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Glycine betaine uptake and metabolism in marine microbial communities

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Summary

Glycine betaine (GBT) is a compatible solute in high concentrations in marine microorganisms. As a component of labile organic matter, GBT has complex biochemical potential as a substrate for microbial use that is unconstrained in the environment. Here we determine the uptake kinetics and metabolic fate of GBT in two natural microbial communities in the North Pacific characterized by different nitrate concentrations. Dissolved GBT had maximum uptake rates of 0.36 and 0.56 nM h^{-1} with half-saturation constants of 79 and 11 nM in the high nitrate and low nitrate stations respectively. During multiday incubations, most GBT taken into cells was retained as a compatible solute. Stable isotopes derived from the added GBT were also observed in other metabolites, including choline, carnitine and sarcosine, suggesting that GBT was used for biosynthesis and for catabolism to pyruvate and ammonium. Where nitrate was scarce, GBT was primarily metabolized via demethylation to glycine. Gene transcript data were consistent with SAR11 using GBT as a source of methyl groups to fuel the methionine cycle. Where nitrate concentrations were higher, more GBT was partitioned for lipid biosynthesis by both bacteria and eukaryotic phytoplankton. Our data highlight unexpected metabolic pathways and potential routes of microbial metabolite exchange.

Introduction

In the ocean, the flow of resources and information carried by organic molecules provides the foundation for complex and globally important microbial ecosystems. The quantity and identity of molecules produced by phytoplankton help shape community structure and influence the balance between organic carbon production and degradation, which together influence carbon storage in the ocean (Gómez-Consarnau et al., 2012; Moran, 2015). Small, polar metabolites released into seawater are exchanged and used by cells in a variety of ways (Poretsky et al., 2010; Vorobev et al., 2018). A molecule might be incorporated directly into biomass as a building block for macromolecules, respired for energy, converted into biosynthetic intermediates, or used for specific biochemical roles as osmolytes, cofactors, cellular signals, antioxidants, or chemical defences. Here we focus on a single compound, the compatible solute glycine betaine (GBT). We elucidate the uptake kinetics and metabolic fate of GBT in two natural microbial communities with different nitrate concentrations.

Once released into seawater through excretion or cell death, compatible solutes are a quantitatively important pool of labile dissolved organic matter (DOM) (Bronk et al., 1994; Welsh, 2000; Bronk, 2002). GBT, a guaternary amine, is one of the most effective compatible solutes and is found in high concentrations inside cells in saline environments (Airs and Archer, 2010; Klähn and Hagemann, 2011; Johnson et al., 2019). GBT comprises a significant portion of intracellular metabolites in the North Pacific subtropical gyre and in the North Pacific transition zone (NPTZ) (Heal et al., 2021) and exhibits diel oscillations indicative of active production and consumption (Boysen et al., 2021). Incubation experiments with ¹⁴C-GBT show that natural microbial communities both respire and assimilate carbon from GBT (Kiene et al., 1998; Kiene and Williams, 1998; Mausz et al., 2022). However, GBT has complex biochemical potential, and previous work has left the flow and fate for GBT-derived carbon and nitrogen mostly unresolved. Furthermore, few studies have investigated the role that

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environmental conditions have on the biochemical fate of GBT.

Across vast swaths of the ocean, inorganic nitrogen availabilitv limits phytoplankton arowth (Moore et al., 2013). In inorganic nitrogen-depleted waters, dissolved organic nitrogen (DON) is the dominant form of nitrogen, sometimes comprising 100% of measurable total dissolved nitrogen (Berman and Bronk, 2003). Microorganisms that can take up and use DON have an advantage in these environments. However, the specific structures of molecules within the DON pool determine which organisms can take it up and how they can use it. Incubations of DOM amended seawater show that GBT transporters are transcribed in response to DOM additions, presumably to take up GBT (Poretsky et al., 2010; Vorobev et al., 2018), but these studies did not directly measure uptake or how GBT was subsequently used.

There are three pathways of GBT metabolism with wellcharacterized enzymes (Fig. 1 pathways 1–3). The demethylation pathway removes methyl groups from GBT one at a time (1 in Fig. 1). Betaine-homocysteine methyltransferase (BHMT) converts GBT and homocysteine to dimethylglycine and methionine, which can fuel methylation throughout the cell via the methionine cycle (Barra *et al.*, 2006). Alternatively, the genes *gbcA* and *gbcB*

encode a Rieske-family oxidase and flavin-linked oxidoreductase that convert GBT to dimethylglycine with the production of formaldehvde (Wargo et al., 2008: Shao et al., 2018). Further oxidation of dimethylglycine to sarcosine and glycine occurs through subsequent demethylations in which methyl groups are transferred to tetrahydrofolate and ultimately oxidized to CO2 in an energy-yielding reaction (Sun et al., 2011). The first of these demethylations is catalysed by either dimethylglycine dehydrogenase (DMGDH) or dimethylglycine oxidase (DMGO) and then the second by sarcosine dehydrogenase (SARDH), monomeric sarcosine oxidase (MSOX), or tetrameric sarcosine oxidase (SoxBDAG) (Wagner et al., 2000; Wagner and Jorns, 2000; Wargo et al., 2008; Lahham et al., 2021). Organisms with the full demethylation pathway, including the ubiquitous marine Alphaproteobacteria clade SAR11, can convert GBT to glycine and use the glycine for respiration, biosynthesis and protein synthesis (Sun et al., 2011; Wargo, 2013; Noell and Giovannoni, 2019). Through this pathway organisms can use GBT to fulfil cellular methylation, glycine, carbon, nitrogen and energy requirements for growth (Lidbury et al., 2015a).

The second pathway with characterized enzymes is the transformation of GBT to choline (2 in Fig. 1). A nonribosomal peptide synthetase-like glycine betaine reductase (ATRR) responsible for this transformation was



Fig. 1. Schematic of formation and degradation pathways for GBT. Circled numbers are GBT metabolism pathways referenced in the text. Solid black arrows indicate pathways of GBT synthesis or degradation that have one or more characterized enzymes. Grey dotted arrows represent select potential fates of GBT-related molecules. Enzyme names are abbreviated as follows: BHMT = betaine-homocysteine methyltransferase; GbcAB = glycine betaine demethylase; DMGO = dimethylglycine oxidase; DMGDH = dimethylglycine dehydrogenase; SARDH = sarcosine dehydrogenase; MSOX = monomeric sarcosine oxidase; SoxBDAG = tetrameric sarcosine oxidase; GSMT = glycine-sarcosine methyltransferase; BSMB = dimethylglycine N-methyltransferase; GbsAB, BetAB, CHDH = choline dehydrogenase; CMO = choline monooxygenase; CodA = choline oxidase; ATRR = non-ribosomal peptide synthetase-like glycine betaine reductase; CDH = carnitine dehydrogenase; BKACE = β -keto acid cleavage enzyme; GrdHI = betaine reductase.

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identified in fungi (Hai et al., 2019), though the presence of this pathway in marine microbes is unexplored. The third characterized pathway of GBT metabolism is GBT reduction to trimethylamine (TMA) via betaine reductase (GrdHI) (3 in Fig. 1). TMA can then fuel anaerobic methanogenesis or be oxidized to trimethylamine-N-oxide (TMAO) (Sun et al., 2019). TMA and TMAO can both be demethylated to ultimately form formaldehyde and ammonium, which can feed back into central carbon and nitrogen metabolism (Sun et al., 2019). Betaine reductase is considered an anaerobic pathway, but ardH homologues have been identified in proteobacteria from open ocean marine metagenomes (Jameson et al., 2016). A fourth GBT metabolism pathway that produces carnitine through GBT-CoA and dehydrocarnitine (4 in Fig. 1) is possible since the enzymes that produce GBT from carnitine [CDH and β -keto acid cleavage enzyme (BKACE)] are theoretically reversible (Sellés Vidal et al., 2018).

Cyanobacteria, heterotrophic bacteria and eukaryotic algae have a variety of abilities to synthesize, transport and metabolize GBT (Keller et al., 1999a; Welsh, 2000; Mou et al., 2007; Torstensson et al., 2019; McParland et al., 2021). As described above, some heterotrophic bacteria can transport and metabolize GBT. Some diatoms, dinoflagellates and haptophytes that produce GBT can take up extracellular GBT under salinity stress or when extracellular concentrations of GBT are elevated, but there is no evidence that these organisms metabolize GBT (Keller et al., 1999b; Torstensson et al., 2019). There are also marine microorganisms that lack both the known synthesis and degradation genes for GBT but still have GBT transporters (McParland et al., 2021). Thus, there are several potential fates for extracellular GBT once taken up by marine microbes. It could be retained as a compatible solute, catabolized to provide cellular energy, used as a methyl donor (1 in Fig. 1), used for biomolecule synthesis including lipids (through choline, 2 in Fig. 1) and proteins and nucleobases (through glycine), or used for the synthesis of other metabolites (3 in Fig. 1). Intermediates along these pathways may also be exchanged between organisms, implicating GBT as a foundation for complex microbial interactions.

Here we explore the fate of GBT in marine surface waters using transcriptomics and stable isotope tracing untargeted metabolomics. We incubated natural seawater collected at two stations in the NPTZ with $^{13}C_5$, $^{15}N_1$ -GBT to measure the kinetics of GBT uptake and determine the fate of the carbon and nitrogen. The NPTZ is a dynamic region characterized by seasonally migrating fronts of temperature, salinity, nutrients and chlorophyll (Ayers and Lozier, 2010; Polovina *et al.*, 2017), and thus a useful region to investigate the effects of environmental conditions on GBT uptake and metabolism. We show that

microbial communities are poised to take up and retain high concentrations of GBT. Once GBT is in cells, we show that the proportion of carbon and nitrogen that flow through each GBT pathway differs between the two sites. The observed differences are consistent with more intense recycling of photosynthate, and, in particular, the nitrogen in GBT, in the southern, more oligotrophic, environment.

Results and discussion

Environmental description

Experiments to determine the uptake kinetics and cellular fate of GBT were performed in spring 2019 at two stations in the NPTZ. The northern station (41.68 °N, -158 °W) was characterized by 0.85 \pm 0.08 µg chlorophyll L⁻¹ and 6.42 µM dissolved inorganic nitrogen (DIN, NO₃⁻ +-NO₂⁻). The southern station (37 °N, -158 °W) was characterized by 0.57 \pm 0.03 µg chlorophyll L⁻¹ and DIN below our limit of detection (near 0.02 µM, Table 1). There was more particulate carbon, heterotrophic bacterial biomass and picophytoplankton biomass in the southern station (Table 1). The elevated chlorophyll in the northern station was likely due to larger phytoplankton (>4 µm diameter) outside the analytical window of the flow cytometer.

Glycine betaine uptake kinetics

GBT uptake was well described by Michaelis-Menten kinetics at both locations (Fig. 2 top row). A non-linear least-squares fit was used to determine two uptake parameters: maximum uptake rate, V_{max}, and half saturation concentration plus in situ GBT concentration, $K_t + S$. The southern station had a $V_{\rm max}$ of 0.56 \pm 0.06 nM h⁻¹ and K_t + S of 11.06 \pm 5.53 nM. The northern station had lower $V_{\rm max}$ of 0.36 \pm 0.06 nM h⁻¹ and higher K_t + S of 79.46 ± 48.26 nM, indicating a lower capacity to take up GBT at low concentrations (Table 2, Supplemental Table S1). The GBT uptake kinetic parameters measured here are within the wide range of published values for amino acids, the DON compounds best described in the literature (Sipler and Bronk, 2015). The half saturation constants of GBT uptake that we estimate are on the high end of those previously measured for free-living bacteria which had $K_t + S$ values of <1 to 17.2 nM, (Kiene et al., 1998; Mausz et al., 2022) and for whole seawater communities in coastal environments (1.8-49 nM for waters with salinity >10 ppt) (Kiene and Williams, 1998). We estimated a turnover time for dissolved GBT of 8.7-37.5 h in the southern station and >100 h in the northern station (Table 2). Compared to the <1-11 h GBT turnover time of coastal and estuarine environments (Kiene

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rable 1. Summary of the environmental conditions where samples were collected in this study

et al., 1998; Kiene and Williams, 1998), these data suggest that open ocean communities are slower to turn over the available pool of GBT.

The presence of more transporters for a given substrate is one way to increase the maximum uptake rate, $V_{\rm max}$, measured at the community scale, and thus biomass is one of the parameters that can directly influence $V_{\rm max}$ (Button, 1998). At the community scale, a low half saturation constant, K_t , is likely due to high-affinity transporters making up a substantial proportion of total transporters, regardless of transporter type. Though the community level K_t value does not necessarily reflect any individual organism or transporter K_t value, it does inform us about the general affinity level that transporters have for a substrate (Kiene et al., 1998). Free-living heterotrophic bacteria were more abundant in the southern station than the northern station (Table 1) and likely contributed to the high V_{max} and low $K_t + S$ values which were near the SAR11 GBT transporter K_t values of 0.89 and 1.85 nM (Noell and Giovannoni, 2019). The microbial community in the northern station had more large eukaryotic phytoplankton, which contributed to the high chlorophyll levels at that location (Table 1). The only characterized GBT transporters in phytoplankton are in sea-ice diatoms, which have Kt values of 189-315 nM (Torstensson et al., 2019) that are near the value measured at the northern station. The high community K_t value in the north indicates that more of the organisms taking up GBT have lower affinity transporters, and we suggest that phytoplankton are substantially contributing to uptake at that station. Our results support the hypothesis that community composition, bacterial abundance and nutrient concentrations influence GBT uptake kinetics (Mausz et al., 2022).

Glycine betaine over time

To assess the fate of GBT once it was taken up into cells, we added $^{13}C_5$, $^{15}N_1$ -GBT to natural seawater communities in excess of the half saturation constants (500 nM) and traced carbon and nitrogen isotopes through metabolic pathways over several days. Samples for the initial time point were filtered directly after the addition of labelled GBT, resulting in an effective incubation time of 15–30 min. Though we expect ambient dissolved GBT concentrations to be lower than 500 nM, these concentrations may be reached in microenvironments such as the phycosphere, where nutrients are exchanged between phytoplankton and their associated bacteria (Stocker, 2012).

In the first 4–6 h, the total particulate GBT concentrations increased from 0.60 to 1.03 nM in the south and from 0.61 to 0.84 nM in the north (Fig. 2 bottom row). Though the particulate GBT concentrations continued to

dard deviation.



Fig. 2. Top row: Uptake of ${}^{13}C_{5}$, ${}^{15}N$ -GBT in natural seawater. Black points are rates calculated from individual bottles, standard error propagated from the uncertainty associated with quantification is shown as the bars, sometimes obscured by the size of the points. Open circles are replicate injections of the same sample. Uptake kinetics curve fit to the data with a non-linear least squares and Monte Carlo randomization, as described in the Experimental Procedures section, is shown in the solid line, with one standard deviation of the best fit $K_t + S$ and V_{max} parameters shown in shading. Dashed vertical line is the best fit $K_t + S$ value. Inset shows the shape of the curve for the 0–200 nM treatments. Bottom row: Particulate GBT over time in the incubation experiments with 500 nM ${}^{13}C_5$, ${}^{15}N$ -GBT. The three most abundant isotopologues are shown here. Others were measured and are shown in Supplemental Fig. S2. Solid lines indicate the mean of triplicate bottles and the error bars represent one stan-

increase over the course of the experiments due to uptake of the added GBT, the composition of the compatible solute pool overall did not change substantially, since homarine dominated the total compatible solute pool at both stations at all time points (Supplemental Fig. S1). Though particulate GBT concentrations continued to increase in the last 2 days of the northern incubation, the fraction of total particulate GBT that was in the $^{13}C_5$, $^{15}N_1$ -GBT isotope form (a value represented by the mass isotopologue distribution, MID) stabilized after 2 days of incubation. $^{13}C_5$, $^{15}N_1$ -GBT, the isotopologue of GBT that was added, reached approximately 85% of the total particulate GBT at both locations (Supplemental Fig. S2).

By searching metabolomics data for compounds enriched in ¹³C and ¹⁵N we were able to identify multiple active pathways of GBT transformation. Unmetabolized GBT dominated ¹³C and ¹⁵N isotopes in aqueous intracellular metabolites (Fig. 3, Supplemental Fig. S3). In the final time point of the northern experiment, GBT isotopologues occupied 85 \pm 19% of ¹⁵N and 80 \pm 17% of ¹³C within the quantifiable particulate metabolites. In

the final time point of the southern experiment, GBT isotopologues occupied 88 \pm 7% of ¹⁵N and 86 \pm 7% of ¹³C within the quantifiable particulate metabolites. Homarine represented a substantial portion of the total ¹³C pool at all time points due to its high concentration and the natural abundance of ¹³C, not due to enrichment from GBT (Supplemental Fig. S1). We underestimate the total isotope uptake and transformation because our analytical window does not allow us to observe every metabolite, not all metabolites we observe can be quantified, and we did not measure metabolic products, including carbon dioxide and ammonium, in the dissolved pool. Nevertheless, our data indicate that a substantial portion of total GBT taken up is retained as a compatible solute.

Our observations contrast studies in coastal environments where 48-h incubations of ¹⁴C-GBT showed that much of the carbon taken up as GBT was incorporated into non-soluble molecules interpreted to be macromolecules (Kiene and Williams, 1998). These previous studies focused on the <0.8 μ m size fraction and concluded that most GBT uptake and metabolism occurred within the

Compound	Location	V_{max} (nmol L ⁻¹ h ⁻¹) ^a	Kt + S (nM) ^a	Turnover time (h) ^a	References
Glycine betaine	N. Pacific (41 N)	0.3–0.42	31.2– 127.72	106–169	This study
Glycine betaine	N. Pacific (37 N)	0.5–0.63	4.81–21.49	8.7–37.5	This study
Glycine betaine	Mobile Bay	0.39–44	1.8–49	0.52–11	Kiene and Williams (1998)
Glycine betaine	Mobile Bay, 0.7 μ m filtrate	2.4–27	0.86–4.5	0.15–1.8	Kiene <i>et al</i> . (1998)
Glycine betaine	Western English Channel, 1.2 µm filtrate	0.6–40.8	1.4–17.2	0.19–6.2	Mausz et al. (2022)
DMSP	Mobile Bay, 0.7 μm filtrate	8.94	3.77		Kiene <i>et al</i> . (1998)
DMSP	NPSG			4.1-7.4	Del Valle et al. (2012)
Choline	Mobile Bay estuarine water	0.5–3.3	1.7–4.1	1.3-3.2	Kiene et al. (1998)
Choline	Western English Channel, 1.2 µm filtrate	0.2–3.4	1.9–5.3	<1–25	Mausz et al. (2022)
Individual DFAA	Sargasso Sea (N. Atlantic)	0.003–0.2	0.1–5.5	1.6–181.9	Suttle et al. (1991)
Alanine	Sargasso Sea (N. Atlantic)	0.015-0.15	0.9–1.8	1.6-13.2	Suttle et al. (1991)
Ornithine	Sargasso Sea (N. Atlantic)			48.7-181.9	Suttle et al. (1991)
Taurine	Off the Iberian Peninsula			6–240	Clifford et al. (2019)
Leucine	Off the Iberian Peninsula			21–6000	Clifford et al. (2019)

Table 2. Summary of uptake kinetic parameters for various metabolites in the marine environment.

^aRanges represent the range of the values in the publication or the best estimate plus and minus one standard deviation, the values for this study include the full range of estimates from the linear transformation and non-linear least squares fit.

bacterial community. We hypothesize that the uptake and retention of GBT, which was greater in the northern experiment than the south experiment, is driven in part by eukaryotic phytoplankton uptake. Cultures of diatoms, haptophytes and dinoflagellates take up GBT when provided with exogenous GBT (Keller et al., 1999b; Torstensson et al., 2019) but lack the known genes for GBT demethylation (McParland et al., 2021). Taking up and retaining compatible solutes has a cost associated with maintaining membrane transporters, but those costs may well be offset by saving the energy, carbon and nitrogen required to synthesize compatible solutes, especially if the transporter can be used for multiple substrates as is the case for some characterized GBT transporters (Kiene et al., 1998; Noell and Giovannoni, 2019). Excess uptake and storage of GBT may be a strategy for surviving patchy and nutrient-limited regions of the ocean. Similar luxury uptake strategies have been proposed for inorganic nutrients and other organic nutrients, including uptake of polyamines by SAR11 (Gaubert et al., 2020; Noell et al., 2021). Future studies of organisms in culture should explore this potential use of GBT and other nitrogen-containing osmolytes as a mechanism for increased fitness in low and fluctuating nitrogen regimes.

Primary pathways of GBT transformation

Isotope tracing metabolomics revealed four potential pathways of GBT metabolism in both incubation

experiments, as summarized in Fig. 1: (i) the demethylation pathway into dimethylglycine, sarcosine, glycine and creatine, (ii) the choline lipid pathway into choline and glycerophosphocholine, (iii) the reduction pathway into TMA and TMAO and (iv) the carnitine synthesis pathway into carnitine and trimethylammoniobutanoate (TMAB). In our environment, the third and fourth potential pathways may be a single pathway, since TMAO can also be produced from carnitine, rather than from GBT directly, as discussed below.

Our stable isotope probing experimental results led us to search for which organisms might be responsible for these various transformations bv searching metatranscriptomes collected from the NPTZ on a cruise in April-May 2016. Unless otherwise specified, we present the gene transcription data as an average of the five stations in the NPTZ, as defined by the second derivative of salinity, following Gradoville et al. (2020). We use the regional average because the stations where metatranscriptomes were collected were not concurrently sampled with those used for the GBT experiments. This analysis reveals which organisms within this region are likely carrying out the observed metabolisms in situ, and thus lets us infer how various microbial community members may rely on GBT and the immediate products of its metabolism. Metatranscriptomes collected from the same region in May-June 2017 suggest that the organisms expressing GBT-related genes are relatively stable from year to year (Supplemental Fig. S4). We identified taxonomic patterns in the transcription of the genes involved



Fig. 3. The total particulate ¹⁵N (top) and ¹³C (bottom) isotope concentrations in the south (left) and north (right) incubation experiments in known, quantifiable metabolites. 'Other' includes all compounds quantified that are not otherwise specified, as listed in Supplemental Table S4. Note that natural abundance isotopes are also included in this total. ¹³C did not accumulate over time in homarine, but the concentration of homarine was high enough that its natural abundance ¹³C content contributes substantially to the total.

in the three steps of the demethylation pathway (*bhmt*, *gbcAB*, *dmg*, *dmgdh*, *sardh*, *sox*, *soxAB*) and the choline synthesis pathway (*atrr*). There were no obvious taxa transcribing homologues of *grdI*, the first step of the reduction pathway, and we therefore hypothesize that TMA and TMAO are formed from carnitine.

GBT demethylation. The pathway of GBT metabolism to dimethylglycine and sarcosine is well characterized in marine heterotrophic bacteria, and this was a major pathway of GBT transformation observed in our experiments (Fig. 4). Unlabelled dimethylglycine was below our limit of detection due to elevated noise in the mass spectrometer signal at the relevant retention time (RT) and mass-to-charge ratio; however, ¹³C₄, ¹⁵N₁-dimethylglycine, the expected product of demethylation of the isotopologue of GBT we added, was detected and accumulated over time in both experiments (Supplemental Fig. S5). ¹³C₃, ¹⁵N₁-sarcosine, the product of ¹³C₄, ¹⁵N₁-dimethylglycine demethylation, also appeared in the second time point of

each incubation experiment and reached a maximum of 30% \pm 8% and 78% \pm 4% of the total sarcosine pool in the north and south experiments respectively (Fig. 4). The accumulation of this sarcosine isotopologue is evidence that organisms were using part of the GBT molecule as an energy source, since characterized enzymes producing sarcosine from dimethylglycine, DMGO and DMGDH, result in complete oxidation of the removed methyl group (Sun et al., 2011; Lidbury et al., 2015a). Future incubations with radiocarbon labelled GBT, such as those conducted by Kiene and Williams, could quantify the proportion of GBT carbon oxidized for energy by microbial communities open ocean (Kiene and Williams, 1998).

In the northern incubation, sarcosine isotopologues did not accumulate to the same extent as the GBT isotopologues despite the general increase in sarcosine concentration in the bottles (Supplementary Figs S6 and S7). This suggests that sarcosine production in the northern incubation was not dominated by the GBT demethylation pathway. Sarcosine can also be produced by the methylation of glycine, a pathway that has been identified in marine bacteria, phytoplankton and invertebrates (Nyyssölä *et al.*, 2000; Lu *et al.*, 2006; Kageyama *et al.*, 2018; Ngugi *et al.*, 2020). This difference of sarcosine production in our two incubations through dominantly a degradative pathway (in the south) and a synthesis pathway (in the north) suggests that the community in the south may be acting in a predominately catabolic mode and an anabolic mode in the north. This difference may be due to the availability of nitrate to support more phytoplankton growth in the northern incubations (Table 1).

Glycine, the product of sarcosine demethylation and a potential precursor to sarcosine synthesis, was measured in the samples but no 13 C or 15 N isotopologues were detected. Based on our limit of quantification, the concentrations of unlabelled glycine measured in each experiment, and our inability to measure the $^{13}C_1$ glycine isotopologue, we estimate that the isotopologues could have been enriched up to 5% of the unlabelled glycine concentration and remain undetectable.

Transcription of *bhmt*, which encodes the first step of GBT demethylation, was dominated by SAR11, with a smaller contribution from Roseobacteria and other Alphaproteobacteria (Fig. 4). Interpretation of the bhmt gene expression should be conservative, as homologues identified in Pseudomonas aeruginosa, which also has gbcAB, do not participate in GBT metabolism (Wargo et al., 2008). Gene expression of gbcA homologues were detected at three of the five NPTZ stations sampled in 2016, while gene expression of gbcB homologues were detected at all NPTZ stations. Expression of gbcA and gbcB was dominated by Gammaproteobacteria and The demethylation Alphaproteobacteria. of dimethylglycine to sarcosine via dimethylglycine oxidase (encoded by dmg) was dominated by transcripts belonging to Gammaproteobacteria with some SAR11 contribution. The same transformation encoded bv dimethylglycine dehydrogenase (dmgdh) had transcription in SAR11, SAR116, Gammaproteobacteria and Roseobacters. The third step of the demethylation pathway, encoded by sardh, monomeric sox (msox), or tetrameric soxBDAG, was transcribed by a variety of both eukaryotic and prokaryotic organisms (Fig. 4). It is possible that sarcosine oxidases catalyse the demethylation of other products in addition to or instead of sarcosine, potentially explaining the two paralogous tetrameric sarcosine oxidase operons and a MSOX in SAR11 (Chlumsky et al., 1995; Sun et al., 2011). Few organisms transcribed all three steps of the demethylation pathway, but exchange of intermediates may have enabled the downstream transformations caused by a wide diversity of organisms.

Creatine production from sarcosine. The ¹³C₃. ¹⁵N₁creatine isotopologue appeared in the second time point in both experiments (Fig. 4, Supplemental Fig. S6), Creatine is primarily thought to be synthesized from glycine and arginine, via glycine aminotransferase and quanidinoacetate N-methyltransferase, and with these enzymes can be produced as a byproduct of the urea cycle (Allen et al., 2006; Joncquel-Chevalier Curt et al., 2015). We expect the accumulation of the ${}^{13}C_2$, ¹⁵N₁-creatine isotopologue because those are the atoms derived from alvcine. However, we instead observed accumulation of the ¹³C₃, ¹⁵N₁-creatine isotopologue (Fig. 4). This indicates that the active production of creatine in these communities was from sarcosine rather than glycine (Fig. 4). These data suggest that either there is an unidentified enzyme that catalyses the reaction from sarcosine to creatine or that creatinase, the enzyme which produces sarcosine from creatine, is reversible (Wyss and Kaddurah-Daouk, 2000).

GBT as a methyl donor. BHMT, one of the enzymes responsible for the first step of the demethylation pathway, generates methionine from homocysteine (Fig. 4) (Lidbury et al., 2015a). Other pathways for methionine synthesis are via methionine synthase (MetH), which uses cobalamin (Vitamin B₁₂) and methyltetrahydrofolate as cofactors, and MetE which directly transfers the methyl from methyltetrahydrofolate without the use of cobalamin. Methionine is used to make the universal methyl donor Sadenosyl methionine (SAM) from S-adenosyl homocysteine via methionine adenosyltransferase. Genomes of both prokaryotes and eukaryotes contain bhmt homologues (Lidbury et al., 2015a; Figueroa-Soto and Valenzuela-Soto, 2018), though no homologues have been functionally verified in eukaryotic phytoplankton. Our analysis only retrieved sequences with homology to experimentally confirmed genes in marine bacteria to avoid including sequences that may encode other types of closely related methyltransferases. Therefore, we only detected bhmt transcripts from heterotrophic bacteria, dominated by SAR11 (Fig. 4). SAR11, originally thought to have a strict glycine and reduced sulfur requirement, can use the methyl donor GBT to satisfy their requirement for glycine through complete GBT demethylation. This pathway also lessens their reliance on exogenous methionine and helps explain the lack of both MetE and the Vitamin B₁₂ dependent MetH pathway for methionine synthesis in SAR11 (Tripp et al., 2008; Sun et al., 2011; Carini et al., 2013). Our data suggest that SAR11 uses GBT as a source of both glycine and methionine in situ.

We did not detect the $^{13}C_1$ natural abundance isotopologues nor label accumulation in methionine or SAM. Instead, we observed the accumulation of ^{13}C in the methyl-acceptors β -alanine betaine and dimethylsulfoniopropionate



Fig. 4. The GBT demethylation pathway and associated metabolic pathways. Shaded squares represent the MID, which is the proportion of a given metabolite that was in the isotopologue shown, where red atoms in molecular structures indicate ¹³C or ¹⁵N derived from GBT. For each molecule, the top row represents the northern experiment and the bottom row represents the southern experiment, with the initial *in situ* conditions as the first square and the final timepoint as the last square. Molecules in grey were not detected in our data but we infer their presence based on the downstream metabolites. Dashed arrows are known metabolic pathways that we did not see evidence for in our isotope-enriched metabolomics data. The listed amino acids are those which increased in their relative isotopologue abundance in the THAAs in the southern experiment. DMSP, β-alanine betaine and trigonelline are compounds that are methylated through SAM-dependent methyltransferases and had significant increases in their ¹³C isotopologues. The pie charts show the proportion of organisms transcribing the genes associated with the demethylation pathway in this region in spring 2016: GBT to dimethylglycine (betaine-homocysteine methyl-transferase, *bhmt*, or glycine betaine oxidase, *gbcAB*), dimethylglycine to sarcosine (dimethylglycine dehydrogenase, *dmgdh*, or dimethylglycine oxidase, *soxBDAG* with *soxA* and *soxB* homologues displayed here). Pie charts show an average across the five stations in the NPTZ, with the exception of *gbcA* which was only detected at three stations. 'Other alphaproteobacteria' includes non-Roseobacter Rhodobacterales, Rhizobiales and Sphingomonadales.



Fig. 5. MIDs (in percent) over time for various isotopologues detected in the metabolites that show evidence of C and N recycling through cellular metabolism. MID is the percent of a given metabolite that was in a particular isotopologue. Isotopologues with only 12 C and 14 N atoms that make up the majority of the MID are not shown so that the isotopologues with atoms derived from the added $^{13}C_5$, $^{15}N_1$ -GBT can be better resolved on the *y*-axis. The *x*-axis is incubation time in hours. For each compound, the northern experiment is on the right and the southern experiment is on the left. Solid lines indicate the mean of triplicate bottles and the error bars represent one standard deviation. Horizontal dashed lines indicate the MID for samples taken at the stations where each experiment was performed.

(DMSP, Figs 4 and 5), providing evidence that GBT fuelled the methionine cycle in these experiments. For example, β-alanine betaine is made through successive methylations of *β*-alanine via a SAM-dependent methyltransferase (Parthasarathy et al., 2019). _β-alanine betaine showed changes in ¹³C isotopologues even though β -alanine did not (Fig. 5, Supplemental Tables S3 and S4), indicating that the ¹³C was transferred as a methyl group derived from SAM. The methylation of SAM from GBT, as demonstrated by these downstream products, may be an important pathway in marine microbial communities. Vitamin B₁₂ (the cofactor for MetH) can be a limiting nutrient for growth in euphotic marine environments (Sañudo-Wilhelmy et al., 2014; Bertrand et al., 2015; Browning et al., 2017). For organisms with BHMT and MetH, storage of GBT could provide a safeguard for methylation potential during temporary Vitamin B₁₂ scarcity.

Because only a subset of organisms can use GBT for methylation (via BHMT), ¹³C accumulation patterns in metabolites synthesized with SAM-dependent methyltransferases can provide insights into which microbes produce them *in situ*. DMSP is synthesized via a SAM-

dependent methyltransferase by both phytoplankton and bacteria (Curson et al., 2017, 2018), and ${}^{13}C_1$ and ${}^{13}C_2$ isotopologues of DMSP accumulated over time, particularly in the southern experiment. Gonvol, potentially produced from DMSP (Trottmann et al., 2020), showed similar isotopologue accumulation (Fig. 5). Though DMSP can be made by heterotrophic bacteria that expressed bhmt in this region (Fig. 4), gonyol synthesis seems to be restricted to eukaryotic plankton (Nakamura et al., 1993; Gebser et al., 2020; Heal et al., 2021, Durham et al., submitted). The appearance of labelled gonyol therefore suggests that either there is a bacterial gonvol source or that some of the unverified eukaryotic bhmt homologues we excluded from our gene expression searches are functional and may include dinoflagellates, as they are considered the main gonyol producers (Gebser et al., 2020; Heal et al., 2021).

Homarine and trigonelline are structural isomers made by different biosynthetic pathways that both involve SAMdependent methyltransferases (Hall and Gurin, 1975; Ashihara, 2008). The $^{13}C_1$ isotopologue of trigonelline increased in both experiments, with a greater increase in

the south than in the north (Fig. 5). We therefore propose that organisms synthesizing trigonelline in the south are heterotrophic bacteria, the organisms dominating bhmt transcription we measured, or possibly the eukaryotes that are likely producing gonyol, as discussed above. In contrast, the homarine ¹³C₁ isotopologue showed no significant change over time in the southern experiment (Fig. 5), indicating that organisms synthesizing the high concentrations of homarine in the south are not using BHMT. Given the high concentration of homarine, some homarine producers may use BHMT, but these organisms are not actively producing enough homarine to cause an observable trend in homarine isotopes. Laboratory cultures implicate phytoplankton as the primary homarine producers (Heal et al., 2021, Durham et al., submitted) and we suggest they may be the dominant organisms producing homarine in these experiments but that those phytoplankton cannot access the methylation potential of GBT through BHMT.

GBT into total hydrolyzable amino acids. Over the course of the experiments, the concentration of total hydrolyzable amino acids (THAAs) increased by \sim 50% in the northern experiment and decreased by $\sim 10\%$ in the southern experiment (Supplemental Fig. S8). ¹³C and ¹⁵N isotopologues of several hydrolyzed amino acids were detectable, but there were no significant differences in the isotopic distributions at the initial and final timepoints [false discovery rate (FDR) corrected p-value >0.1], likely due to high bottle to bottle variability. In the southern experiment there were increases in the proportional contributions of the ¹³C₂ and ¹³C₃ isotopologues of phenylalanine, isoleucine, leucine, and valine and ${}^{13}C_2$, ¹⁵N₁ isotopologues of phenylalanine, isoleucine, and alanine that were significant at a lower threshold (p-value <0.05 without FDR correction, Fig. 6). Some of these isotopologues were below our limit of detection in the initial time point and increased to be above our limit of detection at the final timepoint, despite an overall decrease in the THAA concentrations.

Of the five amino acids that showed isotope enrichment, four (alanine, valine, isoleucine and leucine) are synthesized from pyruvate. Phenylalanine has three carbons derived from phosphoenolpyruvate, which itself can also originate from pyruvate. These synthesis routes and the enrichment of two or three ¹³C atoms in these amino acids provide further evidence that GBT is demethylated to glycine, which can then be converted to serine and then to pyruvate and ammonium through the glycine cleavage system and serine dehydratase (Fig. 4). Meta-transcriptomic (Fig. 4) and genomic evidence (McParland *et al.*, 2021) suggest heterotrophic bacteria were metabolizing GBT through the demethylation pathway to glycine, and thus likely used GBT for protein production.

GBT to choline. Choline is canonically thought of as a precursor of GBT (Kiene, 1998; Lidbury et al., 2015a), but our observations suggest this process is reversible. The labelling pattern we observed $({}^{13}C_{51}, {}^{15}N_1)$ strongly suggests that choline was formed directly from GBT in our experiments (Fig. 7, Supplemental Fig. S6), Choline and glycerophosphocholine were in higher concentrations in the northern experiment than in the south (Supplemental Fig. S7), and ¹³C₅, ¹⁵N₁ isotopes accumulated more rapidly in the north, indicating high flux through this pathway. At the end of the incubation experiment in the north, approximately 25% of the total choline and glycerophosphocholine metabolite pools were in ¹³C₅, ¹⁵N₁ isotopologues. The additional accumulation of the glycerophosphocholine ${}^{13}C_6$, ${}^{15}N_1$ and ${}^{13}C_7$, ${}^{15}N_1$ isotopologues necessitates that the glycerol group was labelled with ¹³C, supporting further metabolic reworking of GBT-derived carbon.

Glycerophosphocholine is made from choline-phospholipids, a lipid class predominantly produced by eukaryotic organisms in marine plankton communities (Van Mooy and Fredricks, 2010; Popendorf et al., 2011). A nonribosomal peptide synthetase-like enzyme (ATRR) that catalyses the reduction of GBT to choline was recently characterized in fungi (Hai et al., 2019). In the metatranscriptome data, transcription of atrr gene homologues was dominated by members of Dinophyceae, Stramenopiles, Haptophyta and Amoebozoa (Fig. 7). These transcripts further support GBT use as an intermediate in lipid biosynthesis by eukaryotes, thus far an underappreciated role for GBT in marine environments. Additional research into the prevalence of atrr in marine ecosystems will highlight where and when this use of GBT may be important.

¹³C₅, ¹⁵N₁ GBT to carnitine. We observed the isotopologue of carnitine in the second time point in both experiments (Fig. 7), suggesting direct synthesis from GBT. Carnitine had a similar total concentration in each experiment (Supplemental Fig. S7) but higher isotope accumulation in the south (Fig. 7, Supplemental Fig. S6). GBT can be formed from carnitine through carnitine dehydrogenase (CDH) and a carnitine-specific BKACE (Meadows and Wargo, 2015). CDH is an oxidoreductase with NAD⁺ as an acceptor, a type of enzyme that has the potential to be reversible under the right chemical conditions (Sellés Vidal et al., 2018). It is therefore possible that these enzymes are responsible for the formation of carnitine from the GBT we observe. However, to avoid misinterpretation of the transcriptomics data we did not search for expression of these enzymes that have only been experimentally demonstrated to synthesize GBT.

TMAB [also known as (3-Carboxypropyl) trimethylammonium and γ -butyrobetaine], is a quaternary

amine that can be a precursor to carnitine or a product of carnitine metabolism (Bremer, 1983; Meadows and Wargo, 2015). The stable isotope accumulation patterns in our experiments suggest that TMAB is derived from carnitine (Fig. 7, Supplemental Fig. S6). The ¹³C₅, ¹⁵N₁-TMAB isotopologue increased throughout the course of both experiments, from 0% to 6% of the total TMAB pool in the north and from 0% to 32% of the total TMAB pool in the south. The isotope accumulation in carnitine and TMAB highlight the relevance of unexplored pathways for compatible solute metabolism in marine microbial communities.

Glycine betaine to trimethylamine-N-oxide. TMAO is a compatible solute and can be an intermediate in the formation of other methylamines and further oxidized to provide energy (Yancey and Siebenaller, 1999; Sun *et al.*, 2011; Lidbury *et al.*, 2014; Lidbury *et al.*, 2015b). The ¹³C₃, ¹⁵N₁-TMAO we observe in our incubations could have been formed from three different pathways (Fig. 7, Supplemental Fig. S6). The most direct route is GBT reduction via betaine reductase (GrdH/GrdI) to form TMA (pathway 3 in Figs 1 and 7), a pathway characterized in anaerobes (Jones *et al.*, 2019). TMAO is then produced from TMA by TMA monooxygenase, which is found in both prokaryotes and eukaryotes (Yancey and Siebenaller, 1999; Welsh, 2000; Yancey, 2005). Another route of TMAO synthesis is from choline via choline

TMA-lyase, CutC, an enzyme in anaerobic organisms (Jameson *et al.*, 2016). Lastly, TMA and TMAO can also be formed from carnitine via carnitine oxygenase, an enzyme in aerobic organisms (Jameson *et al.*, 2016).

In the metatranscriptome sequence data, reads associated with *grd1* were only detected at a single station at low abundance. At that station (35.28 °N), transcription was dominated by sequences taxonomically assigned to a marine Clostridium (Fendrich *et al.*, 1990) which was unlikely to have been dominant in the oxic water column, since the organism was originally described in anoxic sediments. The low number of *grd1* annotations and corresponding transcripts along with the fact that Grd1 and CutC are characterized in anaerobes point to carnitine as the most likely source of TMA for TMAO production in this environment.

Carnitine showed significant labelling in both the northern and southern experiments, as discussed above, and the transformation of carnitine to TMA, encoded by *cntA* and *cntB*, has been demonstrated in Acinetobacter (Zhu *et al.*, 2014) and *cntA* homologues have been found in open ocean metagenomes (Jameson *et al.*, 2016). In searching marine reference genomes and transcriptomes (see Experimental Procedures) for *cntA* and *cntB* homologues, we were unable to identify organisms with highquality homologues for both enzymes and thus refrain from speculating about which organisms might have been responsible for TMAO production in this study.



Fig. 6. The isotopologues of individual THAAs with increases in their isotopologue distribution in the southern experiment (p < 0.05 without FDR correction). The *x*-axis shows incubation time in hours. Replicate samples (n = 3) are in grey and the mean value is in black. No significant increases were observed in the northern experiment.

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Fig. 7. The transformation of GBT to choline, glycerophosphocholine, TMAO, carnitine and TMAB. Circled numbers refer to pathways as in Fig. 1 and as discussed in the text. Shaded squares represent the MID, which is the proportion of a given metabolite that was in the isotopologue shown, where red atoms in molecular structures indicate ¹³C or ¹⁵N derived from GBT. For each molecule, the top row represents the northern experiment, and the bottom row represents the southern experiment, with the initial *in situ* conditions as the first square and the final timepoint as the last square. Molecules in grey were not detected in our data but we infer their presence based on the downstream metabolites. Dashed arrows indicate the three potential routes of TMA synthesis. The pie charts show the microorganisms transcribing *atrr* in the transition zone in spring 2016. The pie chart for *atrr* is an average of the five stations in the NPTZ.

Biomass production and nitrogen re-mineralization

Increases in the ¹³C and ¹⁵N content of amino acids, nucleosides and nucleobases indicated that GBT-derived carbon and nitrogen were used in macromolecular biosynthesis pathways within cellular metabolism (Fig. 5). Significant increases in isotopologue distributions over time (FDR corrected p < 0.05) are discussed below.

GBT nitrogen re-mineralization and reincorporation. The $^{15}N_1$ isotopologues of glutamic acid, glutamine and arginine significantly increased in both experiments (Fig. 5). Glutamine and glutamic acid are the molecules involved in ammonium assimilation, and the appearance of these isotopologues suggests that ammonium was generated from GBT. Other metabolites, including trigonelline,

homarine, taurine, proline and alanine, showed ¹⁵N incorporation in one or both experiments (Fig. 5, Supplemental Tables S3 and S5). The enhanced ¹⁵N isotopologue accumulation trends in many metabolites in the southern station reiterate that, when DIN concentrations are low, organisms are reliant on recycling of organic nitrogen within the photic zone. The small increase in ¹⁵N isotopologues in both trigonelline and homarine underscores that nitrogen from GBT made its way through the microbial community, including organisms that may not be able to degrade GBT themselves, as has been observed in co-culture experiments (Lidbury *et al.*, 2015a; Zecher *et al.*, 2020).

Purine and pyrimidine metabolism. The purines adenine and guanine, and related molecules adenosine,

deoxvadenosine and quanosine, showed enrichment in many isotopologues in both the northern and southern experiments, including ${}^{15}N_1$ alone, ${}^{13}C_1$ alone and ${}^{13}C_2$, ¹⁵N₁ (Fig. 5, Supplemental Table S3). Purines are formed de novo from two molecules of formate, glycine, two molecules of glutamine, HCO₃⁻, and aspartate. The relatively high accumulation of the ¹³C₂, ¹⁵N₁ isotopologues of adenine, adenosine and guanine to 1%, 1.4% and 0.7% in the final time point of the south experiment suggests isotope accumulation from glycine, which contributes two carbons and one nitrogen atom to purines (Fig. 5, Supplemental Table S3, Supplemental Fig. S9). Similar accumulation of the ¹³C₃, ¹⁵N₁ isotopologues suggest significant labelling of tetrahydrofolate within these cells, possibly derived from the second two demethylation steps in the demethylation pathway, in which methyl groups are transferred to tetrahydrofolate and potentially ultimately oxidized to CO₂ (Sun et al., 2011). These pathways for isotope accumulation in the purines suggest that, in organisms that take up and demethylate GBT, there is significant recycling of carbon and nitrogen from GBT throughout primary metabolites, including possibly formate and CO₂.

In contrast to purines, pyrimidines like cytosine are formed *de novo* from glutamine, HCO_3^- and aspartate, without contribution from glycine. We did not observe significant accumulation of cytosine isotopologues with multiple ¹³C and ¹⁵N atoms, but the ¹⁵N₁ isotopologue of cytosine accumulated more in the south compared to the north (Fig. 5), again emphasizing the relative importance of GBT-derived ammonia being re-assimilated for biomass production in low nitrate waters.

Untargeted metabolites with isotope incorporation

Using untargeted metabolomics, we were able to identify several other metabolites that demonstrate the varied fates of GBT-derived carbon and nitrogen. These metabolites are promising targets for future studies into marine microbial metabolism. By searching the high-resolution metabolomics data for evidence of ¹³C and ¹⁵N incorporation in all mass features (unique combinations of massto-charge ratio and RT), we found 74 and 82 high-quality mass features with significant isotope incorporation in the north and south respectively, out of about 2000 initial mass features (see Supplemental Methods). Forty-six of these mass features were shared between the experiments (Supplemental Table S5). In the south, 41% (34/82) of mass features with isotope incorporation were known metabolites, including those discussed above. In the north, 28% (21/74) of mass features with isotope incorporation were known metabolites. The remaining have identities that are either unknown or tentatively identified without an authentic standard for confirmation. Regardless of identification, all mass features represent metabolites that must be direct or indirect products of GBT metabolism. For example, a mass feature in the HILIC-positive mode with an mass-to-charge ratio (m/z) of 184.0734 and RT of 12.8 min had a similar increasing ¹³C₅, ¹⁵N₁ isotopologue trend as choline and glycerophosphocholine (N-FT0422/S-FT391 in Supplemental Fig. S9, Supplemental Table S2). Based on the exact mass and MS² spectra, we identified this metabolite as phosphocholine, which we had not previously investigated in our analysis.

Select unknown metabolites had very similar isotopeaccumulation trends as known compounds. For example, a mass feature in the HILIC-positive mode with a m/z of 175.1077 and a RT of 11.15 min had a similar ¹³C isotopologue decrease, ¹³C₅ isotopologue increase and ¹³C₅, ¹⁵N₁ isotopologue increase as GBT itself (N-FT0365/S-FT328 in Supplemental Fig. S9, Supplemental Table S3). These isotopologue increases and the exact mass suggest this metabolite is GBT-Gly, where a peptide bond has formed between the carboxylic acid of GBT and the amine of glycine. Reports of fungal synthesis of GBT-Ser (Singh *et al.*, 2021) support the possibility of a biologically produced GBT-Gly dipeptide.

We tentatively identified several untargeted metabolites as likely mycosporine-like amino acids (MAAs), using exact masses and common MS² fragments (as identified in Parailloux et al., 2020). Six potential MAAs with isotopologue accumulation in the northern experiment are userjirine (m/z 285.144, N-FT0769), mycosporineglycine (m/z 246.09, N-FT0621/S-FT600), palythenic acid (m/z 329.134, N-FT0887/S-FT807), porphyra-334 (m/z 347.144, N-FT0930/S-FT830), palythine (m/z 245.113, N-FT0614/S-FT596) and shinorine (m/z 333.129, N-FT0899/S-FT814). All but userjirine also had significant isotopologue accumulation in the southern experiment, with enrichment in multiple isotopologues, including ¹³C and ¹³C₂, ¹⁵N₁ (Supplemental Table S2). This latter isotopologue showed similar accumulation patterns to adenine and guanine and is indicative of the incorporation of a GBT-derived glycine (Supplemental Fig. S9). MAAs are primarily thought to be photoprotective and antioxidant compounds produced by algae, fungi and cyanobacteria (Shick and Dunlap, 2002; Llewellyn and Airs, 2010). Though the abundance of MAAs was generally higher in the northern experiment compared to the southern experiment, the ¹³C₂, ¹⁵N isotopologue accumulated more in the southern experiment. In marine phytoplankton cultures, molecular features with m/z and RTs matching the tentatively identified MAAs here were produced dominantly by dinoflagellates (Heal et al., 2021). We did not search for eukaryotic bhmt homologues, as discussed above, and therefore it is possible eukaryotic organisms such as dinoflagellates could be directly

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synthesizing MAAs from GBT. However, if the demethylation of GBT is primarily a heterotrophic bacterial process and MAA synthesis is primarily done by primary producers, then the labelled glycine incorporation into MAAs is indicative of active metabolite exchange. We speculate that dimethylglycine, sarcosine and glycine may be excreted by SAR11 or other bacteria carrying out the GBT demethylation pathway so that they can use the methyl groups in GBT for methylation and energy but maintain homeostasis, since SAR11 need specific ratios of glycine to pyruvate to maintain optimal growth rates (Carini *et al.*, 2013).

Future studies should focus on validating the identities of the molecules tentatively identified here, exploring their cellular roles, and confirming metabolic interactions that they might facilitate. Overall, these results show the promise of using isotope-tracing high-resolution metabolomics to identify metabolites of biological relevance and their biosynthetic origins. We expect that subsequent studies using these techniques with different substrates or microbial communities will lead to more discoveries of ecologically relevant molecules, pathways, and interactions.

Conclusion

Using stable isotope tracing metabolomics, we find that microbial communities in the NPTZ can take up GBT and transform it through both known and previously unrecognized pathways, to ultimately provide methylation capacity, carbon, nitrogen and energy to the community. The uptake kinetics parameters were within the ranges estimated in previous studies of GBT uptake in natural seawater and cultured organisms. The turnover time estimates for dissolved GBT are similar to the turnover time estimates of non-proteinogenic dissolved free amino acids and amino acid-like compounds. The vast majority of the ¹⁵N and ¹³C that we could detect remained in untransformed ${}^{13}C_5$, ${}^{15}N_1$ -glycine betaine. This luxury uptake of DON may have unexplored consequences for how organisms adapt to and survive chronically nutrientlimited biomes, including nitrogen and cobalamin limitation. We also show that a portion of the GBT is transformed to other metabolites on timescales of hours to days. The difference in uptake and fate of GBT between the two stations emphasize that the DIN availability and microbial community composition influence how DON is used. Though the dominant pathways of GBT metabolism differed, at both sites the carbon and nitrogen from GBT appeared in known and unknown metabolic products including amino acids, nucleobases, lipids and MAAs, suggesting both partial and full recycling through demethylation, pyruvate, the citric acid cycle and ammonium assimilation.

Experimental procedures

Sampling strategy

Samples for GBT experiments were collected aboard the R/V *Kilo Moana* in April 2019. Experiments were conducted at two stations: station 4 at 41° 40.85' N and 158° 3.01' W, and station 5 at 37° 0.21' N and 158° 0.20' W (Table 1). Water for samples of the *in situ* conditions and for the incubation experiments were collected with Niskin bottles attached to the CTD from 15 m water depth, which was within the surface mixed layer, in the morning. DON, DIN, particulate carbon and particulate nitrogen samples were analysed according to the standard Hawaiian Ocean Time Series program protocols. Temperature and chlorophyll-a concentrations are from the CTD.

GBT uptake kinetics experiments

To determine the kinetics of GBT uptake, whole seawater was spiked with varying concentrations of ${}^{13}C_5$, ${}^{15}N_1$ -GBT and incubated for 25-42 min. Water was collected into 2 L bottles around 8:00 am local time for both experiments. To minimize the biological transformation of ${}^{13}C_5$, ¹⁵N₁-GBT into other molecules and to limit induction of enzymatic activity, the incubation time with ¹³C₅, ¹⁵N₁-GBT was kept as short as possible given our sampling set up. In order to achieve this, samples were spiked throughout the course of the day since only six samples could be processed at a time. Before and after the addition of ¹³C₅, ¹⁵N₁-GBT, bottles were kept in flow-through incubators with blue shading to be at in situ temperature and approximately mixed layer light conditions. Samples were spiked to have final concentrations of 0, 2, 5, 10, 50, 200, or 2000 nM ¹³C₅, ¹⁵N₁-GBT. Triplicates of each ¹³C₅, ¹⁵N₁-GBT concentration were processed for both experiments, with replicates spread throughout the sampling period. After incubation with the spiked molecule, seawater was filtered onto 47 mm diameter, 0.2 µm pore size PTFE (Omnipore) filters using a peristaltic pump, polycarbonate filter holder and Masterflex PharMed BPT tubing (Cole-Parmer). Filtering time was 10-42 min with an average time of 22 min. Experimental blanks were collected for each spike concentration during the northern experiment by collecting filtrate from one replicate of each treatment and re-filtering filtrate onto a new filter. This provided a measure of dissolved organic compounds adsorbed onto the filter during processing. Filters were frozen in liquid nitrogen immediately after filtration and stored at -80°C until the filters were extracted using the metabolite extraction method described in Boysen et al. (2018).

GBT and ${}^{13}C_5$, ${}^{15}N_1$ -GBT were measured using paired liquid chromatography-mass spectrometry (LC-MS) using a Waters Acquity I-Class UPLC (Waters

Corporation, Milford, MA, USA) with a hydrophilic interaction liquid chromatography column (HILIC SeQuant ZIC-pHILIC column, 5 mm particle size, 2.1 mm × 150 mm, from Millipore) paired to a Waters Xevo TQ-S triple quadrupole. Electrospray ionization in selected reaction monitoring mode with polarity switching was used to monitor for the intensity of ions from $^{12}C_5$, $^{14}N_1$ -GBT and $^{13}C_5$, $^{15}N_1$ -GBT. Standard addition curves were used to quantify GBT as detailed in the supplementary material. Samples were always well above the concentration in the blank, with an average of 63 times greater concentration in the sample than the blank across all treatments. Thus, blank measurements were disregarded for the remainder of the uptake kinetics calculations.

Rates of ${}^{13}C_5$, ${}^{15}N_1$ -GBT uptake were calculated by fitting the data to a Michaelis–Menten equation with a non-linear least-squares method and by using a Wright-Hobbie transformation. Details are in the Supplemental Methods. All methods resulted in similar estimates with overlapping uncertainties (Supplemental Table S1). In the results and discussion, parameters from the nonlinear least-squares estimates are used. We note that the uptake rates we calculate may be underestimates if during the incubations the cells became saturated with GBT and slow their uptake rate (Button, 1998).

GBT fate experiments

Two time-course incubation experiments were performed, one at each of the stations described in Table 1. For each experiment, 2 L bottles were filled with seawater collected with the CTD from 15 m depth (within the mixed layer) at approximately 6:00 am local time. All 2 L bottles were spiked with 500 nM ¹³C₅, ¹⁵N₁-GBT and incubated in on-deck temperature-controlled incubators at 10°C and 14°C for the north and south experiments respectively. The incubators used blue shading to approximate mixed layer light conditions. Samples for the initial timepoint (T0) were filtered directly after being spiked, resulting in actual incubation times of approximately 20 min. At each time point, triplicate 2 L bottles were sampled for analysis of bacterial and picophytoplankton abundance and biomass via flow cytometry, metabolites and THAAs. Timepoints sampled in the north experiment were 0, 4.5, 9, 12.5, 36, 50 and 98 h. Timepoints sampled in the south experiment were 0, 6, 13, 25, 51 and 100 h. For all samples, incubation time was calculated by taking the difference between the time the bottle was spiked and the midpoint time of sample filtration. In the main text, we report values from these experiments as the mean \pm standard deviation of the triplicate bottles.

Flow cytometry samples were taken by aliquoting 2 ml into a cryovial, spiking with glutaraldehyde, incubating in the dark for 20 min, and flash freezing in liquid nitrogen.

Samples were stored at -80°C until analysis on an influx flow cytometer. Flow cytometry samples were analysed on a BD Influx Cell Sorter, gating heterotrophic bacteria, *Prochlorococcus*, *Synechococcus* and other phytoplankton (labelled as picoeukaryotes). The light scattering values were converted to cell size and carbon content using Mie theory (more information available at https:// github.com/seaflow-uw/fsc-size-calibration and https:// github.com/seaflow-uw/fsc-poc-calibration).

Metabolomics samples were collected as described in the kinetics experiment, total volume of 2 L. THAA samples were collected by filtering 200 ml onto glass fibre filters (GF75 grade, 47 mm) using vacuum filtration. Filters were flash-frozen in liquid nitrogen and stored at -80° C until analysis. Experimental blanks for metabolomics samples and THAAs were collected during the northern experiment by re-filtering filtrate from the samples collected at 36 h.

Metabolomics sample processing and data acquisition

Metabolite extractions were performed as in Boysen *et al.* (2018), using a modified Bligh–Dyer extraction that yielded an aqueous and organic fraction. Aqueous soluble metabolites were analysed with two injections for each sample, one with a spike of heavy isotope labelled internal standards and one with an equal volume spike of UPLC grade water. Samples were run using both HILIC and reversed-phase (RP) chromatography on a Waters Acquity I-Class UPLC (Waters Corporation) with injection volumes of 2 and 15 μ I respectively, coupled to a Q Exactive Orbitrap MS (Thermo Scientific). Details of the chromatography and mass spectrometry methods are in Boysen *et al.* (2018) and in the Supplemental Methods.

Targeted metabolite data processing

Skyline-daily (Adams et al., 2020; Pino et al., 2020) was used to integrate the peak areas of the metabolites for which we have authentic standards (Supplemental Table S4), along with the potential isotopologues of those metabolites. Metabolites or their isotopologues were removed from further analysis if they had comparable peak area in the blanks (sample peak area less than two times the blank peak area) or if they were more than 10 ppm mass deviation from expected. MIDs are the percent contribution each individual isotopologue had to the total concentration of all isotopologues for a given metabolite. MIDs were calculated for each metabolite using only the samples that did not receive spikes of isotope standards to ensure that labelled internal all isotopologues were from GBT transformation or ¹⁵N and ¹³C natural abundance and not due to our added internal standards. The significance of MID changes during the

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incubations was assessed using a linear regression of MID over time. Samples from the same stations as the incubation experiments but which were not part of the GBT experiments were analysed to determine the natural isotope abundance.

Using the samples that were not spiked with isotope labelled internal standards, the unlabelled metabolites were normalized using best-matched internal standard normalization (Boysen et al., 2018). If the coefficient of variation for a given metabolite was less than 0.1 with no normalization or if the internal standards did not reduce the coefficient of variation by more than 0.1 then no internal standard normalization was used for that metabolite. Metabolites with internal standards spiked into the samples prior to analysis were quantified directly (L-alanine, L-histidine, L-proline, L-valine, L-isoleucine, L-methionine. L-phenylalanine, L-tryptophan, adenine, cytosine, guanine, arsenobetaine, isethionic acid, sucrose and trehalose). Except for GBT, other metabolites were quantified using the response factors and response factor ratios of authentic standards spiked into water and into representative pooled samples, as described in Boysen et al. (2021).

To calculate the absolute concentration of GBT in the samples, a sample from the cruise that did not come into contact with labelled GBT was used to calculate the response factor in the relevant sample matrix that correlates the peak area of GBT to its known concentration. $^{13}C_5$, $^{15}N_1$ -GBT was spiked into this sample at a final concentration of 500 nM just before analysis. This sample was analysed five times over the course of the sample run and thus provided a robust estimate of the response factor and its variability (coefficient of variation = 0.04). This value was used to determine the concentration of all isotopologues of GBT in each vial, which was then converted into the concentration of GBT in each sample.

Untargeted metabolite data processing

Untargeted metabolite data processing was performed as follows for each of the LC–MS analyses performed on aqueous metabolites (HILIC positive mode, HILIC negative mode, aqueous phase RP positive mode). Samples were processed in R [version 4.0.3, (R Core Team, 2013)] with the xcms package [version 3.10.2, (Smith *et al.*, 2006)] using parameters in Supplemental Table S6. Custom signal-to-noise ratios were calculated for each peak, as the default implementation has known issues (Myers *et al.*, 2017), and were used to filter out likely noise peaks. Briefly, after the initial peak picking step in xcms each peak was fit to an ideal bell curve with varying degrees of skew. For each skew, the standard deviation of the difference between the ideal curve and

the real data was calculated, producing a standard deviation of the residuals as a proxy for noise level and peak similarity. The degree of skew with the minimum residual standard deviation was used as a comparison peak. The Pearson correlation between this comparison peak and the real data was combined with the height of the peak and its residual standard deviation to produce a robust proxy for peak quality. This peak quality score was set arbitrarily at a high threshold that produced an acceptable false-discovery rate of noise peaks, and peaks falling below this threshold were removed from the analysis. Default xcms functions were then used to align, correspond and fill in missing peaks. Custom code was used to identify likely ¹³C and ¹⁵N isotopologues based on calculated m/z match and chromatogram similarity (code for all the above is available at https://github.com/ IngallsLabUW/GBT uptake and use).

Details of mass feature curation and dereplication are in the Supplemental Methods. Metabolites were identified when possible, using an in-house database of authentic standards and MS^2 data for manual comparison against the Metlin database (Smith *et al.*, 2005).

For visualization, the MID values for each metabolite isotopologue were then normalized to have a mean of zero and standard deviation of one. Clustering analyses were performed using R's base 'dist' and 'hclust' functions, and the heatmaply package was used to generate interactive heatmaps for ongoing analysis (Galili *et al.*, 2018). In both cases, the Manhattan distance metric was used due to its better performance with highdimensional data.

Total hydrolyzable amino acid analysis and data processing

Amino acids were hydrolyzed as in Fountoulakis and Lahm (1998) with some modifications specified in the Supplemental Methods. Amino acids were derivatized as in Gray *et al.* (2017) using the AccQ Tag Ultra derivatization kit from Waters (Gray *et al.*, 2017). A 10 μ l aliquot of each sample was transferred into a new glass LC–MS vial and 70 μ l of borate buffer was added to each vial. After vortexing, 20 μ l of AccQ Tag Ultra derivatizing agent was added to each sample and the vials were heated at 55°C for 10 min. Blank filters that had been collected by filtering 0.2 μ M filtrate through the sampling apparatus were collected during the experiment and were analysed alongside the samples.

For analysis of derivatized amino acids, a Waters Acquity UPLC HSS T3 column (1.8 μ m particle size, 2.1 mm \times 150 mm) was used with 0.1% formic acid in water (Solvent A) and 0.1% formic acid in acetonitrile (Solvent B) at a flow rate of 0.6 ml min⁻¹. LCMS parameters are in the Supplemental Methods.

Skyline was used to integrate peak areas of each derivatized amino acid and the isotopologues of each amino acid with 0 or 1 ¹⁵N and 0-5 ¹³C (Adams et al., 2020; Pino et al., 2020). The average peak areas of amino acids in the blank samples were subtracted from the sample peak areas. Amino acids that had peak areas in the samples that were smaller than the blank average were discarded and replaced with 0. To quantify amino acids, a standard addition curve was made using a mix of unlabelled amino acid standards. The standard addition curve was run in one representative sample by spiking in amino acid standards for a final concentration of 0.5, 1 and 2.5 µM after hydrolysis and before derivatization. The standard curve samples were then derivatized and analysed alongside the samples. After quality control. MIDs were calculated for the THAAs. A ttest was used to determine if there was a significant difference in MID between the initial and final timepoints.

Metatranscriptome sample processing

Metatranscriptome samples were collected from a latitudinal transect (23.5-37.3 °N) at approximately 158 °W in the North Pacific during the R/V Ka'imikai-O-Kanaloa Gradients 1 cruise KOK1606 (April/May 2016). These data are used in the main text and figures. To show that springtime gene expression in the NPTZ is relatively consistent from year to year, we also analysed metatranscriptome samples collected from the NPTZ in May and June 2017 aboard the R/V M.G. Langseth Gradients 2 cruise MGL1704 (Supplemental Fig. S4). Sample processing and bioinformatics were nearly identical for the 2 years, with the exception that a 200 µm mesh prefilter was used for the 2016 samples and a 100 µm mesh for the 2017 samples. The data are presented as an average across the stations within the NPTZ, as defined by Gradoville et al. (2020) by using the second derivative of surface salinity as north of 30.7 °N and 32.8 °N in 2016 and 2017 respectively. Environmental conditions for the 2016 and 2017 cruises are published in Gradoville et al. (2020). Plankton samples of ~10 L of seawater collected in the mixed layer (typically from 15 m) were passed through a 200 μ m (2016) or 100 μ m (2017) nitex prefilter followed by sequential collection onto a 3 µm followed by a 0.2 µm polycarbonate filter using a peristaltic pump. This collection scheme yielded two size classes (0.2-3 and 3-200 µm) for metatranscriptome samples. Filters were flash frozen in liquid nitrogen and subsequently stored at -80°C until further processing. To generate quantitative transcript inventories, a set of 14 internal mRNA standards were added at known concentrations at the start of RNA extraction. Total RNA was extracted using the ToTALLY RNA kit (Invitrogen) as described previously (Durham et al., 2019). Briefly, RNA extracts were treated with DNase, and rRNAs were removed using Illumina's Bacteria and Yeast Ribo-Zero rRNA Removal Kits. rRNA-depleted RNA was cleaned using the Zymo RNA Clean and Concentrator Kit. Purified, depleted RNA was then sheared to ~225 bp fragments and used to construct TruSeq cDNA libraries according to the Illumina TruSeq[®]RNA Sample Preparation v2 Guide for paired-end (2 × 150) sequencing using the Illumina NovaSeq 6000 sequencing platform.

Bioinformatics

Raw Illumina data were quality controlled with trimmomatic v0.36 (Bolger *et al.*, 2014) using the parameters MAXINFO:135:0.5, LEADING:3, TRAILING:3, MINLEN:60 and AVGQUAL:20, and matching read pairs were merged using flash v1.2.11 (parameters: *-r 150 -f 250 -s 25*) (Magoč and Salzberg, 2011). FASTQ files were converted to FASTA and translated with seqret and transeq vEMBOSS:6.6.0.0 (Rice *et al.*, 2000) using standard genetic code. Potential coding frames with \geq 40 uninterrupted amino acids were retained for further analysis.

We searched for translated reads corresponding to proteins that carry out demethylation of GBT to [betaine-homocysteine dimethylalycine methyltransferase, BHMT and GBT oxidase, referred to here as GbcAB but sometimes referred to as BmoAB (Shao et al., 2018)], the demethylation of dimethylalycine to sarcosine (DMDGH; DMGO), the demethylation from sarcosine to glycine (SARDH, MSOX, and two components of the tetrameric sarcosine oxidase, SoxBDAG), the reduction of GBT to TMA (betaine reductase, GrdI), the reduction to betaine aldehyde and choline (nonribosomal peptide synthetase-like GBT reductase, ATRR), and the transformation of carnitine to TMA (carnitine oxygenase CntAB). Full-length, experimentally verified protein sequences (locus tags given in Supplemental Table S7) from reference genomes were used as gueries to identify putative orthologues in translated protein sequences from marine prokaryotic and eukaryotic genomes and transcriptome assemblies complied in a custom marine reference database (Coesel et al., 2021) using BLASTp v2.2.31. Genomes and transcriptomes used in this database were derived from Joint Genome Institute (JGI), National Center for Biotechnology Information (NCBI) and the Marine Microbial Eukaryote Transcriptome Sequence Project (MMETSP) (Keeling et al., 2014). A complete list is available in Coesel et al. (2021). Marine sequences retrieved from the BLASTp search were clustered using usearch (Edgar, 2010) and aligned using MAFFT v7 with the E-ISN-I algorithm (Katoh and Standley, 2013). The alignment was trimmed using trimAl v1.2 using -gt .05 -

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resoverlap 0.5 -segoverlap 50 options (Capella-Gutiérrez et al., 2009). The trimmed alignment file was converted to Phylip format, and the best-fit amino acid substitution matrix, among-site rate heterogeneity model, and observed amino acid frequency were determined using ProtTest 3 software (Darriba et al., 2011). A maximum-likelihood phylogenetic reference tree was built using RAxML v8 (Stamatakis et al., 2008), and only those sequences that clustered with experimentally verified enzymes were considered putative homologues. These phylogenetic reference trees served as scaffolds to recruit translated environmental metatranscriptomic reads. An hmm profile was constructed from each reference alignment using hmmbuild, followed by transcript identification and alignment to the reference using hmmsearch (parameters: -T 40 -incT 40) and hmmalign respectively, with the HMMER package v2.1 (Eddy, 2011). NCBI taxonomy was assigned to each environmental sequence using pplacer v1.1. alpha19-0-g807f6f3 (Matsen et al., 2010) based on the read placement with the best maximum likelihood score to the reference tree (parameters: -keep-at-most 1 -maxpend 0.7). Read counts were normalized by recovery of internal mRNA standards to estimate transcript abundance. Size-fractions were summed together for a given biological replicate to yield total transcripts per liter. The NPTZ stations from the 2016 cruise do not match the exact latitudes of those used in the 2019 GBT addition experiments, thus we tried to avoid overinterpretation of detailed latitudinal trends by averaging transcript inventories across the five NPTZ stations, defined as any station with a latitude north of 30.7 °N for the 2016 cruise by Gradoville et al. (2020).

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Data Availability

Metabolomics data are available through Metabolomics Workbench under Project PR001273. Metatranscriptome data from the 2016 cruise are available through NCBI's SRA under BioProject PRJNA492143. Metatranscriptome data from the 2017 cruise are available through NCBI's SRA under BioProject PRJNA816919.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

Appendix S1. Supplementary Information.

Fig. S1. The mean concentration of quantified osmolytes over the course of the GBT fate experiments, in nM C (top) and as a percent of the total C measured in these osmolytes (bottom).

Fig. S2. The particulate concentration of all possible GBT isotopologues over the course of the incubations (top). The mass isotopologue distribution (MID) over the course of the incubations (bottom). Lines are the average of the triplicate bottles, with error bars representing one standard deviation. Dashed lines on the bottom plots indicate the natural isotopic abundance of samples taken from the stations where each experiment took place. Insets show the lower abundance isotopologues.

Fig. S3. The total particulate ¹⁵N (top) and ¹³C (bottom) isotope concentrations in metabolites in the South (left) and North (right) incubation experiments, relative to the total ¹⁵N and ¹³C measured in known, quantifiable metabolites. Other includes all compounds quantified, as listed in Supplemental Table S4.

Fig. S4. Microorganisms expressing GBT-related genes in the NPTZ during springtime cruises in 2016 and 2017. The pie charts show an average of the five stations in the NPTZ for 2016 and the six stations in the NPTZ for 2017, except for *gbcA* which was only detected at three stations in 2016.

Fig. S5. Peak area of ${}^{13}C_4$, ${}^{15}N_1$ -dimethylglycine over the course of both experiments. This isotopologue of dimethylglycine was not detected in the *in situ* samples from the corresponding stations.

Fig. S6. Mass isotopologue distributions (MID, as a percent) over time for various isotopologues detected in the metabolites that are derived directly from glycine betaine in the southern (left) and northern (right) experiments. The *x*-axis is incubation time in hours. Solid lines indicate the mean of triplicate bottles, and the error bars represent one standard deviation. Horizontal dashed lines indicate the MID for

samples taken at the stations where each experiment was performed. Note the different *y*-axis values for each plot.

Fig. S7. The concentration of nitrogen in select metabolites over time in the GBT fate incubation experiments, shown as the concentration of nitrogen in all isotopologues of a given metabolite. The line and error bars are the mean and standard deviation of triplicates.

Fig. S8. The total concentration of carbon in total hydrolyzable amino acids at the initial and final (98 h) time points in the southern and northern GBT fate experiments. Points are the average value of three replicates, with the standard deviation in grey bars. Methionine and histidine were only detected in a few replicates and were therefore excluded from this total.

Fig. S9. Isotopologue accumulation trends for ${}^{13}C_2$. ${}^{15}N_1$ and ¹³C₅,¹⁵N₁ compounds identified in the untaracted metabolomics data for both experiments. Triplicate samples for each time-point are shown as independent rows. Blue colours indicate lower mass isotopologue distribution contribution, yellow colours indicate higher MID. White values indicate data removed during guality control. Compounds with identifiers written as N-FT0000/S-FT000 are unknown or tentatively identified metabolites without authentic standards to confirm their identify, with the names corresponding to the mass feature found in the northern experiment (N-FT) and southern experiment (S-FT) as in Supplementary Tables S2 and S3. Known compounds are shown with data processed through the untargeted pipeline as opposed to the targeted pipeline to make the most robust comparisons to the unknown mass features.

Table S1. The K_s , V_{max} , and turnover time. Uptake kinetics values are reported as a mean and standard deviation calculated as discussed in the Experimental Procedures section. Values are reported as: mean (SD).

Table S2. Results of a significance test to determine if isotopologues changed their distribution over time. Each isotopologue that was detected in >2 samples in a given experiment was fit with a linear model to determine if there was a significant trend with time in the MID. The *p*-value and significance (True if p < 0.05) results are presented here for each isotopologue tested in each experiment. The ¹³C-0 ¹⁵N-0 isotopologues that have a significant trend had MID decreases over time, while the isotopologues with ¹³C or ¹⁵N that were significant typically had MID increases over time. The average and standard deviation of the MID for each isotopologue in each sample are in Table S4. (XLSX)

Table S3. Table of the estimated concentrations (nM) and isotopologue distributions. The timepoint is the approximate sampling time, Incubation time (h) is the actual sampling time after initial spiking. C.lab and N.lab are the number of 13 C and 15 N atoms in the isotopologue. Iso.mz is the mass to charge ratio for the isotopologue. Ionization is the ionization we measured that metabolite in. MID is mass isotopologue distribution, where the number 1 would indicate that in a particular sample 100% of the concentration of a given metabolite was in that specific isotopologue and the number 0 would indicate that in a particular sample, 0% of the concentration of a given metabolite was in that specific isotopologue. (XLSX)

Table S4. Table of targeted metabolites measured in the glycine betaine isotope tracing experiments. 'Isotopologue significant change' column is true if in either the north or the south experiment, any isotopologue of that metabolite showed a significant trend in mass isotopologue distribution with time. 'Isotopologue significant change' column is false if we measured the metabolite but did not detect any ¹³C or ¹⁵N isotopologues, or if we did detect the isotopologues but they did not show any significant enrichment over time (p > 0.05). The details of which isotopologues were had significant MID changes in which experiment are in Table S3. (XLSX)

Table S5. A summary of the number of mass features identified in each LC-MS configuration and those with isotope incorporation. (XLSX)

 Table S6.
 Parameters used in untargeted metabolomics

 data processing (XLSX)
 Image: Compared metabolomic state

Table S7. Reference sequences used for searching meta-
transcriptomic data. For each gene, there is a list of all refer-
ence sequences used (called homologues). The
sequence(s) in red were the annotated/experimentally veri-
fied sequences used to retrieve the rest of the homologues.All these reference sequences/homologues were used to
retrieve short reads from the metatranscriptomes. (XLSX)

Table S8. The K_t (3 parameters) or $K_t + S$ (two parameters), V_{max} , and S values when two versus three parameters (n. params.fit) were fit, reported as a mean and standard deviation as calculated from the Monte Carlo error analysis described in the Experimental Procedures.