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Molecular Commerce on Coral Reefs: Using Metabolomics to Reveal Biochemical Exchanges Underlying Holobiont Biology and the Ecology of Coastal Ecosystems

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The rapidly advancing field of metabolomics encompasses a diverse suite of powerful analytical and bioinformatic tools that can help to reveal the diversity and activity of chemical compounds in individual organisms, species interactions, and entire ecosystems. In this perspective we use examples from studies of coral reefs to illustrate ways in which metabolomics has been and can be applied to understand coastal ecosystems. Examples of new insights that can be provided by metabolomics include resolving metabolite exchange between plants, animals and their microbiota, identifying the relevant metabolite exchanges associated with the onset and maintenance of diverse, microbial mutualisms characterizing unknown molecules that act as cues in coral, reproduction, or defining the suites of compounds involved in coral-algal competition and microbialization of algal-dominated ecosystems. Here we outline sampling, analytical and informatic methods that marine biologists and ecologists can apply to understand the role of chemical processes in ecosystems, with a focus on open access data analysis workflows and democratized databases. Finally, we demonstrate how these metabolomics tools and bioinformatics approaches can provide scientists the opportunity to map detailed metabolic inventories and dynamics for a holistic view of the relationships among reef organisms, their symbionts and their surrounding marine environment.

Keywords: biogeochemistry, coral reefs, coral holobiont, dissolved organic matter, metabolomics, microbial ecology, nutrient cycling

INTRODUCTION

Coastal ecosystems where benthic and pelagic habitats intermingle and the complex ecological dynamics of each biome interact: benthic suspension feeders process the plankton while benthic primary producers exude photosynthate, each variously scouring light, carbon, nutrients and oxygen according to their metabolic needs. Diverse plants, algae, vertebrates, invertebrates, protists, bacteria and viruses bloom at this ecotone, thriving on the physical and chemical complexity born from the interface of land and water. So too does the chemical diversity and interactivity in this environment bloom. Understanding the ecology of coastal organisms demands a window into the cacophony of metabolic resources that perfuse and surround the denizens of the reef: the metabolome.

Over the last two decades, analytical tools to detect, distinguish, characterize, and quantify complex chemotypes from a single environmental sample have emerged. The most prominent of these are ultra-high resolution mass spectrometry (Dittmar and Paeng, 2009; Hawkes et al., 2016; Osterholz et al., 2016) and tandem mass spectrometry [MS/MS, the latter often coupled to various types of liquid and gas chromatographic separation methods (LC, GC)]. Improvements in sample throughput, data management and bioinformatics make it possible to track these compounds increasing and decreasing in abundance through time and space in an open market of chemical dynamics (Steen et al., 2020). These advances are not trivial; in fact, this decade marks the rapid and exponential arrival of cost efficient, interpretable, and deployable untargeted mass spectrometric experimentation (da Silva et al., 2015). **It is the goal of this article to demonstrate to coastal ecologists how metabolomics can inform organismal physiology and ecology as well as reveal the biochemical roles of organisms in ecosystems.** Herein we outline approaches that marine organismal ecologists, coastal ecosystem scientists and resource managers can employ to sample, query and interpret the metabolome of organisms and ecosystems at the interface of benthic and pelagic habitats, using coral reef systems as one concrete example.

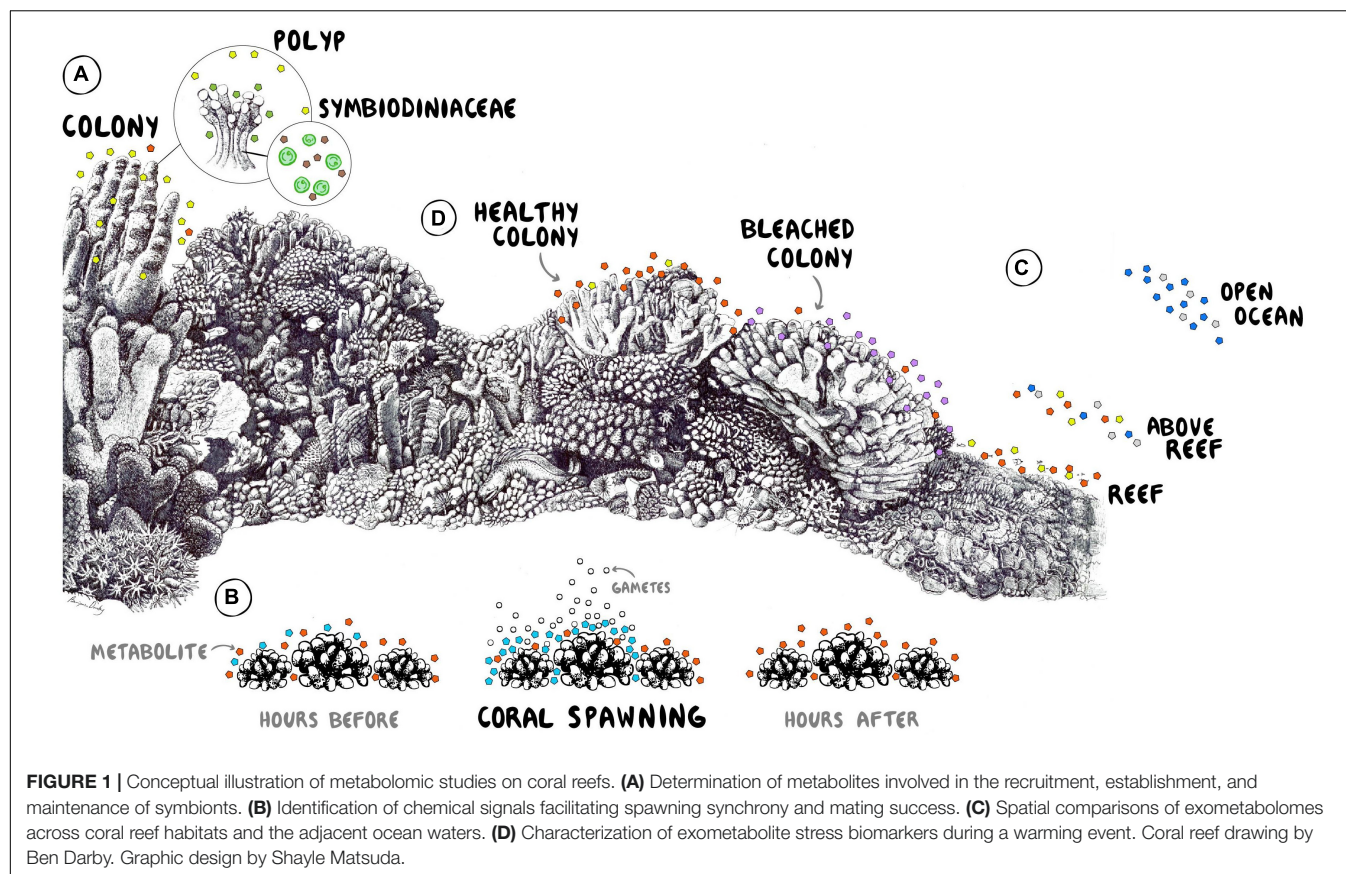
HOLOBIONT METABOLOMICS: IDENTIFYING DRIVERS OF HOST-SYMBIONT PHYSIOLOGY, SYMBIOSIS, AND DISEASE

In marine holobionts, environmental fluctuations and disturbances influence relationships between multicellular hosts and symbionts, with far reaching consequences ranging from physiological variations to dysbiosis and disease (Apprill, 2017). Coral growth and calcification, for example, are influenced by the performance of algal symbionts (Symbiodiniaceae) which are affected by light, temperature and nutrient availability. In addition to the zooxanthellae symbiosis, the coral colony is

host to hundreds, if not thousands, of other viral, bacterial, archaeal, protist, and fungal symbionts that comprise the coral holobiont (Rohwer et al., 2002; Knowlton and Rohwer, 2003; Casas et al., 2004; Kellogg, 2004; Wegley et al., 2004, 2007; Marhaver et al., 2008; Bourne et al., 2016; Ainsworth et al., 2017; Thurber et al., 2017). The diversity and complexity of these host-microbe relationships has become increasingly clear, yet no shortage of questions have emerged regarding the establishment and coexistence of these vital partnerships (Ainsworth and Gates, 2016). The current knowledge about coral microbiomes can be combined with metabolomic studies to reveal how microbial symbionts protect host fitness by conferring resistance to stressors, recovery from bleaching, and prevention of diseases.

A research priority after two decades of genomic research on coral holobionts is understanding the extent to which metabolic complementation occurs between hosts and their microbial symbionts (Gerardo, 2013; Shoguchi et al., 2013). The genomes of host and symbionts frequently lack key genes in important metabolic pathways, indicating an interdependence on the mutualistic exchange of metabolic intermediates between holobiont members (Ferrier-Pagès et al., 2005; Shinzato et al., 2011; Matthews et al., 2018). Recent studies have used metabolomic approaches to explore how functionally divergent endosymbionts impact the nutritional resources and fitness of their hosts (Matthews et al., 2018, 2020a) and careful work at the scale of bacterial-epithelial interactions has demonstrated how even taxonomically homogenous endosymbionts may facilitate spatial heterogeneity in metabolite profiles (Geier et al., 2020). Hermatypic corals colonized by two different species of Symbiodiniaceae exhibit similar metabolite profiles in both host and symbiont, illustrating the potential for more thermally robust strains to colonize diverse hosts without negatively impacting resource allocation (Matthews et al., 2020a). A variety of tools employed to examine the metabolomes of thermally stressed corals have demonstrated significant differences between stressed and control corals (Sogin et al., 2016; Farag et al., 2018; Williams et al., 2021) and Symbiodiniaceae (Petrone et al., 2018; Lawson et al., 2019) while others have demonstrated changes in metabolite exchange between host and symbiont (Hillyer et al., 2017, 2018) or phenotypic differences in whole tissue metabolomes (Roach et al., 2021) or symbiocyte acid-base homeostasis (Innis et al., 2021). In a recent review, Matthews et al. (2020b) posit that Bacteria-Symbiodiniaceae metabolomic interactions that regulate stability between host and algal symbionts are crucial for holobiont resilience to environmental stress, extending areas of investigation into more complex aspects of intra-holobiont metabolomics.

Imbalances in nutrient availability can destabilize symbiotic partnerships with algal symbionts (Wiedenmann et al., 2013) as well as commensal bacteria (Klinges et al., 2019) causing lower bleaching thresholds and higher disease incidence, respectively. Diazotrophic Bacteria and ammonia-oxidizing Archaea associated with corals have been described widely (Beman et al., 2007; Lesser et al., 2007; Pernice et al., 2012; Rädcker et al., 2015) but metabolomics opens up the



capability to investigate the exchange of organic nutrient sources within coral holobionts, providing means to examine myriad questions in the field related to the exchange of metabolomic products between symbionts (Figure 1A). The exchange of metabolites in competitive interactions between corals and macroalgae have been implicated in altering the growth of coral-associated microbiota, inducing dysbioses that potentially initiate proliferation of alternative communities that may diminish holobiont fitness, ultimately reducing competitive dominance at coral-algal interfaces (Roach et al., 2020).

Beyond the metabolomic diversity associated with the microbiome, unique host genotypes also exhibit distinct metabolite profiles that reflect physiological state and can reveal mechanisms of resilience (Lohr et al., 2019b) and photoacclimation (Lohr et al., 2019a). Different coral genera have been shown to exhibit not only unique metabolomes but also distinct profiles of metabolite modifications, hinting at a more nuanced aspect of metabolomic characterization with potential to identify both compounds and reactions that distinguish metabolomes (Hartmann et al., 2017). Comparative metabolomics of coral holobionts interacting with corals or algae demonstrated distinct metabolomes activated by non-self interactions (Quinn et al., 2016), emphasizing that host metabolomes alone may add to the suite of tools we can use for assessing organismal physiology and phenotype.

THE HOLOBIONT-WATER INTERFACE: EXOMETABOLITES REGULATE ORGANISMAL AND HOST-SYMBIONT INTERACTIONS

Communication among marine organisms is primarily dictated by chemical exchange (Hay, 2009). While a diversity of molecules used for communication have been discovered across the tree of life, most signaling compounds that mediate reproduction, larval settlement, symbiont recruitment and pathogen defense remain wholly uncharacterized (Levitan et al., 2011). Organismal interactions ranging from mutualistic to commensal to antagonistic or parasitic are chemically mediated through the water by e.g., allelopathic chemicals, toxins, morphogens, and chemoattractants for symbionts and microbes (Barott and Rohwer, 2012; Sneed et al., 2014; Tout et al., 2015).

In marine organismal ecology, metabolomics is particularly powerful for revealing otherwise invisible signaling systems and uncovering candidate molecules involved therein. For example, the reproductive signaling compounds produced by economically important marine organisms such as corals are largely unknown. Corals worldwide suffer from decreased reproduction due to Allee effects and recruitment failure (Knowlton, 2001; Vermeij et al., 2011), and recent evidence shows that several coral species from the Red Sea have lost mating synchrony (Fogarty and Marhaver, 2019; Shlesinger and

Loya, 2019). A better mechanistic understanding of the chemical cues that synchronize coral spawning could potentially open new avenues for intervention, whether in the field or in captive breeding programs. In hermaphroditic, mass-spawning corals, it has been posited that daylight and sunset times affect the coordination and initiation of the early stages of spawning (including the setting or “staging” of egg-sperm bundles in coral mouths) whereas the precise timing of the release of these gamete bundles appears to be driven by a local threshold concentration of pheromones within a localized patch of coral colonies (Levitán et al., 2011). Metabolomic tools can help to characterize these pheromone systems and determine which coral hormones induce conspecific neighbors to spawn. **Figure 1B** conceptually illustrates how specific compounds can be distinguished from spawning colonies of the grooved brain coral *Diploria* in Curacao (Arts and Haas, unpublished data), demonstrating the potential for untargeted metabolomic studies to identify putative cues involved in reproduction and synchronous spawning.

After spawning and fertilization occur, the behavior of coral larvae and settled juveniles is also influenced by local chemical cues which can either induce or suppress larval settlement, metamorphosis, polyp division and growth (Kuffner et al., 2006; Hadfield, 2011; Tebben et al., 2011; Sneed et al., 2014). Recent advances in gamete collection, *in vitro* fertilization, and larval propagation have created the potential for larger-scale restoration efforts to repopulate reefs with coral juveniles (e.g., Chamberland et al., 2017a,b; van Oppen et al., 2017) yet post-settlement survival remains a major bottleneck to scale and success (Boström-Einarsson et al., 2020; Randall et al., 2020). A mechanistic understanding of the biochemistry that underpins coral settlement and survival can be accomplished by identifying specific metabolites involved in these processes; this could help drive the development of novel settlement substrates and grow-out methods.

Other important metabolic dynamics at the holobiont-water interface include the biochemical strategies used by juvenile coral recruits to acquire and regulate their extra- and intracellular symbionts, which are obtained either vertically from parent colonies or horizontally from the environment; these symbionts may help corals resist pathogens and establish competitive dominance on the reef benthos. For example, the acquisition of Alphaproteobacterial symbionts in the family Rhodobacteraceae by coral offspring both in brooding (Sharp et al., 2012) and broadcast spawning corals (Apprill et al., 2009; Ceh et al., 2013) suggests a key role for these symbionts, but, thus far, we have only a limited understanding of the benefits that these symbionts may confer to their hosts (Sharp and Ritchie, 2012). There is evidence that some coral symbionts produce protective metabolites such as antimicrobial peptides (Ritchie, 2006) and quorum-quenching molecules (Teplitski and Ritchie, 2009; Ma et al., 2018) that could both serve to protect coral spawn and larvae against uncontrolled bacterial colonization. Other bacterial symbionts promote the coral-Symbiodiniaceae symbiosis by supplying vitamin B12, an exogenously acquired cofactor essential for *de novo* synthesis of methionine (Croft et al., 2005; Lawson et al., 2018). A number of metabolomic tools could help to illuminate the biochemical mechanisms that underly these species interactions. For example, untargeted characterization

of metabolites from bacterial isolates collected during spawning or targeted examination of egg bundles and newly developed larvae could help to resolve the protective roles of the microbes associated with each stage of coral reproduction and survival (Thompson et al., 2015).

Experimental evidence has shown that chemical gradients play an important role in bacterial chemotaxis and the establishment of coral symbioses. Specific amino acids (e.g., tryptophan and aspartic acid), nitrogen sources, and dimethylsulfoniopropionate (DMSP) all elicit chemoattractant behavior by natural assemblages of coral-associated bacteria (Tout et al., 2015). These chemotactic cues are presumed to attract beneficial taxa to the coral microbiome, although these gradients may also serve as signals for pathogens to locate coral hosts or even serve as indicators of stress or susceptibility. Compounds such as DMSP are regulators of pathogenesis; for example, the infochemical DMSP stimulates the chemotaxis of the coral pathogen *Vibrio coralliilyticus* toward its coral host (Garren et al., 2014; Sogin et al., 2016; Farag et al., 2018). When corals are heat stressed, concentrations of DMSP are elevated, promoting even greater pathogen recruitment toward susceptible hosts; combined with the release of temperature-regulated virulence factors, this mechanism rapidly promotes coral disease states (Boroujerdi et al., 2009; Kimes et al., 2012; Garren et al., 2014).

THE WATER COLUMN EXOMETABOLOME: COMPLEX MIXTURES OF BIOCHEMICALS INFLUENCE ACQUISITION, EXPORT, DISTRIBUTION, AND SEQUESTRATION OF CARBON AND NUTRIENTS

Highly complex and productive coral reef biomes thrive in regions that are depleted in mineral nutrients. Tightly coupled exchanges and efficient recycling of available resources are therefore crucial. Organic matter provided by primary producers is one of the main vectors for the transfer of energy that fuels the entire reef system. Autotrophic holobionts like corals, sponges, or algae, and even microbes exude a significant proportion of carbon fixed through photosynthesis as dissolved organic matter (DOM) into the surrounding reef waters (Haas et al., 2011, 2013). These exudates are rich substrates for microbial growth and remineralization (Haas et al., 2011; Nelson et al., 2013) and the coupled processes of production and consumption potentially establish a significant flux of organic nutrient recycling. Holobionts have been shown to establish a sphere of microbial associates that extends beyond the narrow boundary layers, variously termed the aurabiome (Walsh et al., 2017) or ecosphere (Weber et al., 2019), that is likely driven by the organic milieu exuded into the surrounding water column by reef primary producers (**Figure 1C**). Seminal work has also shown distinct metabolomic gradients in this sphere of influence that differ between diseased and healthy corals and contain suites of compounds associated with animal-microbe interactions (Ochsenkühn et al., 2018). Establishing the exometabolome of reef ecosystems will unveil a new analytical window to understand their persistence, their ability to support a productive

fishery and ultimately their resilience to both local and global anthropogenic threats such as nutrient pollution, deoxygenation and warming, acidification, respectively.

Extensive research has illustrated the multiple effects of DOM from different exometabolome sources in coral reef ecosystems (Haas et al., 2011; Rix et al., 2017; Kelly et al., 2018, 2019). For instance, exudates derived from corals promote recycling into the benthic community, reducing the loss of energy and nutrients from the reef ecosystem (Wild et al., 2004). Coral exudates potentially foster a distinct microbial community throughout the reef waters enriched in more oligotrophic taxa with higher growth efficiencies (Nelson et al., 2013) that can potentially increase the net productivity of the entire system (Haas et al., 2013) in part through suspension feeding on bacterioplankton (McNally et al., 2017). In contrast, the exometabolome of fleshy algae has been shown to foster less efficient and potentially pathogenic bacterioplankton communities (Nelson et al., 2013) and algal dominated reefs are commonly associated with increased microbial biomass (Dinsdale et al., 2008; Bruce et al., 2012; McDole et al., 2012), reduced trophic transfer to higher trophic levels (Silveira et al., 2015) and higher abundances of copiotrophic bacteria (Haas et al., 2016), leading to greater respiration of organic resources and contributing to the overall degradation of the coral reef ecosystem. The influence of emergent sponges on DOM dynamics in coral reefs has also been shown to be instrumental via the *sponge loop* (De Goeij et al., 2013, 2017; Silveira et al., 2015). While almost all these studies emphasize the importance of the source and composition of DOM in supporting the energetic budgets of reef ecosystems, very few have chemically characterized coral reef exometabolomes (e.g., Haas and Wild, 2010; Nelson et al., 2013; Fiore et al., 2017; Letourneau et al., 2020; Weber et al., 2020; Olinger et al., 2021). Further characterization of DOM exudates is required for elucidating the detailed mechanisms behind these selective processes and could provide insights into enhanced coral growth versus fleshy macroalgae competition and may even guide new conservation approaches. Because our working group and others have reported preliminary evidence of significant changes in microbial activity during warming and deoxygenation events (Johnson et al., in press; Quinlan, unpublished data) and specific coral-derived metabolites have been characterized during bleaching (Sogin et al., 2016; Farag et al., 2018; Williams et al., 2021), we anticipate that stress biomarkers can be measured from reef waters in response to environmental disturbances, providing a new avenue for early detection and monitoring (**Figure 1D**).

Another vital area where characterization of dissolved metabolites can rapidly advance coastal ecosystem ecology is in the realm of anthropogenic pollutants. Terrigenous sources of pesticides and pesticide breakdown products (DDT, DDE, glyphosates, benzamides, imidazolinones) are widespread, and little is known of the distribution or transformation of these compounds in the coastal marine environment (Mackintosh et al., 2016). Additionally, cleaning products, fluoropolymers, pharmaceuticals and a variety of compounds broadly referred to as persistent organic pollutants (POPs) have all been associated with a wide range of toxicities to coastal organisms (Trego et al., 2018). Metabolomic analyses of DOM can provide insight into the prevalence and distribution of POPs and other

xenobiotics that are amenable to the extraction and analysis methods employed (Lara-Martín et al., 2020). Even natural organic inputs from streams, estuaries and groundwater sources have been shown to exhibit unique organic signatures (e.g., Nelson et al., 2015), and untargeted metabolomics of this DOM as it is modified in the marine environment can better define the connectivity of coastal systems and the potential for allochthonous organic matter to play a role in reef nutrient and food web dynamics.

APPROACHES FOR APPLYING UNTARGETED METABOLOMICS IN COASTAL MARINE ECOLOGY FROM HOLOBIONT PHYSIOLOGY TO ECOSYSTEM PROCESSES

Encouraging the application of a complex new methodology to an established area of study demands a pathway to facilitate early adoption. In addition to advances in analytical instrumentation, such as tandem mass spectrometers, the field of metabolomics has made great strides in recent years in the democratization of bioinformatic tools and chemical databases, both of which facilitate the adoption of these methods by disparate fields. Our goal here is to introduce one approach that the authors have used to acquire and interpret biochemical data from coral reef ecosystems by recommending bioinformatic methods bundled into the Global Natural Products Social Molecular Networking (GNPS; Wang et al., 2016) system; we have used these approaches for several years and have had success in elucidating unknown metabolites in diverse ecosystems.

Collecting samples for metabolomic analyses in the field or at remote experimental stations is relatively straightforward and can be done even without access to refrigerants, although use of ultra-low temperature freezers (-80°C or lower) is recommended where possible. Biomass samples should be frozen or solvent-extracted as quickly as possible as metabolites are typically dynamic especially in organismal tissues. In general, metabolomic samples are extracted in organic solvents such as methanol. Small quantity tissue biopsies (mg to g scales) can be directly extracted in small volumes (mL) and chilled or frozen until processing can be undertaken. The extraction of dissolved metabolites from seawater can be done by using the widely adopted approach of Dittmar et al. (2008) where filtered, acidified water is passed slowly (no more than 1 L per h) over a solid phase extraction resin to bind metabolites that are subsequently eluted in methanol: PPL, a hydrophobic non-polar styrene-divinylbenzene polymer (Agilent), is sold in straight barrel cartridges containing a packed bed of resin that has been demonstrated to isolate a wide range of non-polar and semi-polar DOM compounds (Dittmar et al., 2008; Li et al., 2016; Petras et al., 2017). Extraction techniques and data acquisition vary considerably, but for the purposes of this Perspective we assume the extracted metabolites are processed via liquid chromatographic separation and analyzed via electrospray ionization tandem mass spectrometry. This approach is widely applied for untargeted analysis of small to medium sized

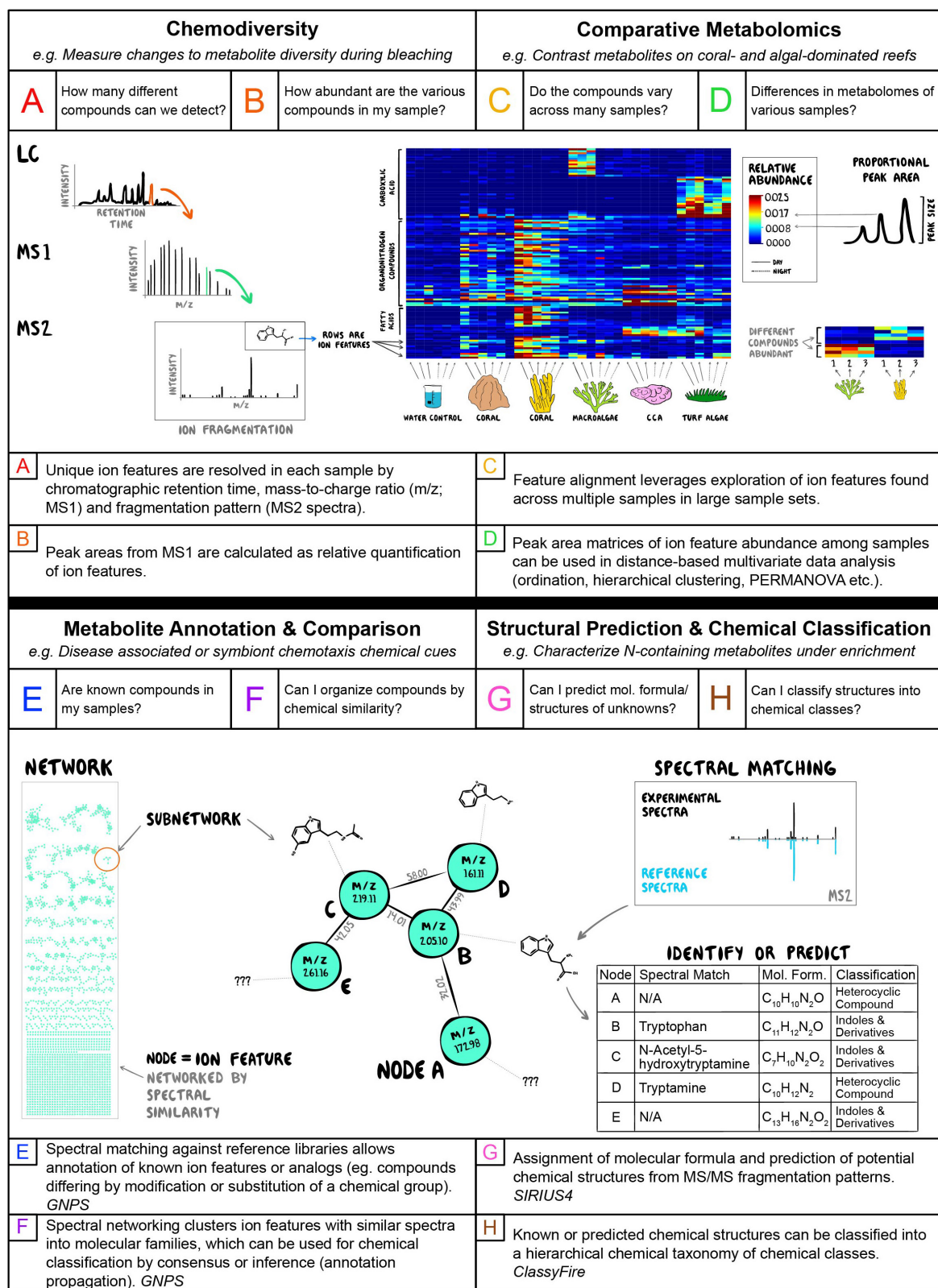


FIGURE 2 | Summary of research areas and tools in coral reef science appropriate for metabolomics analysis, including specific questions asked with metabolomic data (**A–H**, top of panels), and example public access tools and informatic workflows recommended (**A–H**, bottom of panels).

molecules (150–3000 Da), the mass spectrometers are widely available and the analytical methods are well documented (Petras et al., 2017).

Statistical, bioinformatic and visualization analyses of tandem mass spectrometry datasets derived from environmental samples and their application to questions in coastal marine ecology are outlined in **Figure 2**. Questions associated with chemodiversity (**Figures 2A,B**), including basic investigations of the number (richness) and relative abundances (evenness) of biochemicals found in different organisms or habitats are optimized by methods that seek to identify putative compounds and quantify their relative abundances (translate-able to absolute abundances when run with appropriate standards in “targeted” metabolomics). Programs that find and quantify ion features in mass spectrometry data (defined as detected ion signal for an eluting compound with a representative MS/MS spectrum) are diverse and here we apply MZmine 2 as a free, widely used and well-supported option (Pluskal et al., 2010). For example, analyses of metabolite profiles in holobiont tissues may seek to understand when and where chemodiversity increases and decreases as an indicator of organismal response to environmental stressors (**Figure 2A**). Changes in chemodiversity in the water column may be associated with pulses of metabolite exchange associated with signaling or allelopathic interactions (**Figure 2B**). Questions of comparative metabolomics (**Figures 2C,D**), where ion features must be tracked across multiple samples to examine changes in chemical composition, require informatic approaches that leverage large sample sets to better align (Pluskal et al., 2010) and resolve ion features, such as the feature alignment and annotation tools built into the Feature-based Molecular Networking workflow in the GNPS ecosystem (Wang et al., 2016; Nothias et al., 2020). When unknown ion features interpreted as metabolites can be tracked across samples then analyses of beta diversity, including compositional similarity and distance based multivariate approaches, become possible (Oksanen et al., 2019). Resolving holobiont metabolomic profiles associated with pathogen infection or water column metabolite profiles induced by thermal stress is a crucial area of research in coastal benthic ecology, and these comparative approaches may allow identification of both suites of covarying metabolites that illuminate a new aspect of physiological response (**Figure 2C**) and/or wholesale changes in multivariate metabolome structure (**Figure 2D**).

Detecting and annotating unknown metabolites is the holy grail of untargeted metabolomics. While high resolution mass measurements may allow inference of ion feature molecular formulae, tandem mass spectrometry using ion fragmentation (MS2 spectra) patterns can, when coupled with sophisticated informatics, allow characterization of unknown metabolites (**Figures 2E–H**). Analyses of holobiont metabolomes, particularly those of vertebrates and plants, are likely to resolve features with MS2 spectra that are exact or near-exact matches to known compounds, allowing annotation via large public databases facilitated within GNPS (Wang et al., 2016) and unveiling key details of the chemical ecology of host organisms (**Figure 2E**). In more chemically diverse or poorly characterized habitats such as sediments, water or invertebrate holobionts, even

unannotated MS2 spectra can be analyzed via Feature Based Molecular Networking (Wang et al., 2016; Nothias et al., 2020) to organize groups of ion features with similar spectral characteristics into networks (Shannon et al., 2003) that are likely to be structurally related (**Figure 2F**). In studies of chemical signaling in benthic interactions mediated through the water column (**Figure 1**), suites of unknown molecules enriched in areas of, for example, coral-algal interaction could be networked to identify new molecular families important in antagonistic host or microbial processes (e.g., Roach et al., 2020). Another advantage of acquiring MS2 spectra in untargeted metabolomics is the potential for higher confidence assignment of molecular formulas through fragmentation trees and network based algorithms (SIRIUS and ZODIAC; Dührkop et al., 2019; Ludwig et al., 2020) or *in silico* prediction of molecular structures (CSIFingerID, NAP, MS2LDA; Dührkop et al., 2015; van Der Hooft et al., 2016; da Silva et al., 2018). Even when exact structural predictions are limited by database breadth, emerging ontologies for hierarchical chemical classification (Feunang et al., 2016) allow organization of unknown ion features into meaningful chemical classes useful for understanding the chemical ecology of systems (**Figures 2G,H**). Networking and structural prediction can be combined with chemical classification to provide order to the chaos and help illuminate gradients in broad or narrow classes of chemicals that inform coastal ecology. Such approaches are particularly useful for tracking compounds in highly dynamic environments such as the water column, where anthropogenic pollutants, lipid-based signaling compounds, carbohydrate-based photosynthetic exudates and proteinaceous breakdown products intermingle and are metabolized by microbial processes into varied reactants and products.

CONCLUSION

Metabolomic approaches can be used to obtain new knowledge about these precious coastal ecosystems that comprise some of the most critically endangered habitats worldwide. New instruments that are becoming widely accessible combined with the development of open source informatic tools allow for greater capacity to use these biochemical approaches across disciplines where chemodiversity is less well understood. Integrating metabolomic data with experimental and field studies has the potential to:

- (i) Answer specific questions about the establishment and maintenance of symbiotic partnerships between corals and their associated microbiota, including the determination of signaling compounds for the recruitment of symbionts, antimicrobial compounds produced by commensal bacteria, or compounds exchanged between symbionts and host that provide essential nutrients and support energetic budgets;
- (ii) Understand trade-offs between directed establishment of thermal tolerant species of Symbiodiniaceae and host phenotypic responses;
- (iii) Identify chemical signals instrumental in mating synchrony and larval settlement as well as those involved in the acquisition of microbial symbionts;

- (iv) Identify microbial- or host-derived metabolites that serve to protect juveniles against uncontrolled bacterial colonization or pathogens;
- (v) Compare the exometabolomes of coral reefs dominated by reef-builders or fleshy benthic constituents to identify DOM compounds that select for opportunistic pathogens or enrich for copiotrophic microbial guilds which alter ecosystem metabolism; and
- (vi) Track the distribution of terrigenous pollutants and their incorporation into coastal food webs.

Our broad perspective is that knowledge gained from metabolomic approaches will improve mechanistic understanding of relationships between hosts and their microbial consortia, and the chemical exchanges between organisms and their environment.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

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AUTHOR CONTRIBUTIONS

LWK, CN, and AH drafted the manuscript. SM designed the figures. All authors contributed to manuscript revision, read, and approved the submitted version.

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