

Mining the Metabiome: Identifying Novel Natural Products from Microbial Communities

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<http://dx.doi.org/10.1016/j.chembiol.2014.08.006>

Microbial-derived natural products provide the foundation for most of the chemotherapeutic arsenal available to contemporary medicine. In the face of a dwindling pipeline of new lead structures identified by traditional culturing techniques and an increasing need for new therapeutics, surveys of microbial biosynthetic diversity across environmental microbiomes have revealed enormous reservoirs of as yet untapped natural products chemistry. In this review, we touch on the historical context of microbial natural product discovery and discuss innovations and technological advances that are facilitating culture-dependent and culture-independent access to new chemistry from environmental microbiomes with the goal of reinvigorating the small molecule therapeutics discovery pipeline. We highlight the successful strategies that have emerged and some of the challenges that must be overcome to enable the development of high-throughput methods for natural product discovery from complex microbial communities.

Genetically encoded small molecules comprise the single greatest source of available chemotherapeutics. Since 1981, 80% of anticancer agents and roughly 50% of all Food and Drug Administration-approved drugs have been or were derived from natural products (Newman and Cragg, 2012). In addition to their clinical utility, natural products are invaluable tools for advancing basic biological research. With diverse modes of action ranging from antibiotic to anticancer to immunosuppressive and beyond, novel natural products hold promise for the future of medicine, especially for diseases with limited treatment options.

Although the systematic chemical characterization of natural products commenced in the 19th century, the use of environmentally derived substances for medicinal purposes likely predates recorded human history (Solecki, 1975). Early medical texts from Mesopotamia dating to 2600 BC describe plant-derived substances, and Ancient Greek and Chinese cultures documented the application of botanical extracts for medicinal purposes in the 4th century BC (Cragg and Newman, 2005; Ji et al., 2009; Newman et al., 2000). Thus, humans have long recognized that natural products provide an important arsenal with which to combat disease and augment healing (reviewed in Ji et al., 2009).

Antibiotic properties of *Penicillium* fungi were observed as early as 1875 by John Tyndall (Tyndall, 1876), but it was Alexander Fleming's serendipitous rediscovery and partial purification of penicillin (1) in 1928 that first demonstrated that microbes, in this case the *Penicillium rubens* fungus, produce antibacterial substances (Fleming, 1929). This discovery needs little introduction; penicillin ultimately changed the course of human civilization, curing previously debilitating and fatal illnesses. However, it is Selman Waksman who is credited with being the first to systematically explore microbial sources for novel natural products (Sakula, 1988). Waksman's work culminated in 1943 with the isolation of streptomycin from the gram-positive, soil-dwelling actinomycete *Streptomyces griseus*, while working with a graduate assistant Albert Schatz (Schatz et al., 1944; Schatz and

Waksman, 1944). This antibiotic was the first curative therapy for tuberculosis.

The recognition that natural products of microbial origin held clinical potential spurred the discovery of myriad antibacterial natural products from readily cultured environmental bacteria, including tetracycline (*Streptomyces aureofaciens*, 1945); chloramphenicol (*Streptomyces venezulae*, 1947); erythromycin (*Streptomyces erythraea*, 1949, Eli Lilly); vancomycin (2) (*Amycolatopsis orientalis*, 1953, Eli Lilly); and rifamycin (*Streptomyces mediterranei*, 1957). The advances in natural product discovery in the 1940s and 1950s fostered programs focusing on cultivable environmental microorganisms in the pharmaceutical industry. These programs were successful not only for antibiotics, but also for the development of immunomodulatory compounds such as rapamycin (Vézina et al., 1975) and cyclosporin (Stählerlin, 1996), as well as statin development (Endo, 1992), among others. However, as much of the low hanging fruit had been picked, discovery rates inevitably slowed and rediscovery became an impediment to bringing new natural products to market, leading to a waning interest from the pharmaceutical industry by the end of the 20th century (Baker et al., 2007).

Regrettably, the de-emphasis of natural products in the pharmaceutical industry coincided with major innovations in the biological sciences that could have prevented the lapse in the discovery pipelines had the effort been sustained. In the 1980s, advances in genetics and phylogenetic analysis, specifically, the use of ribosomal RNA sequencing as a microbial species profiling tool (Olsen et al., 1986), began to uncover a vast diversity of novel microbes. A combination of 16S rRNA profiling, visual analysis using fluorescent *in situ* hybridization (Langer-Safer et al., 1982), and DNA reassociation kinetics (Torsvik et al., 1990a, 1990b) estimated up to 10⁵ largely uncultured bacterial species per gram of soil and unearthed an array of unique, uncultured bacterial species in marine and thermophilic environments (Britschgi and Giovannoni, 1991; Giovannoni et al., 1990; Liesack and Stackebrandt, 1992; Stahl et al., 1984; Torsvik et al.,

1990a, 1990b; Ward et al., 1990; Weller et al., 1991). Such biodiversity profiling analyses demonstrated that in environmental metabiomes, uncultured species outnumber the cultured ones by two to three orders of magnitude (Rappé and Giovannoni, 2003; Torsvik et al., 1996, 1998). Thus, largely comprised of the uncultured majority, these metabiomes represent a vast reservoir of uncharacterized biodiversity and, by extension, untapped natural product chemistry. These observations led to the development of the field of metagenomics (Handelsman et al., 1998), where DNA extracted directly from environmental samples (environmental DNA, eDNA) is cloned into model cultured hosts for detailed analysis. This approach to studying environmental bacteria has turned out to be a particularly appealing method to use in the search for novel natural products because the genes that encode the biosynthesis of individual secondary metabolites are generally clustered on bacterial chromosomes.

Concurrent with the identification of large numbers of novel environmental bacterial species, molecular biology was undergoing rapid expansion, largely attributable to PCR and DNA sequencing technologies. Whole genome sequencing in the mid-1990s paved the way for the development of the genomics discipline whereby DNA sequences could be analyzed and subsequently probed for phenotypes. While forward genetic approaches had already shown the tendency for bacterial natural product pathways to be encoded by genes clustered together within genomes (Hopwood and Merrick, 1977; Martín and Liras, 1989), genomics enabled reverse genetic and bioinformatic inquiries into natural product biosynthesis. A subsequent, steep drop in DNA sequencing cost and rapid advances in sequencing technologies, driven by the growth of the genomics field, transformed the natural product drug discovery process. The ever-growing number of available bacterial genome sequences (>500/month, most partial, deposited at the National Center for Biotechnology Information (NCBI) in 2013 [NCBI, 2013]), made clear that significant fractions of prokaryotic genomes are devoted to secondary metabolism. Strikingly, genomic analyses have also revealed a large number of cryptic (i.e., silent) gene clusters, even within the genomes of bacterial species that have been extensively studied with respect to natural product chemistry (Bentley et al., 2002; Ikeda et al., 2003).

The realization that the majority of bacterial species remain uncultured and that even previously cultured species contain many cryptic clusters has led to a resurgence in attempts to access environmental biosynthetic diversity. In this review, we discuss new culturing approaches, genomic and metagenomic techniques, and synthetic biology techniques that are being applied to the revitalization of natural product discovery pipelines and outline the future challenges and the promise of mining the metabiomes for biomedically relevant lead structures (Figure 1).

Rethinking Culture-Based Approaches in the Genomics Age

Historically, culture-based approaches followed the same general discovery protocol, bacteria were isolated from the environment, grown in monoculture fermentation broths, subjected to organic extraction, and screened for bioactivity. To maximize the variety of bacteria that could be grown in the laboratory, culturing conditions were varied with respect to parameters

like broth composition, pH, and temperature. As the pool of easily accessible cultured bacteria has dwindled, the field refocused on developing more involved approaches to facilitate cultivation of previously uncultured bacteria and to induce activation of silent gene clusters in previously cultured microbes.

Improving Culturing Techniques

Recognizing that uncultured organisms are likely not intrinsically “unculturable,” several groups have attempted to simulate the native habitat of isolated bacterial species to enable their cultivation in the laboratory. In situ cultivation by enclosing the bacteria within a semipermeable membrane, which allows the passage of nutrients and growth factors, but not cells, and introducing the membrane-enclosed chambers into their native environment, simulated in a laboratory setting, up to 40% of species from marine biofilms were grown as pure culture microcolonies in one such experiment (Kaeberlein et al., 2002). The same group extended this simulation to culture rarely accessed aerobic and anaerobic species from subsurface sediment (Bollmann et al., 2010), and a commercial venture founded on these approaches has reported a number of novel metabolites in the patent literature, including two novel glycosylated macrolactam antibiotics NOVO3 and NOVO4 (**3**) with potent activity against methicillin-resistant *Staphylococcus aureus* (MRSA) and vancomycin-resistant *Enterococci* (VRE) with minimal in vitro toxicity to mammalian cells (Lewis et al., 2009; Peoples et al., 2011). Furthermore, by focusing screening efforts on identifying small molecules with a narrow range of activity against a specific pathogen, the collection of extracts from in situ and prolonged cultivation studies (up to 18 months; Buerger et al., 2012) resulted in the identification of lassomycin, a ribosomal cyclic peptide with potent activity against *Mycobacterium tuberculosis*, from the extract of *Lentzea kentuckyensis* (Gavriš et al., 2014). Other related techniques that have been used to culture isolated bacteria in simulated native environments in the laboratory include culturing on supporting membranes suspended on a soil slurry (Ferrari et al., 2005) and embedding single cells in agarose microdroplets and culturing them in flowing seawater (Zengler et al., 2002) or directly on living coral (Ben-Dov et al., 2009).

Activating Cryptic Metabolism by Fermentation

While the cryptic biosynthetic gene clusters present in the genomes of readily cultured bacteria represent a sizable pool of low hanging fruit for drug discovery, forcing expression of these gene clusters has not been trivial. Mirroring the difficulties associated with culturing environmental microbes, conditions required for the activation of cryptic biosynthetic pathways have been difficult to identify and so far there does not appear to be a “silver bullet” approach for inducing the expression of previously silent gene clusters. One attractive, inexpensive approach to induce expression of cryptic biosynthetic clusters involves media supplementation. While investigating effects of rare earth metals on bacterial physiology, lanthanides scandium and lanthanum were discovered to increase or induce expression of silent gene clusters in several *Streptomyces* species (plural) (spp.) through an as yet undetermined, guanosine pentaphosphate dependent pathway (Kawai et al., 2007; Tanaka et al., 2010). Addition of *N*-acetylglucosamine, a major component of fungal cell walls, was also shown to stimulate production of

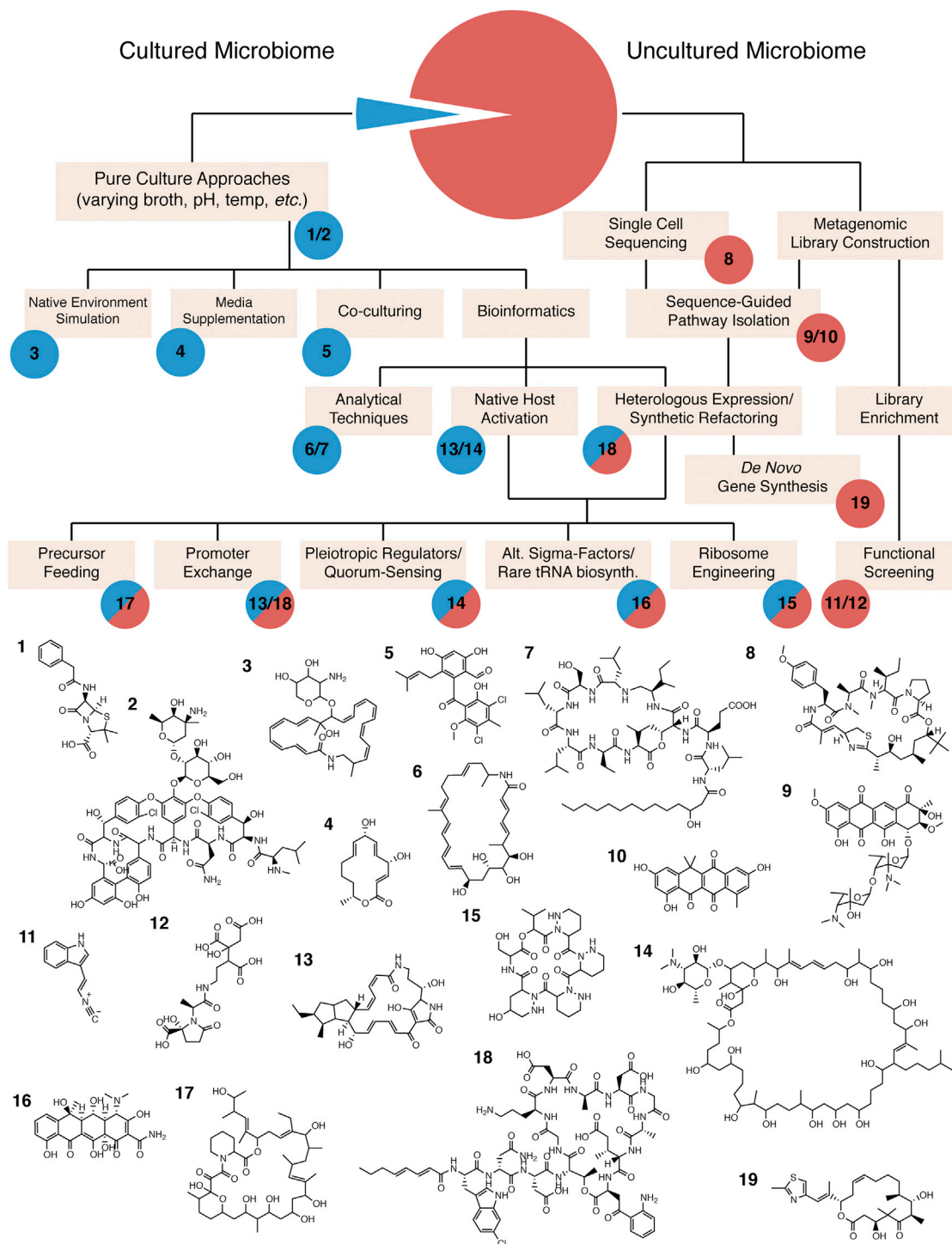


Figure 1. Flow Diagram of the Approaches Used for Metabiome Mining for Natural Product Small Molecules from Cultured and Uncultured Bacteria and Examples of Molecules Identified by Various Techniques

(1) Penicillin G; (2) Vancomycin; (3) NOVO4; (4) Mutolide; (5) Pestalone; (6) Salinilactam A; (7) Orfamide A; (8) Apratoxin A; (9) Arimetamycin A; (10) Tetarimycin A; (11) Isocyanide; (12) Vibrioferrin; (13) 6-*epi*-alteramide A; (14) Stambomycin A; (15) Piperidamycin A; (16) Oxytetracycline; (17) Meridamycin A; (18) Taromycin A; (19) Epothilone C.

several cryptic antibiotic-producing clusters by acting on a pleiotropic transcriptional repressor DasR in *Streptomyces* in minimal media (Rigali et al., 2008). The short chain fatty acids sodium

butyrate and valproic acid, known histone deacetylase (HDAC) inhibitors, have been used to induce expression of biosynthetic genes from *Streptomyces coelicolor* under normally

nonpermissive conditions (Moore et al., 2012). Although HDAC homologs are widespread in bacterial species, the mechanism of secondary metabolism upregulation by HDAC inhibitors is not yet understood. Interestingly, a screen of the Canadian Compound Collection for small molecules that upregulate pigment production in *S. coelicolor* identified at least one set of molecules that function by inhibiting fatty acid biosynthesis (Craney et al., 2012). Other enzyme inhibitors have also been observed to influence activation of biosynthetic pathways. In the fungus *Sphaeropsis* species (singular) (sp.) F-24'707, supplementation with tricyclozole, a 1,8-dihydroxynaphthalene biosynthesis inhibitor, resulted in the rediscovery of mutolide (4), a rare fungal macroide originally discovered by UV mutagenesis of the wild-type strain (Bode et al., 2000, 2002).

Coculturing has been yet another effective approach for the induction of cryptic secondary metabolite pathways. As with media supplementation, it aims to elicit production of secondary metabolites in a target organism by providing a missing regulatory cue. Often, this technique has employed two specific bacterial strains that are known to interact. This was the case for identification of pestalone (5), which was produced by a marine fungus of *Pestalotia* sp. only in the presence of a marine deuteromycete CNL-365 (Cueto et al., 2001) and for athepsin K inhibitors F-9775A and F-9775B, whose production was induced in *Aspergillus nidulans* in the presence of soil actinomycetes (Cueto et al., 2001; Schroeckh et al., 2009). A more general method was recently developed by Onaka et al. who found that mycolic acid-producing bacteria *Tsukamurella pulmonis* TP-B0596, *Rhodococcus erythropolis*, and *Corynebacterium glutamicum* alter metabolite biosynthesis in over 85% of *Streptomyces* strains tested (Onaka et al., 2011). Although these percentages reflect the combined increase and decrease in secondary metabolite production, these three species were sufficient to induce biosynthesis of over 35, 34, and 23 different metabolites, respectively.

Exploiting Genomic Sequence Data in Cultured Bacteria

While fermentation-based activation approaches represent largely trial and error efforts to activate cryptic metabolism, a number of groups have worked to take advantage of full genome sequence data to carryout more directed gene cluster induction and molecule detection studies. The ready availability of sequence data has led to the development of user-friendly bioinformatics tools, like antiSMASH, NP.searcher, and ClustScan, which identify biosynthetic gene clusters in assembled genomes and predict structural elements of their molecular products based upon substrate specificities of their conserved biosynthetic domains (Li et al., 2009; Medema et al., 2011; Starcevic et al., 2008). This type of analysis is now routinely applied to newly assembled genomes, allowing researchers to rapidly prioritize silent gene clusters for molecule discovery efforts. For example, bioinformatic analysis of the draft genome of *Salinispora tropica* CNB-440 predicted the presence of biosynthetic clusters for a broad range of unknown polyketide synthase systems, as well as nonribosomal peptide synthetases and hybrid clusters (Udwy et al., 2007). Structure predictions afforded by bioinformatics are likely to become an ever more important part of future efforts to identify structurally and functionally novel natural products.

In addition to revealing the presence of silent biosynthetic gene clusters in cultured bacteria, bioinformatic analysis of sequence data is an effective tool for identification and characterization of their undetected products. For example, genomic sequence analysis of *Salinispora tropica* CNB-440 predicted that a cryptic 10-module PKS gene cluster would incorporate polyene units into the final product, producing characteristic UV absorption. Subsequent screening of culture broth extracts by UV absorption led to the isolation of a series of new polyene macrolactams, exemplified by salinilactam A (6) (Udwy et al., 2007). In a different study, bioinformatic analysis of silent NRPS and PKS-NRPS hybrid gene clusters encoded in the *Pseudomonas fluorescens* Pf-5 gene enabled the prediction of the amino acid precursors used as substrates by these cryptic clusters. By feeding stable isotope labeled precursors to *P. fluorescens* Pf-5 cultures and then tracking their incorporation by 2D nuclear magnetic resonance, this well studied bacterium was found to produce the orfamides (7), a novel family of macrolytic lipopeptides (Gross et al., 2007). Similarly, bioinformatic prediction of a cryptic tetrapeptide siderophore sequence in the genome of the industrially important bacterium *Saccharopolyspora erythraea* facilitated the use of radiolabeled precursors to identify the hydroxamate-type siderophore antibiotic erythrochelin A (Robbel et al., 2010).

In the case of NRPS peptides, a systematic genome mining technique, termed natural product peptidogenomics (NPP) has been developed that combines the predictive power of bioinformatics with the detection sensitivity of tandem mass spectrometry (MSⁿ) (Kersten et al., 2011). This method extends the repertoire of imaging mass spectrometry (IMS) based techniques that have been long used to study bacterial chemistry and, in particular, antibacterial peptides (extensively reviewed in Watrous and Dorrestein, 2011). It relies on comparing the peptide structures, determined from the MSⁿ fragmentation patterns, with the sequence-based predictions to match the secondary metabolites present in the sample with the biosynthetic clusters responsible for their production. In proof of principle NPP experiments, detection of novel ribosomal and nonribosomal peptide natural products by MALDI-TOF imaging combined with de novo MSⁿ structure elucidation allowed researchers to link known molecules to the gene clusters responsible for their production in previously studied bacteria, as well as identify 14 new peptide natural products from various sequenced streptomycetes (Kersten et al., 2011). Moreover, this technique was shown to work for parallel identification of three new peptide natural products from daptomycin-producing *Streptomyces roseosporus* NRRL 15998, demonstrating its potential for high-throughput small molecule discovery. More recently, a similar approach has been applied to glycosylated metabolites with the development of natural product glyco-genomics. This technique further extends the application of IMS to the identification of novel glycosylated natural products and their corresponding biosynthetic gene clusters via MSⁿ characterization of O- and N-glycosyl groups (Kersten et al., 2013).

Developing Culture Independent Approaches, Metagenomics

Metagenomics provides the means of accessing bioactive small molecules synthesized by environmental bacteria through the

direct cloning of DNA from essentially any environmental source, thereby bypassing the requirement of cultivation. Metagenomic approaches for natural product discovery fall into two general categories that are generically referred to as sequence-based and function-based. As the names imply, sequence-based methods rely on the sequencing and subsequent bioinformatics analysis of a metagenomic sample to identify biosynthetic gene clusters of interest, while functional methods rely on the direct functional screening of metagenomic clones for discrete activities to identify clones of interest.

Sequence-Driven Metagenome Mining

Many of the bioinformatics tools employed for genome mining have also been successfully applied to partially assembled DNA sequences from metagenomic sources to predict natural product compounds encoded within the genomes of uncultured bacteria. However, high degree of sequence similarity and the repetitious nature of biosynthetic domains render the assembly of sequences corresponding to natural product gene clusters from metagenomic sequences difficult, and this has limited these straightforward sequence-based genome mining applications to the lower complexity metagenomes of uncultured symbionts, albeit with considerable success. Using a combination of targeted PCR screening of crude DNA isolates from zooids inhabiting the marine tunicate *Lissoclinum patella* and shotgun sequencing of the fraction containing genes associated with patellazole biosynthesis, a symbiotic α -proteobacterium *Candidatus Endolissoclinum faulkneri* was identified as a previously unknown source for a family of these potent cytotoxins (Kwan et al., 2012). Whole genome sequencing (WGS) of the lichen *Peltigera membranacea* led to the identification of nosperin, a novel polyketide of the pederin family of anticancer compounds, produced by its photobiont *Nostoc* sp. (Kampa et al., 2013). This finding expanded both the structural diversity and the known distribution of this family of secondary metabolites. In another example, a biosynthetic pathway encoding for the clinically promising cytotoxin apratoxin (**8**) was identified from a complex marine microbiome associated with the cyanobacterium *Lyngbya bouillonii* by physical micromanipulation of the symbiont to dislodge individual cells and whole genome amplification (WGA) to obtain partial genomic sequence from single cells (Grindberg et al., 2011). In addition to identifying new metabolites, metagenomic sequence data assembly recently led to the identification of a “metabolically talented” marine sponge endosymbiont genus *Entotheonella*, which is believed to be responsible for nearly all bioactive molecules that have been isolated from its host (Wilson et al., 2014).

Unfortunately, even partial genome assembly has been difficult for metagenomes of higher complexity, which may contain as many as 10^4 – 10^5 unique species, as is the case for a typical soil metagenome (Pop, 2009; Rappé and Giovannoni, 2003; Torsvik et al., 1990a). However, the biosynthetic capacity of complex environments can be mined using a PCR-based “sequence tag” approach (Seow et al., 1997). Sequence tag approaches take advantage of the fact that much of the biosynthetic diversity arises from a relatively small number of biosynthetic classes (e.g., nonribosomal peptide synthase [NRPS], polyketide synthase [PKS], isoprene, sugar, shikimic acid, alkaloid, and ribosomal peptide) related through common, highly

conserved biosynthetic domains (Dewick, 2009) and focuses next-generation sequencing efforts on PCR amplicons of the most commonly used biosynthetic domains (i.e., sequence tags). In general, gene clusters encoding structurally related metabolites are predicted to share common ancestry and, therefore, high sequence identity among conserved domains. Correspondingly, a high degree of identity among sequence tags predicts that the biosynthetic domains they correspond to are involved in biosynthesis of structurally related small molecules. By extension, sequence tags with no close relatives might be predicted to arise from gene clusters encoding structurally novel classes of metabolites (Reddy et al., 2014). These correlations form the basis of bioinformatics tools, eSNaPD and NaPDoS, developed to facilitate functional classification of novel secondary metabolite biosynthetic gene clusters from metagenomic data (Owen et al., 2013; Reddy et al., 2014; Ziemert et al., 2012). Remarkably, short sequence tags (~400 base pairs) accurately predict the structural family of the molecule encoded by a gene cluster, as validated by the recovery and subsequent analysis of metagenomic DNA clones predicted to produce novel glycopeptide, lipopeptide, and bis-intercalator natural products (Owen et al., 2013).

The sequence tag approach is a powerful tool for rapid, low cost profiling of biosynthetic capacity of (meta)genomes, without the requirement for complete genome sequencing or chemical isolation. It has been used to genotype uncultured marine *Streptomycetaceae*, as well as to profile a collection of cultured *Actinobacteria* from marine sediment (Gontang et al., 2010; Prieto-Davó et al., 2013), and it is scalable to profiling metagenomes of any complexity. The true utility of the sequence tag approach, though, is in its ability to identify gene clusters encoding congeners of valuable bioactive compounds and potentially novel chemistries that have remained hidden in the environment (Charlop-Powers et al., 2014; Owen et al., 2013; Reddy et al., 2014). In a direct comparison of NRPS and PKS biosynthetic domain detection from metagenomic samples, this PCR-based method was shown to be 10 to 100 times more sensitive than shotgun sequencing in identifying unique sequences (Woodhouse et al., 2013). Sequence tag-based profiling of soil metagenomes revealed thousands of unique adenylation and ketosynthase domains, with only minor overlap among distinct microbiomes (Charlop-Powers et al., 2014; Owen et al., 2013). Profiling of environmental samples can be performed on purified metagenomic DNA, facilitating identification of the most biosynthetically rich samples for library construction.

The sequence tag strategy has been used to identify and heterologously express gene clusters from both soil and marine environments that encode known natural products, congeners of high value natural products, and completely novel natural products. This includes the tryptophan dimer-based cytotoxins borregomycin and erdasporine (Chang and Brady, 2013; Chang et al., 2013); congeners of a potent antitumor compound pederin (Zimmermann et al., 2009); the pharmacologically important antitumor polyketide psymberin (Fisch et al., 2009); arimetamycin A (**9**), a natural doxorubicin analog, that is active against multidrug resistant cancer cell lines (Kang and Brady, 2013); a number of variants of the highly selective protease inhibitor microviridin (Gatte-Picchi et al., 2014; Ziemert et al., 2010) and structurally diverse aromatic polyketides including the MRSA active,

tetramycin A (**10**), as well as the MRSA and VRE active fasamycins A and B (Feng et al., 2012; Kallifidas et al., 2012); and a penatacyclic compound erdacin, with a novel ring system (King et al., 2009).

Functional Metagenome Mining

Sequence-based (meta)genome mining is rapidly becoming a relatively mature field with a plethora of tools at its disposal for identifying, capturing, and expressing cryptic clusters from sequenced genomes and novel biosynthetic clusters from metagenomes. In contrast, functional metagenomics methods have developed more slowly. Functional approaches aim to accelerate the rate of bioactive small molecule discovery from metagenomes by directly screening environmental DNA clones for bioactivities of interest. These approaches are akin to high-throughput screening efforts of broth extracts from culture collections, with the added dimension of roughly two to three orders of magnitude deeper biosynthetic diversity potential (Rappé and Giovannoni, 2003; Torsvik et al., 1998).

Early Functional Metagenomic Studies and Their Derivatives

To date, functional metagenomics has largely consisted of very simple screens performed on cosmid libraries hosted in a small number of model bacterial hosts. In one of the earliest efforts, direct high-performance liquid chromatography-electrospray ionization mass spectrometry screening of fermentation broths derived from a small (1,020-member) soil DNA library hosted in *Streptomyces lividans* yielded terragline A and a family of related molecules, the first novel family of small molecules from a metagenomic source (Wang et al., 2000). It quickly became apparent that this kind of screen was impractical for the larger metagenomic libraries that would be required in order to identify diverse molecules from metagenomic sources. Consequently, the majority of functional assays for secondary metabolites have since relied on easily identifiable phenotypic traits that are commonly associated with secondary metabolite production, such as pigmentation and antibiosis. Screening of approximately 700,000 clones from soil metagenomic mega-library in *E. coli* for antibiosis against *B. subtilis* in overlay assays produced a large family of long-chain N-acyl amino acids, which were the first novel antibiotics discovered using functional metagenomics (Brady and Clardy, 2000). The same approach was used to identify long-chain N-acyl palmitoylputrescine from a bromeliad tank water metagenome (Brady and Clardy, 2004) and an isocyanide antibiotic (**11**) from a soil metagenome (Brady and Clardy, 2005). In a screen of soil metagenomic libraries, hosted in *E. coli*, a combination of pigmentation and activity against *B. subtilis* helped identify metagenomic clones encoding production of indirubin, indigo blue, and other related molecules (Lim et al., 2005; MacNeil et al., 2001). In a chrome azurol S (CAS) colorimetric assay (Schwyn and Neilands, 1987), designed to report on siderophores production, new gene clusters encoding known siderophores vibrioferrin (**12**) and bisucaberin were found in libraries constructed from marine sources (Fujita et al., 2011, 2012).

Improving Functional Access to Natural Products

Functional metagenomics is a potentially powerful approach for identification of bioactive natural products encoded by environ-

mental microbiomes in a high-throughput manner. However, to reach the full potential of this approach a number of methodological advances are required. Improvements in the size of cloned environmental DNA are needed to facilitate the capture of complete gene clusters on individual clones. Diversification of the heterologous hosts is required to enable native expression of a more diverse collection of genes clusters. Finally, enrichment of metagenomic libraries for biosynthetic clones is necessary to increase the efficiency of the screening process.

A number of groups have recently reported advances geared at solving these problems. Several groups have reported bacterial artificial chromosome/clone (BAC) libraries from soil samples, with average insert sizes of up to 90 kb (Liles et al., 2008; Liu et al., 2010; Yu et al., 2008); however, the reported libraries remain small in size and are not yet capable of saturating coverage of the environments from which they are derived. A variety of broad host-range vectors have been developed to facilitate heterologous expression in diverse model hosts. For example, vectors based on Φ C31 and Φ BT1 phage, site-specific integrase systems are able to facilitate integrative transfer of cloned DNA into a variety of *Streptomyces* spp. (Gregory et al., 2003; Kuhstoss and Rao, 1991). For transfer of libraries into diverse alpha-, beta-, and gammaproteobacterial species, several RK2-derived vectors have been recently developed (Aakvik et al., 2009; Kikirde et al., 2011). The utility of these vectors was demonstrated by parallel functional metagenomic library screens carried out in *Agrobacterium tumefaciens*, *Burkholderia graminis*, *Caulobacter vibrioides*, *E. coli*, *Pseudomonas putida*, and *Ralstonia metallidurans*, using a single broad-host-range vector. This study identified orthogonal sets of clone-specific small molecules expressed in each strain with minimal overlap (Craig et al., 2010). On average, less than 2% of the bacterial metagenome is devoted to secondary metabolism (Garcia et al., 2011), which means that metagenomic libraries are only sparsely populated with gene clusters encoding bioactive small molecules. Increasing the biosynthetic density of metagenomic libraries will therefore translate directly into more efficient screening. The majority of existing enrichment techniques rely on some variation of probe hybridization (Kalyuzhnaya et al., 2006; Meyer et al., 2007; Morimoto and Fujii, 2009; Uchiyama et al., 2005; Zhang et al., 2009). Recently, enrichment strategies based on genetic complementation have shown promise as tools for library scale enrichments. In particular, the promiscuity of 4'-phosphopanthetheinyl transferases and their universal requirement to generate holo-NRPS and PKS enzymes has been used to develop successful gene cluster enrichment strategies (Charlop-Powers et al., 2013; Owen et al., 2012; Yin et al., 2007). While all of these methods remain works in progress, recent advances in the field suggest that functional metagenomics may one day reach the promise envisioned for it at its inception (Handelsman et al., 1998).

Activating Silent Pathways through Host and Cluster Engineering

From minimally invasive techniques that employ minor modifications to the native host to induce gene cluster activation, to more invasive heterologous expression and gene cluster refactoring approaches, molecular biology, bacterial genetics, and more recently synthetic biology have proved essential to many

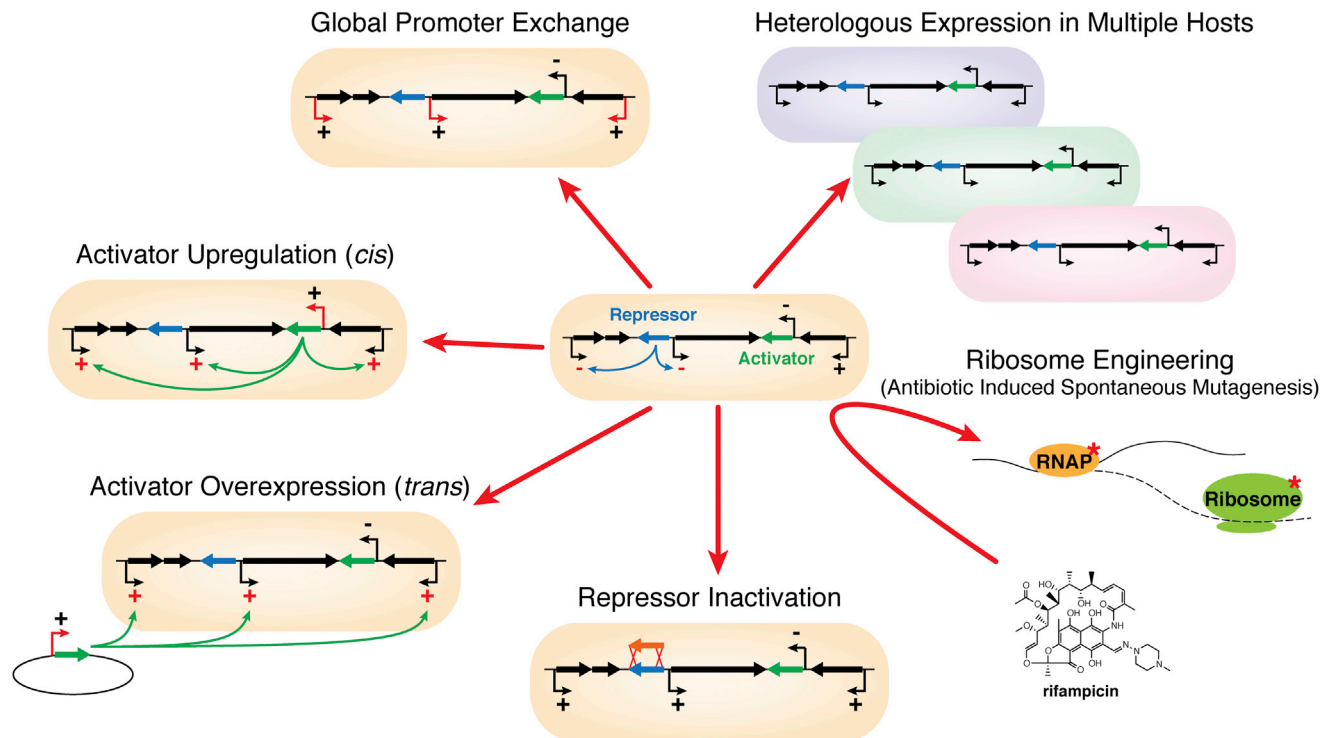


Figure 2. Common Approaches for Activation of Cryptic Biosynthetic Gene Clusters

aspects of the modern natural product discovery process. When applied to the natural product discovery, these methods represent a diverse and growing toolbox for activating biosynthetic pathways that are identified by either genomic approaches in cultured bacteria (i.e., cryptic) or metagenomic approaches (Figure 2).

Cryptic Cluster Activation in Native Hosts

The simplest and the most effective approaches to activating cryptic secondary metabolite pathways rely on minimal modifications made directly in the native host organism. When sequence data are available it is possible to perform a detailed bioinformatic analysis of the cryptic cluster of interest and design rational strategies for their potential activation. Strategic insertion of strong constitutive promoters in front of either biosynthetic or regulatory genes in cryptic clusters has been one such productive strategy for activating biosynthesis. For example, insertion of a strong constitutive *ermE*⁺*p* promoter from *Saccharopolyspora erythraea* into bioinformatically identified cryptic clusters in the well studied strain *S. albus* J1074 led to the expression of an NRPS encoded blue pigment and two novel compounds 6-*epi*-alteramide A (**13**) and B, encoded by a hybrid type I PKS-NRPS cluster (Olano et al., 2014). In cases where putative transcription activators are identified in a cluster, induction of their constitutive expression has been shown to activate an entire cluster. For example, cloning out the large ATP-binding LuxR-type regulatory gene and reintroducing it under control of a constitutive *ermE*⁺*p* led to identification of four novel, 51-membered, glycosylated macrolides stambomycins A-D (**14**) with activity against human cancer lines in *Streptomyces ambo-*

faciens ATCC23877 (Laureti et al., 2011). Not all regulatory elements are inducers of secondary metabolism, as was observed when mutagenic inactivation of the quorum-sensing regulator in *Burkholderia thailandensis* led to activation of a cryptic cluster encoding a novel polyketide metabolite, thailandamide lactone (Ishida et al., 2010).

Ochi and coworkers have worked on a more general, sequence-agnostic method for cryptic cluster activation by exposure of bacteria of interest to antibiotics and selecting for drug-resistant mutants. With this approach, which they have termed “ribosome engineering,” they observed that mutations in the endogenous RNA polymerase and ribosomal protein S12 lead to strong activation of cryptic clusters and improve antibiotic expression in *Streptomyces* spp., myxobacteria, and fungi (reviewed in Ochi and Hosaka, 2013). This work led to the identification of 3,3'-neotrehalosadiamine, a novel aminosugar antibiotic, in *B. subtilis* and piperidamycin (**15**), a novel antibacterial class, in *Streptomyces* sp. 631689 (Hosaka et al., 2009; Inaoka et al., 2004).

Capturing Gene Clusters for Heterologous Expression in Model Hosts

To take advantage of many newly developed natural products discovery approaches it is often necessary to clone a gene cluster from its native host for subsequent introduction into a heterologous host. Secondary metabolite gene clusters vary in size from less than 10 kb to over 150 kb, with an average size around 50 kb (Starcevic et al., 2008), and thus cloning using standard “cut and paste” techniques can be challenging. If the sequence of the target gene cluster is known, a variety of

recombination-based techniques (e.g., transformation-assisted recombination [TAR] in the budding yeast *Saccharomyces cerevisiae* [Kim et al., 2010; Larionov et al., 1997], Red/ET recombination in *E. coli* [Bian et al., 2012; Fu et al., 2012; Muylers et al., 2001], and *oriT*-directed capture [Chain et al., 2000; Kvitko et al., 2013]) can now be used to directly clone intact gene clusters from genomic DNA. For metagenomic samples, which are significantly more complex than individual genomes and lack sequence data, metagenomic cosmid or fosmid libraries containing 30–40 kb inserts are first constructed and large gene clusters are assembled using TAR from overlapping metagenomic clones (Kim et al., 2010). Theoretically BAC libraries, capable of carrying inserts of up to 250 kb, can be used to circumvent this limitation; however, as mentioned above, more efficient cloning methodologies are needed to generate useful BAC libraries from biosynthetically rich metagenomic source.

Host Modifications

At the host level, problems with heterologous expression can arise from issues associated with transcription and translation of the exogenous DNA and from insufficient availability of precursors. Most studies addressing translation and transcription issues have focused on *E. coli*, as it is a rapidly growing and genetically tractable organism. Overexpression of an alternative sigma factor, σ^{54} , in *E. coli* enabled expression of a heterologous oxytetracycline polyketide (**16**) encoded by a type II PKS gene cluster (Stevens et al., 2013). In addition, *E. coli* strains expressing rare tRNAs have been engineered to alleviate translation problems (e.g., stalling, premature termination, and frameshifts) due to the imbalance of available tRNAs caused by the differences in codon usage (Novy et al., 2001). In the same vein, expression of chaperones to facilitate proper folding of biosynthetic enzymes has been beneficial for heterologous expression studies in *E. coli* (Hoertz et al., 2012; Mutka et al., 2006). Limited precursor availability can often be a result of the heterologous host not producing sufficient amounts of the precursor or of precursor depletion by endogenous biosynthetic pathways. Precursor feeding has been used successfully to increase production of meridamycin (**17**), a 27-membered macrolide with potent neuroprotective properties, in *S. lividans* (Liu et al., 2009). Whereas, deletion of the competing pathways has generally led to a dramatic increase in the levels of heterologous expression of secondary metabolites in *S. lividans* (Pestov et al., 1999), *S. avermitilis* (Komatsu et al., 2010), and *S. coelicolor* (Gomez-Escribano and Bibb, 2011).

Refactoring of Cloned Gene Clusters

At the gene cluster level, lack of recognition of promoters and ribosome binding sites by the heterologous host frequently leads to low or no expression of the biosynthetic enzymes and a corresponding lack of small molecule expression. This challenge is most commonly overcome by replacing the native transcriptional regulatory elements within the biosynthetic gene cluster with ones known to function in the heterologous host. For small clusters, this can be accomplished by PCR amplification of individual genes and subcloning them under the control of such promoters, a strategy that yielded a family of novel tryptophan dimer cytotoxins erdasporines A–C from a

metagenomic clone refactored for expression in *E. coli* (Chang et al., 2013). For larger clusters, Red/ET- and TAR-based tools can be used to replace promoters driving the expression of each polycistronic operon. Using Red/ET recombineering in *E. coli*, the myxochromide S biosynthetic gene cluster from myxobacteria *Stigmatella aurantiaca* was placed under the control of the toluic acid inducible P_m promoter suitable for use in pseudomonads. Transfer of this construct into *Pseudomonas putida* resulted in high expression level of myxochromide S, as well as two new derivatives that have not been previously observed (Wenzel et al., 2005). In two examples of promoter replacement in a multioperon cluster, a powerful plug-and-play “DNA assembler” method based on homologous recombination in yeast (Shao et al., 2009, 2011) was used to activate and characterize cryptic clusters encoding four novel polycyclic tetramate macrolactams from *S. griseus* (Luo et al., 2013) and a daptomycin-like dichlorinated lipopeptide antibiotic taromycin A (**18**) from the marine actinomycete *Saccharomonospora* sp. CNQ-490 (Yamanaka et al., 2014).

Success of complete refactoring of biosynthetic gene clusters as a strategy for heterologous activation of cryptic metabolism prompted the development of new tools to simplify and accelerate this process. Gibson (Gibson et al., 2009) and Golden Gate (Engler et al., 2008) assembly techniques allow rapid assembly of multiple PCR generated fragments in a specific order determined by the user-designed overlapping sequences. These methods are particularly useful for assembling codon-optimized synthetic constructs into functional gene clusters. Seamless ligation cell extract is an *ex vivo* recombineering method that allows both cloning and genetic manipulation of large DNA fragments using cell extracts from an *E. coli* strain expressing an optimized lambda prophage Red/ET system (Zhang et al., 2012). This method accelerates the process by eliminating multiple transformation and cell growth steps. Also recently introduced is the clustered regularly interspaced short palindromic repeats (CRISPR)/Cas system, which functions via an RNA-guided nuclease that, in isolation, can be programmed to target any sequence by providing a short (20–50 nucleotide) synthetic RNA sequence (Wiedenheft et al., 2012). It offers ease of use, high specificity, simplicity, and speed for targeting virtually any DNA sequence. These characteristics have led to an immense amount of interest in developing a variety of genome editing tools based on CRISPR/Cas system, which are certain to revolutionize synthetic biology (Baker, 2014). Although these latter tools have not yet yielded novel molecules, they are very likely to turn out many new natural products in the coming years.

Even with appropriate transcriptional elements in place, the DNA sequence itself varies significantly among species, and differences in GC content and codon usage can lead to decreased translation efficiency, early transcription termination, and self-resistance, if the final product is toxic to the host (Gustafsson et al., 2004). *De novo* synthesis with codon optimization and refactoring is possible, as the most aggressive approach to cryptic cluster activation, but has predominantly been used to increase yields of existing clinically important compounds by engineering their production in a heterologous host. In one such example, the gene cluster encoding production of the promising anticancer metabolites epothilone C and D (**19**)

from the slow growing and genetically intractable myxobacterium *Sorangium cellulosum* was completely resynthesized to optimize codon and promoters usage to allow production in *E. coli* (Mutka et al., 2006).

The Future of Metabiome Mining

Transformational changes in the field of DNA sequencing and molecular and synthetic biology (i.e., genome manipulation) are drastically altering the way we pursue the discovery of natural products. The ever increasing availability of sequenced genomes will undoubtedly continue to fuel targeted approaches to access cryptic clusters in cultured bacteria. Although many of the most important clinical therapeutic agents have been discovered in cultured bacteria, these organisms represent only the tip of the microbial diversity iceberg. Metagenomics is taking advantage of the sequencing revolution to bypass many of the traditional barriers to drug discovery by providing the means to profile and access the biosynthetic capacity of the environmental microbiome. Promising new technologies have the potential to extend the power of WGS to metagenomes. Long-read nano-pore-based, single molecule sequencing alleviates the problems inherent in metagenomic sequence assembly (Koren et al., 2013; Mavromatis et al., 2012), while additional new techniques can now obtain nearly complete genome sequence from a single cell (Fitzsimons et al., 2013; Lasken, 2012). At this point, a collection of heterologous expression strategies exist that appears capable of inducing expression of effectively any gene clusters, given sufficient expended effort. The focus of heterologous expression studies has also clearly shifted from fine-tuning, aimed at increasing the production of industrially important metabolites, to discovery, aimed at the detection of cryptic metabolites. In the future, the challenge will be to develop more universally applicable heterologous expression strategies, which can streamline efforts to activate the diverse cryptic gene cluster now known to be present in the Earth's microbiome.

ACKNOWLEDGMENTS

This work was supported by NIH (grant number GM077516). J.S.S. is supported by the Helmsley Postdoctoral Fellowship for Basic and Translational Research on Disorders of the Digestive System. S.F.B. is a Howard Hughes Medical Institute Early Career Scientist.

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