we also did not find significant ectoine synthesis in the eukaryotic transcriptomes surveyed here.

3.2.7. GABA

The synthesis and breakdown of Gamma-aminobutyric acid (GABA) was also found to be relatively rare compared to other osmolytes in both prokaryotes and eukaryotes (Figure 1). Most prokaryotes were only capable of GABA breakdown ($n = 225$), compared to a smaller number capable of synthesis only ($n = 55$), or both synthesis and breakdown ($n = 52$) (Figure 1). A majority of Actinobacteria (73%) and Alphaproteobacteria (63%) were capable of GABA breakdown (Figure 2). GABA synthesis was also common in Actinobacteria (52%), but less so in other groups (e.g., Alphaproteobacteria 4%). The ability to only breakdown GABA was most common in eukaryotes ($n = 125$) (Figure 1), but some proportion of every supergroup was capable of GABA

synthesis or breakdown, ranging from 5% to 61% synthesis and 8%–83% breakdown across supergroups (Figure 3). In particular, a large portion of Archaeplastida were capable of synthesis (57%), but breakdown was relatively constrained (27%). GABA has been identified in certain species of marine yeast at much higher concentrations than commercial yeast (Masuda et al., 2008), and is an indicator of ecosystem health in snapper (Goode et al., 2020). In marine bacteria, greater quantities of GABA are released from cells in response to decreased sodium chloride concentrations or increased pH (Mountfort and Pybus, 1992). GABA is also used as a settlement cue for marine invertebrates, and around 1/3 of bacteria isolated from a potential settlement site were also found to metabolize GABA and have high- and low-affinity transporters for the substrate (Kaspar et al., 1991). This suggests a complex role in marine environments where GABA serves as a signaling compound, osmolyte, and carbon substrate.

3.3. Amine Oxide

3.3.1. TMAO

Only one pathway for synthesis and breakdown of TMAO was identified within the KEGG framework (Table 2), which limited our analysis of TMAO synthesis, breakdown, and transport as more pathways likely exist. We identified synthesis pathways for trimethylamine N-oxide (TMAO) in a few prokaryotic groups: Actinobacteria (5%), Alphaproteobacteria (14%), Bacteroidetes (2%), Cyanobacteria (3%), and Gammaproteobacteria (5%).

However, this is likely due to limitations of the pathways annotated in KEGG. TMAO is found in deep sea animals including teleosts, skates, and crustaceans where it is thought to play an additional role as a protectant from the high hydrostatic pressure of the deep ocean (Yancey et al., 2002). Nanomolar concentrations of TMAO have been measured in Antarctic surface waters as well, indicating a role throughout the water column (Gibb and Hatton, 2004), and copepods have been shown to produce TMAO from trimethylamine (Strom, 1979).

3.4. Sugars and Sugar Alcohols

3.4.1. Sorbitol

Both the breakdown and synthesis of sorbitol were equally distributed in prokaryotes and eukaryotes, though a larger percentage of eukaryotes had the ability to breakdown and synthesize sorbitol (Figure 1). More than 40% of all Actinobacteria, Firmicutes, Alphaproteobacteria, and Gammaproteobacteria were capable of sorbitol breakdown and synthesis (Figure 2 and Supplementary Figure 4). Cyanobacteria appeared to be missing all of the single step pathways for sorbitol breakdown and sorbitol synthesis (Figure 2, Table 2 and Supplementary Figure 4). Sorbitol synthesis by prokaryotes was more common than expected as prokaryotes typically do not use polyols (e.g., sorbitol and mannitol) as compatible solutes (Kinne, 1993; Empadinhas and Costa, 2008). Both sorbose and sorbitol were previously identified as metabolites that are rapidly consumed from the dissolved pool (Vorobev et al., 2018). Here, the breakdown of sorbose to sorbitol was annotated as a sorbitol synthesis pathway, and therefore it is possible that many of the prokaryotes identified to be capable of sorbitol synthesis actually utilize this pathway to breakdown sorbose. Additionally, the majority of prokaryotes that were capable of sorbitol transport ($n = 100$) also contained sorbitol breakdown and synthesis pathways (Figure 4). Only a small number of prokaryotes ($n = 7$) contained all subunits of the sorbitol/mannitol transporter without the ability to synthesize or breakdown sorbitol. Sorbitol synthesis and breakdown was broadly and commonly distributed across the eukaryotic supergroups surveyed (Figure 3). An average $\sim 70\%$ of each eukaryotic group was also capable of sorbitol breakdown and/or synthesis, except for the Excavata (Figure 3). The widespread ability to synthesize and breakdown sorbitol in eukaryotes was expected as polyols have previously been observed to function as compatible solutes in eukaryotes (Burg and Ferraris, 2008), and sorbitol has been observed to increase with salinity in microalgae (Brown and Hellebust, 1978).

3.4.2. Mannitol

Mannitol breakdown and synthesis was less common in both prokaryotes and eukaryotes (Figure 1). Mannitol synthesis and breakdown was broadly found across all Proteobacteria ($<60\%$) and Actinobacteria ($\sim 30\%$), whereas more Firmicutes had the potential to breakdown mannitol (65%) compared to synthesis (11%) (Figure 2 and Supplementary Figure 4). Mannitol was also not expected to be widespread across prokaryotes as polyols are uncommon in prokaryotes (Empadinhas and Costa, 2008), and mannitol synthesis has been previously observed

in prokaryotes but in a limited number of taxa. Specifically, a soil Gammaproteobacterium synthesized mannitol *de novo* when other osmolyte sources were not provided exogenously (Sand et al., 2013). As observed for sorbitol, few prokaryotes ($n = 26$) were capable of sorbitol/mannitol transport only, and the majority of prokaryotes capable of mannitol transport were also able to both synthesize and breakdown mannitol ($n = 124$) (Figure 4). The presence of the transporter alongside breakdown and synthesis pathways suggests that (1) sorbitol and mannitol are not transported for osmotic function only, and (2) the transporter may actually be important for efflux with respect to internal recycling of the osmolytes. Mannitol utilization was even less common in eukaryotes (Figure 3). Eukaryotic synthesis and breakdown of mannitol was most common in Rhizaria ($> 50\%$), but not common in their potential Hacrobia symbiotic partners (1%). Mannitol was previously found to be present in Archaeplastida (Kirst, 1989), and was shown to increase with salinity in three Archaeplastida strains of Chlorophytes (Dickson and Kirst, 1987b). Relative to Rhizaria though, mannitol synthesis was limited in Archaeplastida (only 15%), though this supergroup encompasses a large diversity of eukaryotes.

3.4.3. Glycerol

Glycerol synthesis and breakdown was common across prokaryotes and eukaryotes, but the presence of breakdown and synthesis was not even in prokaryotes (Figure 1). While Actinobacteria were almost equally capable of synthesis (98%) and breakdown (100%), Proteobacteria were 2-fold more capable of breakdown (93%) than synthesis (51%) (Figure 2 and Supplementary Figure 4). In contrast, Cyanobacteria were 2-fold more capable of synthesis (92%) than breakdown (50%). Globally, 31% of prokaryotes were only capable of glycerol breakdown, lacking the ability to synthesize it (Figure 1). Prokaryotes that can transport glycerol ($n = 139$) were almost exclusively limited to Proteobacteria and were all capable of synthesis and/or breakdown of glycerol (Figure 4). In eukaryotes, 78% were capable of both synthesis and breakdown of glycerol, with the ability broadly spread across groups (Figures 1, 3). Glycerol is one of the major byproducts of photosynthesis that is often released and shared within algal-host symbioses and production has also been found to be induced under osmotic stress in model symbiotic algae such as *Symbiodinium* (Mayfield and Gates, 2007; Suescún-Bolívar et al., 2016). Moreover, glycerol has been found to increase the activity and abundance of mixotroph-associated bacteria (e.g., Alphaproteobacteria and Gammaproteobacteria), and may act as a key currency of exchange between protists and prokaryotes in the marine system (Poddar et al., 2018).

3.4.4. Sucrose

The breakdown of sucrose was at least 2-fold more common than the synthesis of sucrose in all prokaryotic groups, except for Thermotogae (Figure 1). All of the references in the Thermotogae group surveyed in MarRef ($n = 20$) are capable of both synthesis and breakdown of sucrose. This group includes known hyperthermophilic chemoorganotrophic

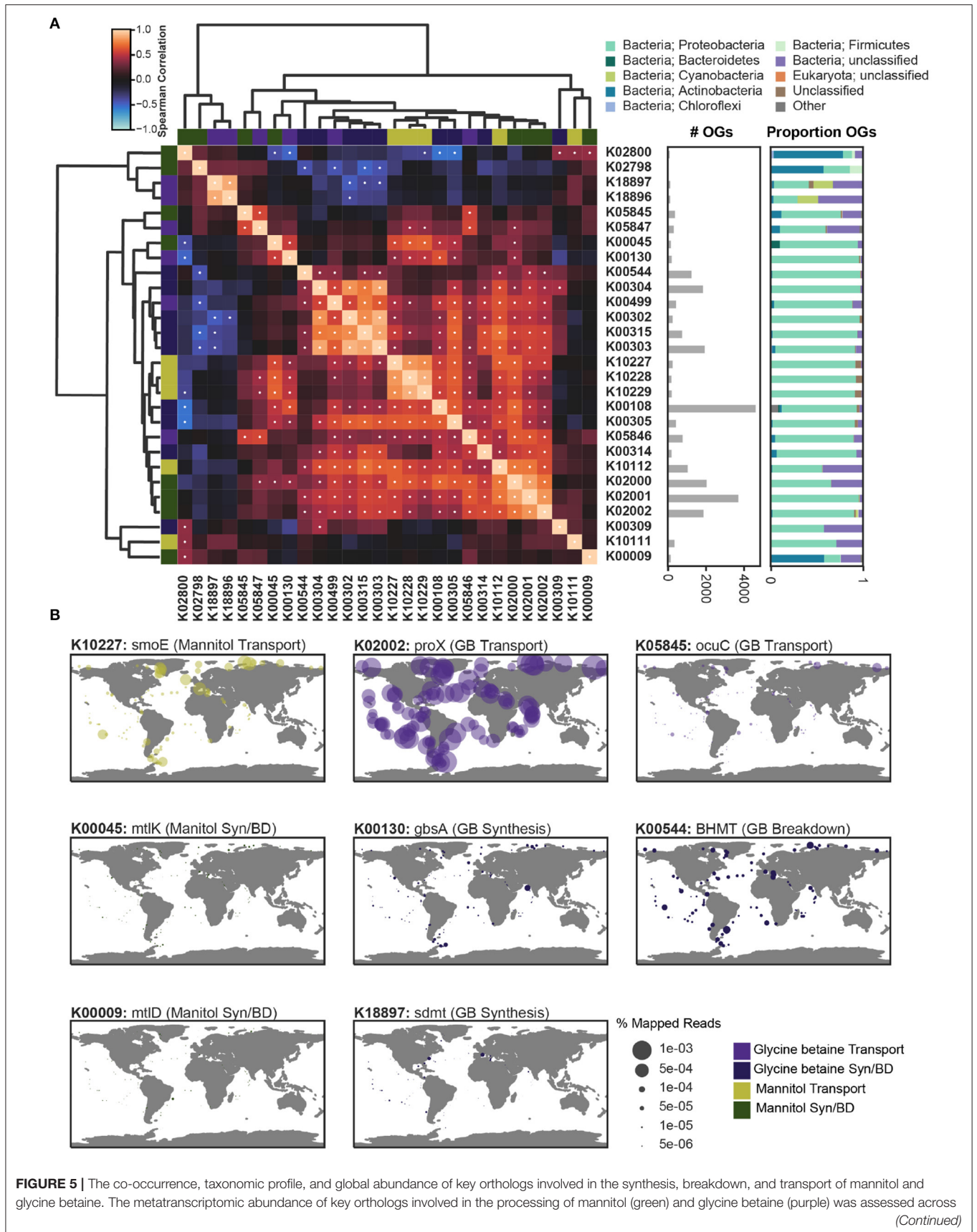


FIGURE 5 | the prokaryotic metatranscriptomic data from Tara Oceans using the OM-RGC_v2 dataset (**Supplementary Figures 5, 6**). **(A)** The relative correlation of metatranscriptomic profiles for each of the orthologs considered was assessed by taking the Spearman's correlation of the sum of all orthologs at a given site. The Spearman's ρ is depicted with a heatmap that is clustered based on Bray-Curtis similarity. Significant correlations were determined with a Bonferroni multi-testing p -value correction, and significant correlations (defined as $p < 0.001$) are depicted as a white dot. The total number of orthologs (OGs) is depicted as a bar graph and the taxonomic breakdown of the OGs is given as a proportion. **(B)** The relative abundance of key orthologs of interest is plotted as percent mapped reads across all surface samples from the Tara Oceans sampling efforts. Genes involved in mannitol metabolism include: K10111: malK, K10112: msmK, K10227: smoE, K10228: smoF, K10229: smoG, K00009: mtlD, K00045: mtlK, K02800, mtlA, K02798: cmtB. Genes involved in glycine betaine metabolism include: K02000: proV, K02001: proW, K02002: proX, K05845: opuC, K05846: opuBD, K05847: opuA, K00130: gbsA, K00499: CMO, K18896: gsmt, K18897: sdmt, K00108: betA, K00315: DMGDH, K00302: soxA, K00303: soxB, K00304: soxD, K00305: soxG, K00309: dmg, K00544: BHMT, K00314: SARDH.

organisms such as *Thermotoga maritima*, which are known to metabolize many simple and complex carbohydrates (Nelson et al., 2001). The general bias toward sucrose breakdown may suggest that prokaryotes primarily rely on other organisms for sucrose supply. Although sucrose has been shown to serve an osmotic function in Cyanobacteria and other photosynthetic organisms (Reed et al., 1986; Klähn and Hagemann, 2011), the sucrose synthesis pathway was unexpectedly incomplete based on our three-step definition (**Supplementary Material 1**) in Cyanobacteria genomes surveyed in MarRef ($n = 38$), which included *Synechococcus*, *Prochlorococcus*, known nitrogen fixers, and others. All Cyanobacteria genomes were missing the pathway's first step (conversion of glucose-1-phosphate to UDP-glucose). The majority of *Synechococcus* and *Prochlorococcus* genomes contained the glucosyltransferase required for the second step (conversion of UDP-glucose to sucrose-6-phosphate), but not the phosphatase required for the third step (sucrose-6-phosphate hydrolysis). The few other Cyanobacteria (*Calothrix* and *Nodularia*) ($n = 9$) did contain the phosphatase for the third step but were missing the enzymes for the first two steps. Sucrose utilization was common in eukaryotes ($n = 586$), and the majority were capable of both synthesis and breakdown (**Figure 1**). Amoebozoa, Hacrobia, Rhizaria, and Stramenopiles were all most frequently capable of both synthesis and breakdown (**Figure 3**). In contrast, Alveolata had 3-fold more synthesis pathways than breakdown, suggesting that sucrose may potentially be a significant osmolyte for dinoflagellates.

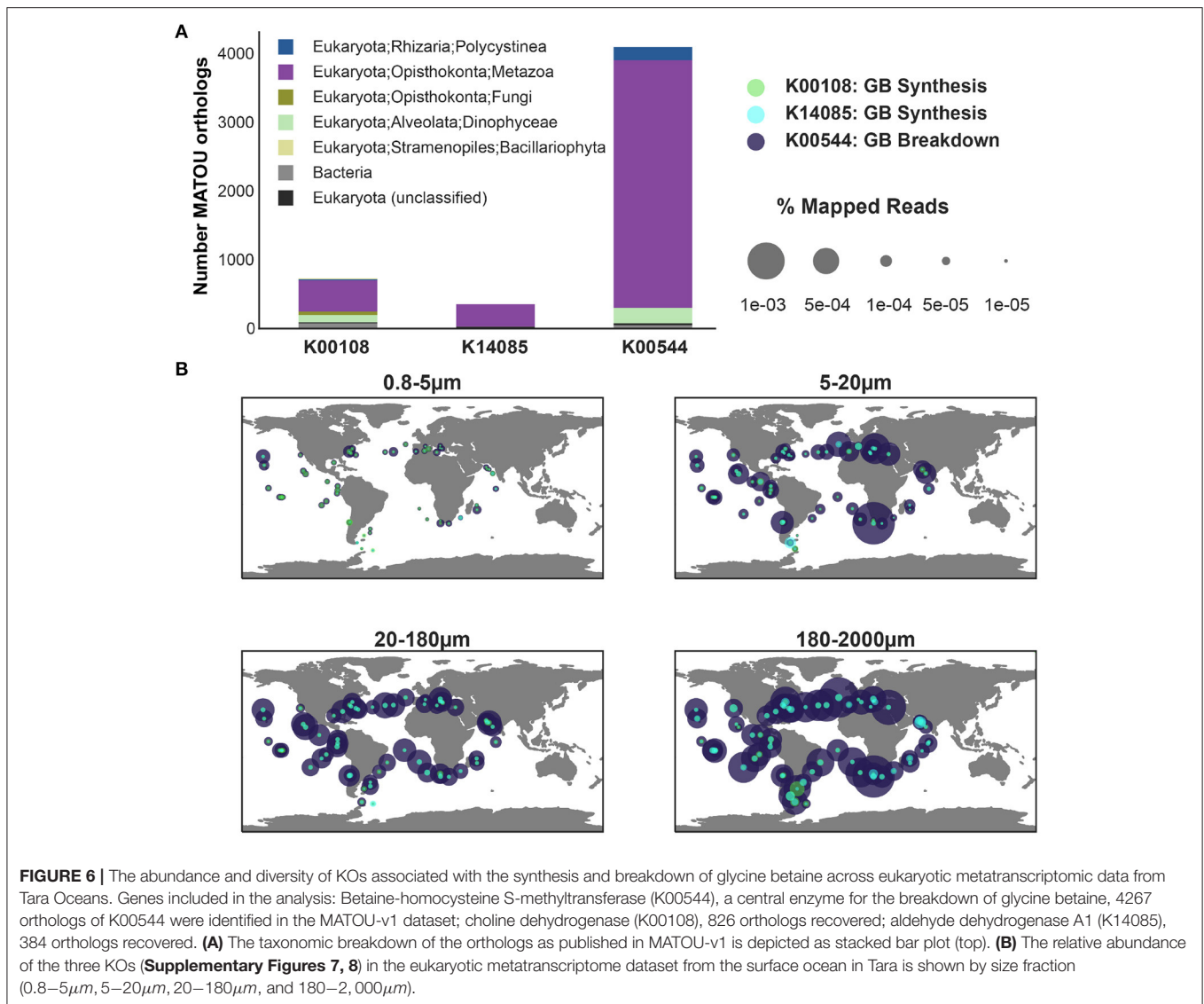
3.4.5. Trehalose

The breakdown and synthesis of trehalose exhibited unique trends across the major prokaryotic groups surveyed, where either only synthesis or only breakdown were more common than the ability to both synthesize and breakdown (**Figures 1, 2** and **Supplementary Figure 4**). This suggests that trehalose is less likely to be internally recycled, but rather it is either synthesized as an osmolyte, or consumed as a carbon source. Trehalose synthesis was more common in Actinobacteria (86%), Alphaproteobacteria (26%), Archaea (19%), and Cyanobacteria (42%), whereas trehalose breakdown was more common in Bacteroidetes (30%), Firmicutes (63%), and Gammaproteobacteria (55%) (**Figure 2** and **Supplementary Figure 4**). Interestingly, 90% of *Thermotogae* surveyed were capable of trehalose synthesis, but no trehalose breakdown. Trehalose was found to be a primary osmolyte in *Crocospaera watsonii*, instead of glucosylglycerol which is typically used by marine Cyanobacteria (Pade et al., 2012).

Many non-marine bacteria use this osmolyte, suggesting that this capability in *Crocospaera* may have been obtained through horizontal gene transfer (Pade et al., 2012). A new trehalose synthase gene that converts maltose to trehalose has also been identified in a marine species of *Pseudomonas* (Gao et al., 2013). Intracellular concentrations of trehalose have been found to fluctuate on a diel cycle in the North Pacific subtropical gyre (Boysen et al., 2020), suggesting that it might be important in the physiology of oligotrophic bacteria, such as nitrogen fixers. Trehalose breakdown relative to trehalose synthesis was at least 5-fold more common in the potentially mixotrophic or heterotrophic eukaryotic groups of Alveolata (43%), Amoebozoa (75%), and Excavata (71%). The other eukaryotic groups (Archaeplastida, Hacrobia, and Stramenopiles) were more often capable of both breakdown and synthesis (**Figures 1, 3**). Trehalose has been found to be abundant and to follow a diel cycle intracellularly in *Ostreococcus tauri* (Hirth et al., 2017). Seaweeds, and other marine plants also potentially use trehalose as an osmolyte (Xuan et al., 2012; Danaraj et al., 2020).

3.5. Metatranscriptomic Evidence of Osmolyte Cycling Across the Surface Ocean

In addition to assessing the genetic potential for osmolyte metabolism in marine microbe monocultures, the metabolism of two key osmolytes, glycine betaine, an amino acid derivative representative, and mannitol, a sugar representative, across the global ocean was assessed using the Tara prokaryotic and eukaryotic metatranscriptomic data (Carradec et al., 2018; Salazar et al., 2019). We decided to assess the metatranscriptomic signatures of genes associated with targeted osmolytes, as metatranscriptomics provides the ability to assess the transcription and potential activity of these genes across populations (Alexander et al., 2015, 2020; Hu et al., 2018; Salazar et al., 2019). The Ocean Gene Atlas (Villar et al., 2018) was used to assess (1) the diversity of *in situ* communities capable of transport, synthesis, and breakdown of each osmolyte, and (2) the geographical patterns of osmolyte cycling across the surface ocean. Representative KOs were chosen for different subpathways of glycine betaine and mannitol transport, breakdown, and synthesis, and correlations were calculated for the total abundance of orthologs across samples (**Supplementary Material 4**). Broadly, we found an alignment in the taxonomic distribution of transcripts recovered with our survey of cultivated genomes and transcriptomes (**Figures 2, 5A**). Additionally, we show that transporters were much more



highly abundant in the metatranscriptomic datasets compared to either synthesis or breakdown pathways (**Figure 5**). We also highlight the importance of metazoans in the cycling of glycine betaine (**Figure 6**).

Subunits for the different transport systems significantly correlated ($p < 0.001$) with each other across samples (**Figure 5A**). Proteobacteria dominated both the total expression and the recovered orthologs of both osmolyte transport systems. For mannitol, the substrate-binding subunit (K10227) and permease subunits (K10228, K10229) of the mannitol transport system clustered together and were significantly correlated ($\rho > 0.79$) (**Figure 5A**), suggesting that they are co-expressed across the ocean. By contrast, the ATP-binding subunit (K10111, K10112) of the mannitol transport system did not cluster with the other subunits, though one (K10112) was significantly correlated with the associated subunits ($\rho > 0.57$) (**Figure 5A**). Two different ABC transporter systems in KEGG were associated with glycine betaine. The glycine betaine/proline transporter was more common than the osmoprotectant transporter (**Figure 5A**).

All subunits (K02000, K02001, K02002) for the glycine betaine/proline transport system clustered together and were significantly correlated ($\rho > 0.72$). The ATP-binding subunit (K05847) and the substrate-binding subunit (K05845) of the osmoprotectant transporter were significantly correlated ($\rho = 0.59$). The permease subunit (K05846) for the osmoprotectant transporter did not cluster with the other subunits, but was significantly correlated with the ATP-binding and substrate-binding subunits ($\rho > 0.56$) (**Figure 5A**).

Mannitol is often recycled internally, and therefore two mannitol-associated KOs (K00045, K00009) were annotated as being capable of both mannitol breakdown and synthesis (recycling). The recycling of mannitol and fructose (K00045) was highly correlated with three subunits of the mannitol transport system ($\rho > 0.60$). Both the expression and recovered orthologs for K00045 were dominated by Proteobacteria, but also included the greatest representation of Bacteroidetes relative to all other KOs (**Figure 5A**). The recycling of mannitol and fructose-6-phosphate (K00009) was not significantly correlated

with any other orthologs, except for K02800 ($\rho = 0.41$) (Figure 5A). Both the recovered transcripts and expression of mannitol and beta-d-fructose-6-phosphate recycling, as well as the breakdown of mannitol via a phosphotransferase (K02798, K02800), were uniquely dominated by Actinobacteria. The orthologs for synthesis and breakdown of glycine betaine were overall more abundant than those for mannitol (Figure 5B). The methyltransferases involved in glycine betaine synthesis from glycine (K18896) and sarcosine (K18897) were significantly correlated ($\rho = 0.88$), and the expression of these orthologs uniquely included a significant portion of Cyanobacteria. Glycine betaine has been previously detected in Cyanobacteria (Fiore et al., 2015; Heal et al., 2020). Specifically, glycine betaine was characterized in *Synechococcus* sp. WH8102 with the two methyltransferases required for synthesis (Lu et al., 2006). The KO for glycine betaine synthesis from choline (K00108) was the most highly expressed (Supplementary Figure 5). The expression of orthologs associated with this synthesis pathway (K00499, K00108, K00130) were dominated by Proteobacteria, but were not all significantly correlated with each other. The breakdown of glycine betaine to glycine is a three-step pathway and the associated orthologs were all significantly correlated (K00315, K00302, K00303, K00304) ($\rho > 0.73$). This glycine betaine breakdown cluster was almost exclusively expressed by Proteobacteria and was significantly anti-correlated with the orthologs for glycine betaine transport ($\rho > 0.40$) (Figure 5A). Interestingly, glycine betaine breakdown was negatively correlated with richness across all samples, suggesting that this pathway is limited to a niche-group, as was observed in the prokaryotic genomes (Figure 1 and Supplementary Figure 9).

Globally, the mannitol/sorbitol transporter (K10227) was detected in most samples, but expression levels were particularly high toward the poles (Figure 5B). The latitudinal trend was supported by a significant negative correlation between temperature and mannitol/sorbitol transporter expression ($\rho = -0.66$) (Supplementary Figure 9). Salinity conditions at the poles are significantly different, and therefore may represent sites of osmolyte recycling that are unique from the majority of the open ocean (Bano et al., 2004). Both mannitol recycling genes (K00045, K00009) were expressed at comparatively low levels (Figure 5B and Supplementary Figure 6). Mannitol is generally not considered to be commonly used by prokaryotes (Empadinhas and Costa, 2008), but it has been experimentally identified as an osmolyte in several bacteria (Kets et al., 1996; Zahid et al., 2015), though not in any marine bacteria. However, marine bacteria clearly transport mannitol (and/or sorbitol) actively, while expression of the associated cellular recycling genes are orders of magnitude lower (Figure 5B). While the potential role of mannitol in marine bacteria is not entirely clear, mannitol, like other sugars and polyols, is used as a cryoprotectant in industrial applications where bacterial cells must be preserved through freezing (Savini et al., 2010). We hypothesize that the increased expression of the mannitol/sorbitol transporter in the bacterial size fraction at the poles may be linked to the availability of mannitol through production by eukaryotic phytoplankton, and its cryoprotectant properties for the bacteria.

Both glycine betaine transporters (the glycine betaine/proline transporter, K02002, and the osmoprotectant transporter, K05845) were present in bacterial transcripts. However, the glycine betaine/proline transporter was much more highly expressed throughout the surface ocean and did not show any clear differentiation between high and low latitudes or temperature (Figure 5B and Supplementary Figure 9). This expression was predominantly derived from unknown or unclassified bacteria and Proteobacteria. Previous work has shown that the Proteobacteria SAR11 have a high-affinity glycine betaine transporter (Noell and Giovannoni, 2019). *Vibrio parahaemolyticus* species contains two proU ABC transporters, but have also been found to transport glycine betaine, along with dimethylglycine, choline, ectoine, and proline with a BCCT transporter which was not linked to glycine betaine in KEGG (Gregory et al., 2020). Unlike mannitol, a glycine betaine breakdown gene (K00544) was expressed at an order of magnitude higher than the synthesis gene (K18897), and was, again, predominantly expressed by Proteobacteria (Figure 5B and Supplementary Figure 5). The glycine betaine transport and breakdown expression suggests that after transport into the cell, glycine betaine may be more commonly recycled through anabolic pathways or catabolized, rather than used as an osmolyte, at least compared to mannitol. Proteobacteria appear to be the most significant prokaryotic group metabolizing glycine betaine in Tara. Nanomolar concentrations of glycine betaine are rapidly scavenged from seawater, and most of this activity is assumed to be driven by bacteria (in the $< 1\mu\text{m}$ fraction) (Kiene and Hoffmann Williams, 1998). Radiolabeled glycine betaine was rapidly incorporated in its existing molecular structure, but after 46 h of incubation, 46% of the radioactivity was transformed into carbon dioxide (Kiene and Hoffmann Williams, 1998), indicating substantial remineralization of glycine betaine as a carbon (and possibly a nitrogen) substrate.

3.6. Zooplankton and the Cycling of Glycine Betaine

Glycine betaine synthesis and breakdown was also present in eukaryotic unigenes (Carradec et al., 2018). The taxonomy of glycine betaine synthesis (K00108, K14085) and breakdown was dominated by Arthropoda (specifically, copepods), but was also found in Vertebrata (fish) and protists (Dinophyceae and Rhizaria) (Figure 6). A similar protistan signature was observed in MMETSP transcriptomes, where glycine betaine breakdown was exclusively observed in Dinophyceae (Alveolata) (Figure 3). The observation of Rhizaria capable of glycine betaine breakdown (Figure 6) likely results from Rhizaria (Polycystinea) transcriptomes that were used to supplement the MMETSP transcriptomes as references for taxonomy assignment in the MATOU (Carradec et al., 2018). Additionally, genes for glycine betaine synthesis from glycine (K18896, K24071) in Tara were positively correlated with the Dinophyceae pigment marker ($\rho > 0.22$), peridinin, across all samples (Supplementary Figure 10). As was observed in prokaryotes (Figure 5), glycine betaine synthesis from choline was also most abundant across eukaryotes (Supplementary Figure 7). However, across all size fractions, total expression by eukaryotes was dominated by glycine betaine breakdown to glycine (Figure 6). The protist contribution to

glycine betaine breakdown expression is most likely represented in the 5 – 20 μ m, whereas the highest expression was contributed by the Arthropoda and Vertebrata in the 180 – 2000 μ m fraction.

Copepod excretion is a potentially important source of labile substrates (Maas et al., 2020). Previous studies have described taurine as an important osmolyte in marine copepods (Clifford et al., 2020). Although glycine betaine has been described across all three domains of life (Yancey, 2005), to our knowledge, the potential for glycine betaine synthesis and breakdown has not been described previously for marine planktonic Arthropoda. The expression patterns described here for eukaryotic unigenes hint at an important role for not only protists, but also higher-order marine eukaryotes in the recycling of this osmolyte. The disconnect between breakdown and synthesis across all eukaryotic size fractions suggests that eukaryotes are capable of obtaining glycine betaine from the environment. In the case of multicellular organisms, glycine betaine could be obtained through grazing or the microbiome (Shoemaker and Moisaner, 2017). The dominance of breakdown could in part be due to the conservative threshold used here to filter ortholog hits causing a higher recovery of breakdown KOs and/or removal of synthesis genes. The expression of both synthesis and breakdown of glycine betaine in the 180 – 2000 μ m fraction suggests that multicellular organisms, particularly copepods, could be an important source or sink of this osmolyte that is not currently considered.

4. CONCLUSIONS

Osmolytes are core cellular metabolites that are required for all marine microbes living in the high salinity environment of the ocean. Across the three domains of life, we found that osmolyte metabolic capabilities could be broken down into two main categories: (1) globally ubiquitous, and (2) mosaically present across groups. The synthesis and breakdown of three amino acid osmolytes surveyed (glutamate, glutamine, and proline) were common across all prokaryotes and eukaryotes. The ubiquity of synthesis and breakdown of these core metabolites likely indicates an important role for internal recycling. Although it is difficult to directly attribute osmotic function without experimental evidence, we hypothesized that osmolytes in genomes with synthesis only, or genomes with transport only were more likely used for osmotic functions. The genomic evidence for a potential osmotic function (synthesis or transport in the absence of breakdown) was most apparent for the classically known osmolyte, glycine betaine, which appeared to be a unique instance of osmolyte specificity in the absence of central metabolic importance. The dominance of synthesis or breakdown of the other osmolytes surveyed differed significantly across prokaryotes and eukaryotes, suggesting that osmolyte sources and sinks are taxa dependent. Critically, the ability to breakdown was most common relative to synthesis or transport for the majority of the osmolytes surveyed, suggesting that these small molecules serve a central metabolic function and are likely important substrates in the microbial loop.

We took a two pronged approach in our survey of osmolyte metabolism within marine taxa: (1) examining metabolic

potential within reference genomes and transcriptomes and (2) assessing transcriptional activity from mixed communities across the global ocean. When we compared the potential of taxonomic groups to synthesize, breakdown, and transport osmolytes to the expression of the genes associated with these pathways in the environmental metatranscriptomic data, we found good agreement between the species with the potential to perform these metabolic functions, and those that were observed to transcribe the relevant genes in the environment. In the environment, bacterial transcription of transporter genes for both mannitol and glycine betaine were an order of magnitude higher than their respective genes for synthesis and breakdown. Although only four Alphaproteobacteria genomes were found to have the potential for glycine betaine breakdown, the relative abundance of transcription of the breakdown pathway was significantly higher than the breakdown pathway for mannitol. One of the strains capable of breakdown of glycine betaine, *Candidatus Pelagibacter ubique*, is a highly abundant taxa found broadly throughout the oligotrophic ocean (Morris et al., 2002), which perhaps explains the high level of transcription. This suggests that a niche based on glycine betaine as a carbon source may be occupied by these species without competition from other bacterial species. Interestingly, although synthesis of glycine betaine by eukaryotic size fractions was widespread, breakdown was most highly expressed in the larger size fractions, particularly by metazoa. Thus, not only are protists likely important sinks of the osmolyte glycine betaine, but also higher-order, multicellular eukaryotes should also be considered as potential sinks. This finding would not have been possible without probing the Tara metatranscriptomes. Our initial investigation of two key osmolytes within the Tara metatranscriptomes highlights the fact that both eukaryotes and prokaryotes are actively using and recycling osmolytes *in situ*, and that osmolytes are potentially being exchanged between pelagic bacteria, phytoplankton, and also zooplankton. Marine protists, particularly phytoplankton, are most often emphasized as potential sources of marine osmolytes, but clearly connections with higher order trophic levels are also important in the global cycling of carbon and should be examined more in depth (Durham et al., 2019; Clifford et al., 2020).

The ocean is a dynamic and variable environment, and the organisms that inhabit it must be prepared to survive sudden environmental fluctuations. Here we surveyed and reviewed the osmoadaptive capabilities in the form of the synthesis, breakdown, and transport of various osmolytes across cultured organisms with sequenced genomes or transcriptomes as well as within a global metatranscriptomic dataset. We believe that the work here serves as a first step in the targeted and informed examination of osmoregulation by ecologically-relevant groups in the microbial loop. In particular, further work examining the presence and absence of these genes across metagenome assembled genomes (MAGs) or single celled amplified genomes (SAGs) may provide greater insights into the osmoadaptive capabilities of uncultivated lineages. Additionally, examining rates of uptake and loss of osmolytes from cells, coupled with more comprehensive characterization of their dissolved concentrations in the ocean, would support our understanding

of their potentially significant influence on labile carbon cycling in the ocean. Broadly, we show that while some osmolytes (e.g., the amino acids glutamate and glutamine) are ubiquitously synthesized and broken down across the tree of life, the ability to synthesize, breakdown, and transport most osmolytes is present mosaically across organisms. The mosaicism of the genomic traits of osmolyte metabolism may suggest that many of the osmolytes surveyed here are central currencies of exchange between organisms (Moran et al., 2016), between domains or within domains.

DATA AVAILABILITY STATEMENT

Publicly available datasets were analyzed in this study. They can be found in the following repositories: MMETSP (<https://zenodo.org/record/1212585>), Tara Oceans Gene Atlas (<https://tara-oceans.mio.osupytheas.fr/ocean-gene-atlas/>), KEGG homologs (<https://www.genome.jp/tools/kofamkoala/>), and MarRef v5 (<https://mmp.sfb.uit.no/downloads/>). Accession numbers are listed in **Supplementary Material 2 and 3**.

AUTHOR CONTRIBUTIONS

All authors contributed equally to the development of this project, analysis of the data, and writing of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fmars.2021.689306/full#supplementary-material>

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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