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Metagenomics: Application of Genomics to Uncultured Microorganisms

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Metagenomics: Application of Genomics to Uncultured Microorganisms

Jo Handelsman*

Department of Plant Pathology, University of Wisconsin-Madison, Madison, Wisconsin

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INTRODUCTION

Microbiology has experienced a transformation during the last 25 years that has altered microbiologists' view of microorganisms and how to study them. The realization that most microorganisms cannot be grown readily in pure culture forced microbiologists to question their belief that the microbial world had been conquered. We were forced to replace this belief with an acknowledgment of the extent of our ignorance about the range of metabolic and organismal diversity.

This change fomented a revolution in microbiological thought. At the heart of this revolution was the convincing demonstration that the uncultured microbial world far outsized the cultured world and that this unseen world could be studied (105–108). This change in thinking was prompted by another, equally important realization: microorganisms underpin most of the geochemical cycles and many human health conditions that were previously thought to be driven by inorganic processes and stress, respectively. The glimmers of insight into the influence that microorganisms exert on the world propelled microbiologists to pursue the uncultured

world. In 1931, Waksman optimistically believed that "a large body of information has accumulated that enables us to construct a clear picture...of...the microscopic population of the soil" (145), and in 1923 *Bergey's Manual* stated categorically that no organism could be classified without being cultured (133).

By the mid-1980s, however, microbiologists had lost this confidence, and the language and practice of microbiology changed to accommodate the vast unknown of uncultured life. Concepts, assumptions, images, and words needed to be replaced when it became evident that they were based upon the premise that microorganisms did not exist unless they could be cultured. Pace and colleagues highlighted the need for nontraditional techniques to understand the microbial world: "The simple morphology of most microbes provides few clues for their identification; physiological traits are often ambiguous. The microbial ecologist is particularly impeded by these constraints, since so many organisms resist cultivation, which is an essential prelude to characterization in the laboratory" (107).

In the ensuing years, microbiologists dedicated intense effort to describing the phylogenetic diversity of exotic and ordinary environments—ocean surfaces, deep sea vents, hot springs, soil, animal rumen and gut, human oral cavity and intestine. Many new lineages were classified based on their molecular signatures alone. The next challenge was to elucidate the func-

^{*} Mailing address: Department of Plant Pathology, University of Wisconsin-Madison, Madison, WI 53706. Phone: (608) 263-8783. Fax: (608) 265-5289. E-mail: joh@plantpath.wisc.edu.

tions of these new phylotypes and determine whether they represented new species, genera, or phyla of prokaryotic life. This challenge spawned various techniques, including metagenomics, the genomic analysis of assemblages of organisms. In a few years, the study of uncultured microorganisms has expanded beyond asking "Who is there?" to include the difficult question "What are they doing?"

The outcomes of the recognition of uncultured microorganisms are worthy of examination. One of these outcomes, metagenomics, is further shaping microbiology. Metagenomics has already opened new avenues of research by enabling unprecedented analyses of genome heterogeneity and evolution in environmental contexts and providing access to far more microbial diversity than has been viewed in the petri dish. This review will explore the origins of metagenomics and examine its recent application to microbial ecology and biotechnology.

HISTORY OF THE CULTURE DIVIDE

The current excitement about the uncultured world may make students of modern microbiology wonder why this aspect of microbiology was largely ignored for so long. It is worth tracing the origins of microbiology, which did not rely on culturing, and examine the reasons for the shift to culturing and the subsequent discoveries that rekindled interest in the uncultured world. This review will use the term uncultured microorganisms to capture the entire spectrum of organisms that are not cultured in a specific experiment. These may include microorganisms that we have not attempted to culture and those that have been resistant to culturing efforts but may submit to culturing in the future.

Early Microbiology and the Microscope

The roots of microbiology are firmly associated with the microscope. The first record of a human being's seeing a bacterial cell is in 1663. Antonie van Leeuwenhoek watched bacteria that he recovered from his own teeth through his homemade microscope. He was a keen observer and an outstanding maker of microscopes, and his observations and detailed illustrations of microbial life prompted many other observers (both scientists and nonscientists) to take an interest in the microscopic world. His colorful descriptions of bacteria made their study compelling; in his descriptions of the many shapes of the bacteria he sampled from his teeth, he marveled that one "shot through the water like a pike does through water" (30), firmly establishing that these tiny objects were, indeed, alive. For 200 years, microscopy enabled microbiologists to view heterotrophs, autotrophs, and obligate parasites alike.

Among the advances during this period of microbiology was the work of botanist Ferdinand Cohn, who classified many bacteria and described the life cycle of *Bacillus subtilis* based on his microscopic observations (60). Although mycologists such as Franz Unger had understood the concept of pure culture as early as the 1850s, it was in large part the emphasis on disease causality that solidified pure culture as the standard bacteriological technique for laboratory microbiology (49, 96). Robert Koch's postulates and his own innovation in developing culture media were instrumental in this shift, and from the 1880s forward, the microbiological world was divided into the

cultured and the uncultured. Microbiologists were attracted to the power and precision of studies of bacteria in pure culture, and as a result, most of the knowledge that fills modern microbiology textbooks is derived from organisms maintained in pure culture.

Modern Microbiology—a Pure Culture Is Not Enough

Because culturing provided the platform for building the depth and detail of modern microbiological knowledge, for a long time microbiologists ignored the challenge to identify and characterize uncultured organisms. They focused instead on the rich source of discovery found in the readily cultured model organisms, and this contributed to the explosion of knowledge in microbial physiology and genetics in the 1960s to mid-1980s. Meanwhile, the study of uncultured microorganisms remained in the hands of a few persistent scientists who began to accumulate hints that flitted at the edge of the microbiological consciousness, suggesting that culturing did not capture the full spectrum of microbial diversity.

One of the indicators that cultured microorganisms did not represent much of the microbial world was the oft-observed "great plate count anomaly" (135)—the discrepancy between the sizes of populations estimated by dilution plating and by microscopy. This discrepancy is particularly dramatic in some aquatic environments, in which plate counts and viable cells estimated by acridine orange staining can differ by four to six orders of magnitude (66), and in soil, in which 0.1 to 1% of bacteria are readily culturable on common media under standard conditions (138, 139).

Brock and colleagues encountered microorganisms in Yellowstone hot springs that could not be cultured and others whose behavior in culture did not reflect their activities in situ. Many of the organisms could not be cultured on agar medium because their temperature requirements exceed the melting point of the agar. Therefore, elucidating the physiological function of microorganisms without culturing them required ingenuity. Brock's central technique involved the immersion of microscope slides in the spring for 1 to 7 days, followed by microscopic examination and often staining with fluorescent antibodies raised against cultured members of the taxonomic groups suspected to inhabit the environment (17, 21). This approach estimated in situ population sizes and growth rates, which indicated, for example, that certain strains of Sulfolobus grew in the hot springs at temperatures well below the optima in pure culture (103). The expanding body of evidence indicating that it was imperative to study physiology in the environment led Brock's group to determine which organisms in the hot spring were responsible for photosynthesis. To do so, they placed an opaque cover over the spring for a week. The spring lost its pink color, leading them to infer that the genus Synechococcus, typically pink in culture, was a major contributor to photosynthesis (25, 26).

Further evidence that drew attention to the uncultured world accumulated during the 1970s and 1980s. A study of oligotrophs indicated that incubation times longer than 25 days enhanced the recovery of certain organisms in culture (147). The food industry generated intense interest in "injured bacteria" in food—live organisms that cannot be cultured following stressful treatments such as heat, chilling, or desiccation

but represent a significant risk to human health (41). The concept of organisms that were viable but not culturable emerged from the work of Colwell and colleagues, who showed that strains of *Vibrio cholerae* were indeed alive and virulent when isolated from aquatic environments (8) but did not grow in culture until after passage through a mouse or human intestine (35–37).

The confluence of these and many other scientific and technical advances steadily drew attention to the unculturable microbial world, but two discoveries figured significantly in the sharpened focus. The first was work on the diversity of soil bacteria, which demonstrated with DNA-DNA reassociation techniques that the complexity of the bacterial DNA in the soil was at least 100-fold greater than could be accounted for by culturing. This work suggested that the diversity of the uncultured world exceeded previous estimates (138). The second discovery was the demonstration that Helicobacter pylori causes gastric ulcers and cancer. Although spiral bacteria had been observed in the gastric mucosa of dogs in 1893 and in humans in 1906 (29), and correlations between the appearance of the bacteria and peptic ulcers were noted in 1938 (48), it was not until H. pylori was cultured that its role in disease was accepted (94, 95). Culturing was accompanied by the satisfaction of Koch's postulates on a human volunteer (94), providing definitive evidence for the causal relationship between the bacterium and ulcers.

Ironically, culturing was not that difficult. Plates accidentally incubated for 5 days instead of 3 revealed colonies later shown to be *H. pylori* (29). The fact that strong microscopic evidence for the role of *H. pylori* long preceded culturing and might have served as the basis for successful treatment decades earlier, perhaps reducing human suffering and mortality due to ulcers and cancer, did not escape the notice of microbiologists, medical practitioners, and the public. Whereas the studies of the complexity of the soil DNA demonstrated the diversity of the unknown world, the connection of uncultured bacteria and ulcers provided a striking example of the power of the undetected organisms. These discoveries provided compelling evidence that drew microbiologists to wrestle with the daunting challenge of devising strategies to access these organisms.

THE PARADIGM SHIFT

In 1985, an experimental advance radically changed the way we visualize the microbial world. Building upon the pioneering work of Carl Woese, which showed that rRNA genes provide evolutionary chronometers (148), Pace and colleagues created a new branch of microbial ecology (83, 134). They used direct analysis of 5S and 16S rRNA gene sequences in the environment to describe the diversity of microorganisms in an environmental sample without culturing (107, 134). The early studies were technically challenging, relying on direct sequencing of RNA or sequencing of reverse transcription-generated DNA copies. The next technical breakthrough arrived with the development of PCR technology and the design of primers that can be used to amplify almost the entire gene (62). This accelerated the discovery of diverse taxa as habitats across the earth were surveyed by the new technique (6, 50, 62, 126).

The application of PCR technology provided a view of microbial diversity that was not distorted by the culturing bias and revealed that the uncultured majority is highly diverse and contains members that diverge deeply from the readily culturable minority. Today, 52 phyla have been delineated, and most are dominated by uncultured organisms (Fig. 1) (114). The application of phylogenetic stains (Fig. 2)—nucleic acid probes with fluorescent labels that facilitate visualization of single cells in situ—led to a recrudescence of microscopy as a central tool of microbiology and microbial phylogeny (43, 63). Whereas traditional microscopy provides little phylogenetic information and fluorescent antibody studies require prior knowledge and culturing of an organism or one closely related to it to raise antibodies (17, 18), phylogenetic stains require only an rRNA sequence, which can be derived from an environmental sample without culturing. Phylogenetic stains corroborated evidence from PCR-based studies but provided quantitative information as well, because the findings are based on direct observation that is not subject to the skewing of organism abundance potentially observed with PCR (137).

rRNA Analysis and Culturing

In addition to providing a universal culture-independent means to assess diversity, 16S rRNA sequences also provided an aid to culturing efforts. Bacteria may be recalcitrant to culturing for diverse reasons—lack of necessary symbionts, nutrients, or surfaces, excess inhibitory compounds, incorrect combinations of temperature, pressure, or atmospheric gas composition, accumulation of toxic waste products from their own metabolism, and intrinsically slow growth rate or rapid dispersion from colonies (131). Testing myriad conditions requires focus on the critical variables, is challenging and laborious, and can only succeed if there is a sufficiently quantitative assay available to determine whether the organism of interest is enriched under a specific set of conditions.

Nucleic acid probes labeled with fluorescent tags provide such an assay, facilitating quantitative assessment of enrichment and growth. As a result, culturing efforts have intensified recently, and successes have included pure cultures of members of the SAR11 clade, now termed the genus Pelagibacter (34, 38, 113), which represents more than one-third of the prokaryotic cells in the surface of the ocean but was known only by its 16S rRNA signature until 2002 (38, 102, 113). The corollary to SAR11 in terrestrial environments is the Acidobacteria phylum (76, 121). Acidobacteria are abundant in soil, typically representing 20 to 30% of the 16S rRNA sequences amplified by PCR from soil DNA, but until recently only three members had been cultured (7, 56, 79, 81, 89, 97, 129, 132). Once again, the culture-independent indications that it was prevalent in the environment led to intensive efforts to culture members of the Acidobacteria phylum. The current efforts to culture new microorganisms will be advanced by the information that metagenomics can reveal about uncultured organisms

Given that many organisms will not be coaxed readily into pure culture, a critical advance is to extend the understanding of the uncultured world beyond cataloging 16S rRNA gene sequences, and microbiologists have striven to devise methods to analyze the physiology and ecology of these diverse, uncultured organisms.

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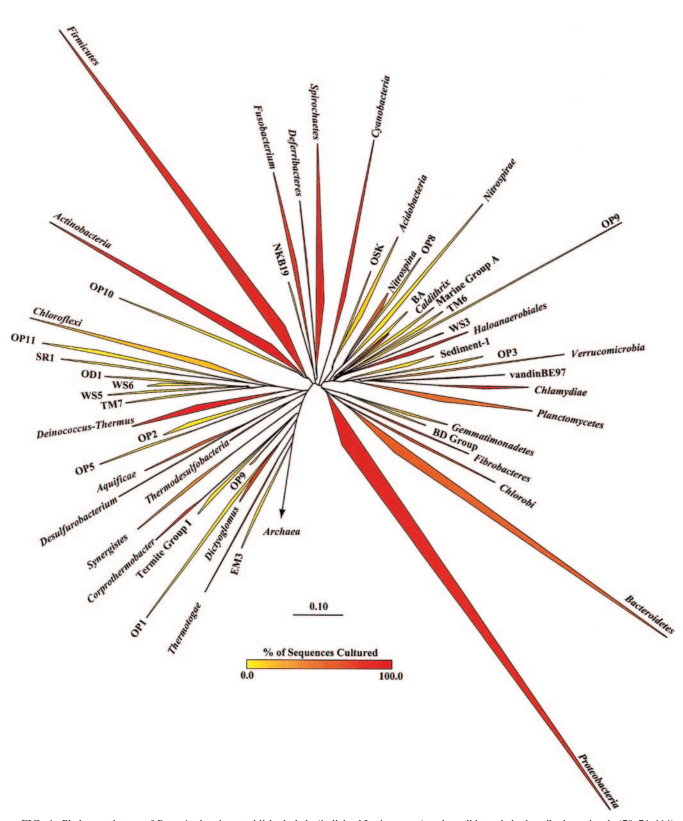


FIG. 1. Phylogenetic tree of *Bacteria* showing established phyla (italicized Latin names) and candidate phyla described previously (70, 74, 114) with the November 2003 ARB database (http://arb-home.de) (90) with 16,964 sequences that are >1,000 bp. The vertex angle of each wedge indicates the relative abundance of sequences in each phylum; the length of each side of the wedge indicates the range of branching depth found in that phylum; the redness of each wedge corresponds to the proportion of sequences in that phylum obtained from cultured representatives. Candidate phyla do not contain any cultured members.

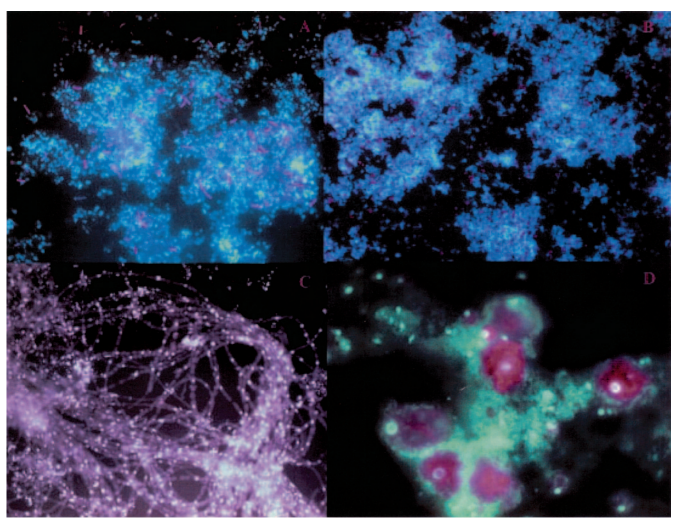


FIG. 2. Phylogenetic stains. Fluorescent in situ hybridization biofilm samples from Iron Mountain Mine, Calif. Nucleic acid probes were labeled with indodicarbocyanine, and DNA was stained nonspecifically with 4',6'-diamidino-2-phenylindole. The nucleic acid probes are specific for (top left) *Sulfobacillus* spp., (top right) *Archaea*, (bottom left) *Archaea* on fungal filaments, and (bottom right) *Eukarya*. Reproduced from reference 5 with permission of the publisher.

METAGENOMICS—CULTURE-INDEPENDENT INSIGHT

Among the methods designed to gain access to the physiology and genetics of uncultured organisms, metagenomics, the genomic analysis of a population of microorganisms, has emerged as a powerful centerpiece. Direct isolation of genomic DNA from an environment circumvents culturing the organisms under study, and cloning of it into a cultured organism captures it for study and preservation. Advances have derived from sequence-based and functional analysis in samples from water and soil and associated with eukaryotic hosts.

The word metagenomics was coined (69) to capture the notion of analysis of a collection of similar but not identical items, as in a meta-analysis, which is an analysis of analyses (64). (Community genomics, environmental genomics, and population genomics are synonyms for the same approach.) The idea of cloning DNA directly from environmental samples was first proposed by Pace (108), and in 1991, the first such cloning in a phage vector was reported (126). The next advance was the construction of a metagenomic library with DNA de-

rived from a mixture of organisms enriched on dried grasses in the laboratory (71). Clones expressing cellulolytic activity were found in these libraries, which were referred to as zoolibraries, a term that has not been used widely in the field (71). The work of DeLong's group defined the field when they reported libraries constructed from prokaryotes in seawater (136). They identified a 40-kb clone that contained a 16S rRNA gene indicating that the clone was derived from an archaeon that had never been cultured. Construction of libraries with DNA extracted from soil lagged due to difficulties associated with maintaining the integrity of DNA during its extraction and purification from a soil matrix (14, 69, 80, 118) but eventually produced analyses analogous to those from seawater (39, 72, 118).

APPROACHES TO METAGENOMIC ANALYSIS

Metagenomic analysis involves isolating DNA from an environmental sample, cloning the DNA into a suitable vector,

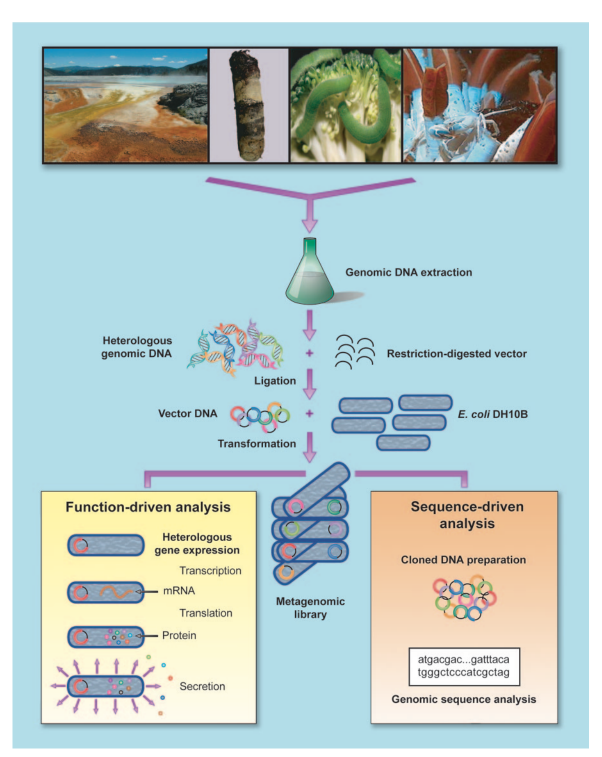


FIG. 3. Construction and screening of metagenomic libraries. Schematic representation of construction of libraries from environmental samples. The images at the top from left to right show bacterial mats at Yellowstone, soil from a boreal forest in Alaska, cabbage white butterfly larvae, and a tube worm.

transforming the clones into a host bacterium, and screening the resulting transformants (Fig. 3). The clones can be screened for phylogenetic markers or "anchors," such as 16S rRNA and *recA*, or for other conserved genes by hybridization

or multiplex PCR (136) or for expression of specific traits, such as enzyme activity or antibiotic production (39, 44, 61, 78, 87, 88, 91, 93, 118, 125), or they can be sequenced randomly (141, 142). Each approach has strengths and limitations; together

these approaches have enriched our understanding of the uncultured world, providing insight into groups of prokaryotes that are otherwise entirely unknown.

Sequence-Based Analysis

Sequenced-based analysis can involve complete sequencing of clones containing phylogenetic anchors that indicate the taxonomic group that is the probable source of the DNA fragment. Alternatively, random sequencing can be conducted, and once a gene of interest is identified, phylogenetic anchors can be sought in the flanking DNA to provide a link of phylogeny with the functional gene. Sequence analysis guided by the identification of phylogenetic markers is a powerful approach first proposed by the DeLong group, which produced the first genomic sequence linked to a 16S rRNA gene of an uncultured archaeon (136). Subsequently, they identified an insert from seawater bacteria containing a 16S rRNA gene that affiliated with the γ -Proteobacteria. The sequence of flanking DNA revealed a bacteriorhodopsin-like gene. Its gene product was shown to be an authentic photoreceptor, leading to the insight that bacteriorhodopsin genes are not limited to Archaea but are in fact abundant among the Proteobacteria of the ocean (11,

A promising application of phylogenetic anchor-guided sequencing is to collect and sequence many genomic fragments from one taxon. In more complex environments and taxa, reassembly of a genome may not be feasible, but inference about the physiology and ecology of the members of the groups can be gleaned from sequence data. This approach has been initiated with clones from diverse soils carrying 16S rRNA genes that affiliate with the *Acidobacteria* phylum, which is abundant in soil and highly diverse (7, 28, 58, 76, 121) and about which little is known (85, 112). Complete sequencing of the estimated ≈500 kb of *Acidobacterium* DNA in metagenomic libraries may provide insight into the subgroups of bacteria in this phylum that have never been cultured.

The alternative to a phylogenetic marker-driven approach is to sequence random clones, which has produced dramatic insights, especially when conducted on a massive scale. The distribution and redundancy of functions in a community, linkage of traits, genomic organization, and horizontal gene transfer can all be inferred from sequence-based analysis. The recent monumental sequencing efforts, which include reconstruction of the genomes of uncultured organisms in a community in acid mine drainage (141) and the Sargasso Sea (142), illustrate the power of large-scale sequencing efforts to enrich our understanding of uncultured communities. These studies have made new linkages between phylogeny and function, indicated the surprising abundance of certain types of genes, and reconstructed the genomes of organisms that have not been cultured. These studies will be discussed in detail in the section entitled Biogeochemical Cycles.

The use of phylogenetic markers either as the initial identifiers of DNA fragments to study or as indicators of taxonomic affiliation for DNA fragments carrying genes of interest because of their function is limited by the small number of available markers that provide reliable placement in the Tree of Life. If a fragment of DNA that is of interest for other reasons does not carry a dependable marker, its organism of origin

remains unknown. The collection of phylogenetic markers is growing, and as the diversity of markers increases, the power of this approach will also increase, making it possible to assign more fragments of anonymous DNA to the organisms from which they were isolated. Moreover, as more genomes are reconstructed, more genes will be linked to phylogenetic markers even though they were not cloned initially on the same fragment (141, 142).

Functional Metagenomics

Heterologous expression. A powerful yet challenging approach to metagenomic analysis is to identify clones that express a function. Success requires faithful transcription and translation of the gene or genes of interest and secretion of the gene product, if the screen or assay requires it to be extracellular. Functional analysis has identified novel antibiotics (39, 61, 91, 142, 146), antibiotic resistance genes (44, 115), Na⁺(Li⁺)/H⁺ transporters (93), and degradative enzymes (71– 73) The power of the approach is that it does not require that the genes of interest be recognizable by sequence analysis, making it the only approach to metagenomics that has the potential to identify entirely new classes of genes for new or known functions. The significant limitation is that many genes, perhaps most, will not be expressed in any particular host bacterium selected for cloning. In fact, there is an inherent contradiction in this approach—genes are cloned from exotic organisms to discover new motifs in biology, and yet these genes are required to be expressed in Escherichia coli or another domesticated bacterium in order to be detected. The diversity of the organisms whose DNA has been successfully expressed in *E. coli* is surprising (16, 33, 45, 57, 118–120), but heterologous expression remains a barrier to extracting the maximum information from functional metagenomics analyses.

Identifying active clones—screens, selections, and functional anchors. The frequency of metagenomic clones that express any given activity is low. For example, in a search for lipolytic clones derived from German soil, only 1 in 730,000 clones showed activity (73). In a library of DNA from North American soil, 29 of a total of 25,000 clones expressed hemolytic activity (118). The scarcity of active clones therefore necessitates development of efficient screens and selections for discovery of new activities or molecules. Just as bacterial genetics relies on selections to detect low-frequency events, metagenomics will be advanced by seeking selectable phenotypes to increase the collection of active clones that can be compared, analyzed, and used to build a conceptual framework for functional analysis.

Several selections have proved to be fruitful. For example, the Daniel group designed a clever selection for Na⁺(Li⁺)/H⁺ antiporters that requires complementation of an *E. coli* mutant deficient in the three Na⁺/H⁺ antiporters (*nhaA*, *nhaB*, and *chaA*) enabling growth on medium containing 7.5 mM LiCl (93). This powerful selection facilitated the discovery of two novel antiporter proteins in a library of 1,480,000 clones containing DNA isolated from soil. Another selection strategy involved complementation of an *E. coli* mutant deficient in biotin production, which led to the isolation of seven new

operons for biotin synthesis from enrichment cultures derived from samples of soil or horse excrement (52).

Selection for antibiotic resistance led to the isolation of a tetracycline resistance determinant from samples of the microbiota from the human mouth (44) and aminoglycoside resistance determinants from soil (115). The selection for aminoglycoside resistance identified nine clones, six of which encoded 6'-acetyltransferases that formed a new cluster based on sequence analysis. These genes were discovered in libraries containing a total of 4 Gb of DNA, or approximately 1 million genes, and thus their infrequent representation would have made it prohibitively laborious to discover them by a screen without a selection. This example illustrates the power of functional metagenomics—genes that are expressed in an ordinary host such as *E. coli* may be extraordinary and novel.

High-throughput screens can substitute when the functions of interest do not provide the basis for selection. For example, on certain indicator media, active clones display a characteristic and easily distinguishable appearance even when plated at high density. With the indicator dye tetrazolium chloride, Henne et al. (72) detected clones that utilize 4-hydroxybutyrate in libraries of DNA from agricultural or river valley soil. Very rare lipolytic clones in the same libraries were detected by production of clear halos on media containing rhodamine and either triolein or tributryin (73).

The discovery of new biological motifs will depend in part on functional analysis of metagenomic clones. Functional screens of metagenomic libraries have led to the assignment of functions to numerous "hypothetical proteins" in the databases. Innovation will be required to identify and overcome the barriers to heterologous gene expression and to detect rare clones efficiently in the immense libraries that are needed to represent all of the genomes in complex environments, such as soil. An emerging and powerful direction for metagenomic analysis is the use of functional anchors, which are the functional analogs of phylogenetic anchors. Functional anchors are functions that can be assessed rapidly in all of the clones in a library. When a collection of clones with a common function is assembled, they can be sequenced to find phylogenetic anchors and genomic structure in the flanking DNA. Such an analysis can provide a slice of the metagenome that cuts across clones with a different selective tool, determining the diversity of genomes that contain a particular function that can be expressed in the host carrying the library. Technological developments that promote functional expression and screening will advance this new frontier of functional genomics.

ECOLOGICAL INFERENCE FROM METAGENOMICS

The exigent questions in microbial ecology focus on how microorganisms form symbioses with eukaryotes, compete and communicate with other microorganisms, and acquire nutrients and produce energy. Thus far, metagenomics has provided insights into each of these areas, but in each instance, the challenge is to link the genomic information with the organism or ecosystem from which the DNA was isolated. Expression of a gene in a cultured host can establish gene function, but without the appropriate biological context, circumspection is required in drawing ecological inferences. Future technical

innovations are needed to extend insights from metagenomics from inference to mechanistic analyses.

Symbiosis

Many bacterial symbionts that have highly specialized and ancient relationships with their hosts do not grow readily in culture. Many of them live in specialized structures, often in pure or highly enriched culture, in host tissues, making them ideal candidates for metagenomic analysis because the bacteria can be separated readily from host tissue and other microorganisms. This type of analysis has been conducted with Cenarchaeum symbiosum, a symbiont of a marine sponge (111), a Pseudomonas-like bacterium that is a symbiont of Paederus beetles (110), Buchnera aphidicola, an obligate symbiont of aphids (1), the Actinobacterium Tropheryma whipplei, the causal agent of the rare chronic infection of the intestinal wall (13, 31), and the Proteobacterium symbiont of the deep sea tube worm Riftia pachyptila (75). These systems provide good models for metagenomic analysis of more complex communities and thus warrant further attention in this review, although the term metagenomics typically connotes the study of multispecies communities. Therefore, the following section focuses on two of these obligate symbionts and the insight into their lifestyles offered by metagenomic analysis.

Buchnera-aphid symbiosis. The first genome reconstruction of an uncultured organism was that of Buchnera aphidicola, the endosymbiont of aphids. The relationship between the bacterium and the insect is ancient, leaving each partner unable to function independently of the other, as is reflected in the genomic analysis. Moran's group isolated bacterial DNA from the insect and sequenced and reassembled the bacterial genome. The genus Buchnera contains a "reduced" genome of 564 open reading frames. Upon comparison with a reconstructed ancestral genome, 1,906 genes appear to have been lost. Most of the functions are associated with biosynthetic pathways contributed by the host, suggesting that the genome shrinkage is the result of the symbiotic lifestyle, which has become obligate because of gene loss (1, 42, 101).

The reconstruction of *B. aphidicola*'s genome provided insights into the evolution of the symbiosis between the insect and bacterium, the biochemical mutual dependence that they have developed, and the mechanisms of genome shrinkage and rearrangement. The success of genome reconstruction with a single uncultured species provided part of the impetus needed to propose sequencing and reconstructing genomes in more complex assemblages.

Proteobacterium-tube worm symbiosis. Riftia pachyptila, the deep sea tube worm, lives 2,600 m below the ocean surface, near the thermal vents that are rich in sulfide and reach temperatures near 400°C. The tube worm does not have a mouth or digestive tract, and therefore it is entirely dependent on its symbiotic bacteria, which provide the worm with food. The bacteria live in the trophosome, a specialized feeding sac inside the worm (32). The bacteria and trophosome constitute more than half of the animal's body mass. The bacteria oxidize hydrogen sulfide, thereby producing the energy required to fix carbon from CO₂, providing sugars and amino acids (predominantly as glutamate) that nourish the worm (55, 84). The worm contributes to the symbiosis by collecting hydrogen sul-

fide, oxygen, and carbon dioxide and transporting them to the bacteria on hemoglobin-like molecules (3, 46, 149–153).

The bacterium is a member of the γ -Proteobacteria, as identified by 16S rRNA gene sequence (47). The bacteria have not been grown in pure culture in laboratory media, but they provide an excellent substrate for metagenomics because they reach high population density in the trophosome and exist there as a single species. Hughes et al. (75) isolated DNA from the bacterial symbiont and constructed fosmid libraries from it that were used to understand the physiology of the bacteria. Robinson et al. (116) identified a gene with similarity to ribulose-1,5-bisphosphate carboxylase/oxygenase (RubisCO) from the same fosmid library. All of the residues associated with the active site are conserved in the protein sequence deduced from the DNA sequence, and it has highest similarity with the RubisCO from *Rhodospirillum rubrum*. The characterization of this gene lends further support to the premise that the chemoautotrophic bacterial symbiont in R. pachyptila fixes carbon for its host.

The libraries were also screened for two-component regulators with a labeled histidine kinase gene as a probe. They identified a two-component system whose components complemented an envZ and a phoR creC double mutant, respectively. The discovery of a functional envZ homologue indicates that the symbiont carries a response regulator that is typical of γ -Proteobacteria, although the signals eliciting responses from these proteins have not yet been identified.

Genomic analysis of the symbiont also led to the identification of a gene encoding flagellin, which was expressed in *E. coli* and shown to direct the synthesis of flagella that are immunologically cross-reactive with *Salmonella* flagella. The presence of genes for flagella suggested to the authors that the endosymbiont has a free-living stage in its life cycle and may infect each generation of tube worms rather than being passed maternally (99).

Competition and Communication

What can metagenomics tell us about microbial competition and communication? Competition for resources among community members selects for diverse survival mechanisms, including antagonism and mutualism among the members. Understanding these mechanisms is central to advancing the definition of principles that govern microbial community structure, function, and robustness. Historically, genetics has provided the most convincing evidence for traits contributing to microbial fitness. Classic mutant analysis has revealed genes required for microbial competition (9, 15, 20, 27, 82, 98, 100, 140, 143), antagonism (40, 51, 77, 117, 130), and mutualism (10, 54, 86, 92). Mutant analyses have provided the greatest advances in knowledge because screening mutants containing random mutations for effects on fitness has led to the identification of genes that would not have been predicted to play a role in microbial competition or mutualism.

Genes for competition and cooperation are hard to recognize based on sequence alone because the utility of their functions is entirely dependent on ecosystem context and the nature of the resources that are limiting. Therefore, genomics by itself does not provide a means to test ecological hypotheses or identify genes that confer fitness, but it can provide the basis for forming hypotheses. Ecological hypotheses are difficult to

test in microorganisms that cannot be cultured or for which there are no genetic tools; however, functional genomics coupled with chemical ecology can yield informative answers. Chemical ecology involves the identification of small molecules with biological activity and proposed ecological function. These compounds can be identified through a variety of methods, including metagenomics. The addition of these molecules to communities can provide the basis for postulating their ecological roles in the community by measuring perturbations of community function. The following sections explore the discovery of small molecules in metagenomic libraries and postulate the ecological functions of these molecules in the organisms producing them.

Role of small molecules. Small-molecule discovery by functional metagenomics has concentrated on antibiotics, which are of interest for their pharmaceutical applications as well as for their roles in ecosystem function. Traditional antibiotic screens for molecules that inhibit bacterial growth have led to the discovery of antibiotics in metagenomic libraries (Fig. 4). They have not been a rich source of novel antibiotics, likely because of the experimental limitations associated with the search. In studies that report frequencies, antibiotic-producing clones are detected at a frequency of approximately 1 producer per 10⁴ clones (23, 24, 61). This low frequency hinders discovery because space and labor are required to conduct typical antimicrobial screens.

With standard inhibition assays, a Mycobacterium-inhibiting antibiotic, terragine, was discovered from a soil metagenomic clone maintained in Streptomyces lividans (146) and acyltyrosines from a clone maintained in E. coli (22, 24). Colored antibiotics represent a disproportionate share of those discovered because they can be identified visually. For instance, a clone noticed for its brown pigment was found to produce melanin, which masked orange and red pigments, two novel antibiotics, turbomycin A and turbomycin B (61). A purplepigmented clone (23) produced violacein, previously shown to be an antibiotic made by the soil bacterium Chromobacterium violaceum. The sequence of the genes on the metagenomic clone diverged substantially from the C. violaceum violacein biosynthetic operon despite similar genetic organization (4, 23, 109), suggesting that the pathway on the metagenomic clone was derived from an organism other than C. violaceum. Osburne's group identified structurally related compounds, indirubin and indigo blue, in a soil metagenomic DNA library based on their blue color (91).

Sequence-based screening for small molecules. The first polyketide synthases, enzymes involved in synthesis of polyketides, the broad class of antibiotics that includes erythromycin, epithilone, and rifamycin, were first cloned from soil with a PCR-based approach. Seow et al. (128) designed primers that hybridize with the highly conserved region of polyketide synthase genes and amplified novel polyketide synthase homologues directly from soil. This approach was adapted for screening metagenomic libraries by Osburne's group, who screened a 5,000-member metagenomic library for conserved regions of genes encoding type I polyketide synthase. Primers directed toward a conserved region of polyketide synthase I genes that flanks the active site of the ketoacyl synthetase domain were used to screen pools of 96 clones. The screen yielded 11 new polyketide synthase homologues that contained significant se-

Long-chain N-Acyl Amino Acid Antibiotics

FIG. 4. Antibiotics discovered in metagenomic libraries.

quence similarity to polyketide synthase genes from cultured organisms. In addition, screening clones in both *E. coli* and *Streptomyces lividans* by chemical means revealed two novel compounds, fatty dienic alcohol isomers (Fig. 4).

Antibiotics as signal molecules. If antibiotics evolved as mediators of functions other than warfare (42a), such as communication, antibiotic discovery will be expedited by screening metagenomic clones for signaling compounds as well as inhib-

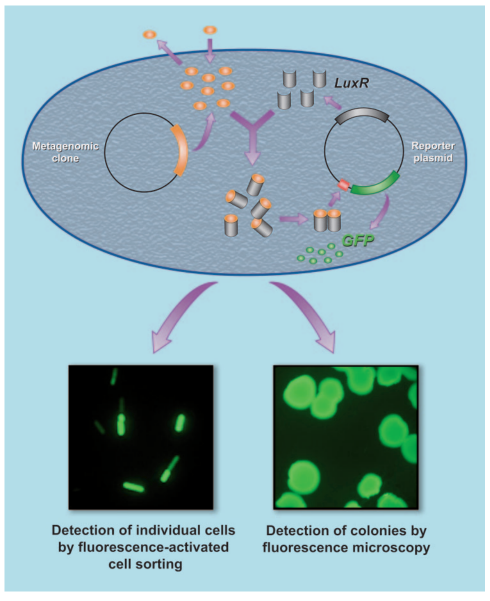


FIG. 5. Intracellular screen for quorum-sensing inducers. The biosensor, which detects molecules that induce *luxR*-regulated genes, resides inside the same cell as the metagenomic clone. An active clone fluoresces due to accumulation of green fluorescent protein and can be detected by fluorescence microscopy or captured by fluorescence-activated cell sorting.

itory compounds. The challenge is to develop assays that detect signaling by many compounds. A surprising result from the Davies group indicated that subinhibitory concentrations of many antibiotics induce quorum sensing despite no resemblance in structure to the acylated homoserine lactones that appear to be the natural inducers (65). This result presents a propitious opportunity—a single screen might capture molecules that are quorum-sensing inducers as well as antibiotics.

This opportunity was investigated by designing a highthroughput screen to identify compounds that induce the expression of genes under the control of a quorum-sensing promoter. The screen is intracellular, meaning the metagenomic DNA is in the same cell as the sensor for quorum-sensing induction (Fig. 5). The sensor is comprised of the *luxR* promoter, which is induced by acylated homoserine lactones, linked to gfp, and resides on a plasmid in an E. coli strain that did not induce quorum sensing itself (2). If an inducer of the luxR-mediated transcription of gfp is expressed from the metagenomic DNA, the cell fluoresces and can be captured by fluorescence-activated cell sorting or as a colony observed by fluorescence microscopy. Conversely, this sensor system can detect inhibitors of quorum sensing if acylated homoserine lactone is added to the medium and fluorescence-activated cell sorting is set to collect the nonfluorescent cells (Fig. 5). Metagenomic libraries from microbiota of the soil and from the midgut of the gypsy moth have been subjected to this screen, and an array of genes have been identified. Their products are under analysis, and some appear to differ from previously described quorum sensing inducers (L. Williamson, C. Guan, B. Borlee, and J. Handelsman, unpublished data).

Biogeochemical Cycles

Acid mine drainage. An exciting potential of metagenomics is to provide community-wide assessment of metabolic and biogeochemical function. Analysis of specific functions across all members of a community can generate integrated models about how organisms share the workload of maintaining the nutrient and energy budgets of the community. The models can then be tested with genetic and chemical approaches. The best example of such an analysis is the nearly complete sequencing of the metagenome of a community in acid drainage of the Richmond mine, which represents one of the most extreme environments on Earth. The microbial community forms a pink biofilm that floats on the surface of the mine water. The drainage water below the biofilm has a pH of between 0 and 1 and high levels of Fe, Zn, Cu, and As (317, 14, 4, and 2 mM, respectively). The solution around the biofilm water is 42°C and microaerophilic. There is no source of carbon or nitrogen other than the gaseous forms in the air. The community is dominated by a few bacterial genera, Leptospirillum, Sulfobacillus, and sometimes Acidomicrobium, and one archaeal species, Ferroplasma acidarmanus, and other members of its group, the *Thermoplasmatales*. The mine is rich in sulfide minerals, including pyrite (FeS₂), which is dissolved as a result of oxidation, which is catalyzed by microbial activity (5,

The simple community structure made it possible for Tyson et al. (141) to clone total DNA and sequence most of the community with high coverage. The G+C content of each clone provided a good indicator of its source because the G+C content of the genomes of the dominant taxa in the mine differ substantially (19) (Fig. 2). Sequence alignment of 16S rRNA and tRNA synthetase genes confirmed the organismal origins of the clones. Nearly complete genomes of *Leptospirillum* group II and *Ferroplasma* type II were reconstructed, and substantial sequence information for the other community members was reported.

The metagenomic sequence substantiated a number of significant hypotheses (Fig. 6). First, it appears that Leptospirillum group III contains genes with similarity to those known to be involved in nitrogen fixation, suggesting that it provides the community with fixed nitrogen. This was a surprise because the previous supposition was that a numerically dominant member of the community, such as Leptospirillum group II, would be responsible for nitrogen fixation. However, no genes for nitrogen fixation were found in the Leptospirillum group II genome, leading the authors to suggest that the group III organism is a keystone species that has a low numerical representation but provides a service that is essential to community function. Ferroplasma type I and II genomes contain no genes associated with nitrogen fixation but contain many transporters that indicate that they likely import amino acids and other nitrogenous compounds from the environment.

Energy appears to be generated from iron oxidation by both *Ferroplasma* and *Leptospirillum* spp. The genomes of both groups contain electron transport chains, but they differ significantly. The genomes of *Leptospirillum* group II and III contain putative cytochromes that typically have a high affinity for oxygen. The cytochromes may play a role in energy transduc-

tion as well as in maintaining low oxygen tension, thereby protecting the oxygen-sensitive nitrogenase complex.

All of the genomes in the acid mine drainage are rich in genes associated with removing potentially toxic elements from the cell. Proton efflux systems are likely responsible for maintaining the nearly neutral intracellular pH, and metal resistance determinants pump metals out of the cells, maintaining nontoxic levels in the interior of the cells.

The acid mine drainage community provides a model for the analysis of other communities. Determining the origin of DNA fragments and assigning functions may be more difficult for communities that are phylogenetically or physiologically more complex (59), but the approach will be useful for all communities.

Sargasso Sea. The Sargasso Sea is a complex and physically sprawling ecosystem compared with the contained acid mine drainage system. The inputs and outputs are more difficult to quantify, and the phylogeny of the community members has not been exhaustively surveyed. Venter et al. (142) embarked on the largest metagenomics project to date (affectionately dubbed megagenomics), in which they sequenced over 1 billion bp and claim to have discovered 1.2 million new genes. Intriguing inferences could be drawn because of the sheer size of the data set. They placed 794,061 genes in a conserved hypothetical protein group, which contains genes to which functions could not be confidently assigned. The next most abundant group contained 69,718 genes apparently involved in energy transduction. Among these were 782 rhodopsin-like photoreceptors, increasing the number of sequenced proteorhodopsin genes by almost 10-fold. Linkage of the rhodopsin genes to genes that provide phylogenetic affiliations, such as genes encoding subunits of RNA polymerase, indicated that the proteorhodopsins were distributed among taxa that were not previously known to contain light-harvesting functions, including the Bacteroides phylum (142).

The Sargasso Sea data set is a gold mine for further analysis. Intriguing hints about many aspects of ecosystem function abound and await further exploration. For example, an intriguing initial observation is that many of the genomes in the Sargasso Sea contain genes with similarity to those involved in phosphonate uptake or utilization of polyphosphates and pyrophosphates, which are present in this extremely phosphatelimited ecosystem. The phosphorus cycle is not well understood, and this collection of genomes provides a new route for discovery of the mechanisms of phosphorus acquisition and transformation. The understanding of nutrient cycling will be advanced by reconstruction of the genomes and the type of function-species analysis that Tyson et al. applied to the acid mine drainage community. The sequence data set from the Sargasso Sea provided the means to reassemble a number of genomes with criteria that include the depth of sequencing coverage, oligonucleotide frequencies, and similarity to previously sequenced genomes. The structures of these genomes individually and collectively will no doubt inform the development of models for nutrient cycling.

Population Genetics and Microheterogeneity

Metagenomic analysis has revealed that even apparently uniform populations contain substantial microheterogeneity.

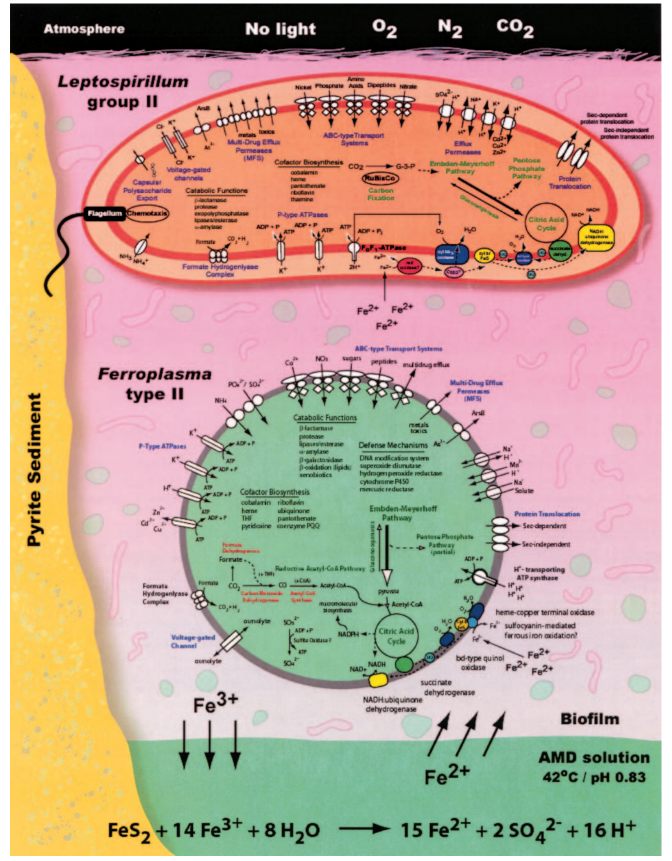


FIG. 6. Metagenomics-based model of biogeochemical cycles mediated by prokaryotes in acid mine drainage. Cell metabolic cartoons constructed from the annotation of 2,180 open reading frames identified in the *Leptospirillum* group II genome and 1,931 open reading frames in the *Ferroplasma* type II genome. The cells are shown within a biofilm that is attached to the surface of an acid mine drainage stream. Reproduced from reference 141 with permission of the publisher.

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Within the population of Cenarchaeum symbiosum associated with the marine sponge, the rRNA genes are highly conserved, showing >99.2% identity, which indicates that the population comprises a single species. In the genomic regions flanking the rRNA genes, however, the DNA sequence identity drops to 87.8% (123). Similarly, Tyson et al. (141) found that the genomes of the species or groups in acid mine drainage varied in their uniformity. They found a high frequency of single-nucleotide polymorphisms among strains of the same species. The Ferroplasma type II group appears to contain a composite genome, with segments derived from three sources. In contrast, the Leptospirillum group II genome contained very few single-nucleotide or large-scale genome polymorphisms. These studies point to the importance of conducting genomic analysis on mixtures of strains to obtain a portrait of the heterogeneity within the species. In fact, metagenomics may provide insight into genome variation of organisms that can be readily cultured. If genetic variation in the environmental population is of interest, it may be more productive to clone the genome from the natural population than analyze the genomes of individuals cultured from it.

CONCLUSIONS AND FUTURE DIRECTIONS

Metagenomics has changed the way microbiologists approach many problems, redefined the concept of a genome, and accelerated the rate of gene discovery. The potential for application of metagenomics to biotechnology seems endless. Functional screens have identified new enzymes (39, 52, 53, 67, 68, 72, 73, 88, 93, 104, 118, 122, 124, 144) and antibiotics (22, 23, 39, 61, 69, 91, 110, 118, 146) and other reagents in libraries from diverse environments. A number of barriers have limited the discovery of new genes that provide insight into microbial community structure and function or that can be used to solve medical, agricultural, or industrial problems.

Realizing the potential for discovery from metagenomics is dependent on the advancement of methods that are central to library construction and analysis. For sequence-based approaches, the speed and cost of nucleotide sequencing will be a barrier of rapidly diminishing significance as sequencing technology continues to improve. Sequence-based assignment of function will also benefit from advances in detection of homology, which will increasingly rely on the tertiary structures of predicted proteins rather than simply on primary sequence. Advances that will facilitate the management and analysis of large libraries include bioinformatics tools to analyze vast sequence databases and reassemble multiple genomes rapidly and affordable gene chips for library profiling (127) or that readily distinguish clones that are expressing genes from those clones that are silent. Functional analysis will require more innovation in method development. Most important among these are strategies to improve heterologous gene expression and approaches for efficient screening of large libraries.

Microbiology has long relied on diverse methods for analysis, and metagenomics can provide the tools to balance the abundance of knowledge attained from culturing with an understanding of the uncultured majority of microbial life. Myriad environments on Earth have not been studied with culture-independent methods other than PCR-based 16S rRNA gene analysis, and they invite further analysis. Metagenomics

may further our understanding of many of the exotic and familiar habitats that are attracting the attention of microbial ecologists, including deep sea thermal vents; acidic hot springs; permafrost, temperate, desert, and cold soils; Antarctic frozen lakes; and eukaryotic host organs—the human mouth and gut, termite and caterpillar guts, plant rhizospheres and phyllospheres, and fungi in lichen symbioses. With improved methods for analysis, funding stimulated by recent triumphs in the field, and attraction of diverse scientists to identify new problems and solve old ones, metagenomics will expand and continue to enrich our understanding of microorganisms.

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AUTHOR'S CORRECTION

Metagenomics: Application of Genomics to Uncultured Microorganisms

Jo Handelsman

Department of Plant Pathology, University of Wisconsin-Madison, Madison, Wisconsin

Volume 68, no. 4, p. 669–685, 2004. Page 682: The Acknowledgments section was inadvertently omitted and should appear as shown below.

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