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Metagenomic tools in microbial ecology research

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Ability to directly sequence DNA from the environment permanently changed microbial ecology. Here, we review the new insights to microbial life gleaned from the applications of metagenomics, as well as the extensive set of analytical tools that facilitate exploration of diversity and function of complex microbial communities. While metagenomics is shaping our understanding of microbial functions in ecosystems via gene-centric and genome-centric methods, annotating functions, metagenome assembly and binning in heterogeneous samples remains challenging. Development of new analysis and sequencing platforms generating high-throughput long-read sequences and functional screening opportunities will aid in harnessing metagenomes to increase our understanding of microbial taxonomy, function, ecology, and evolution in the environment.

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Introduction

Over 20 years ago the term ‘metagenomics’ [1] emerged, announcing a new frontier in exploring the cross-section of biology and chemistry. From its first application in soil, metagenomics has become a driving force for discoveries in microbial ecology and biotechnology and a key method in exploring the microbial universe. As sequencing technologies became cheaper, faster, easier to use and higher-throughput, our ability to survey microbial diversity and functional potential in any ecosystem increased dramatically. Successful applications of metagenomics are closely tied to the availability and capability of the computational methods. In this review, we highlight the new knowledge gleaned from applications of metagenomics in Earth’s different ecosystems as well as the analytical tools that enable such discoveries.

A journey from genes to ecosystem functions

Metagenomics, direct sequencing and analysis of DNA from microbial assemblages, has rapidly become a routinely employed method to characterize the functional potential of microbial communities. In its most straightforward application, DNA is extracted, prepared into libraries, and sequenced either on short-read (Illumina, Roche 454, Ion Torrent) or long-read (PacBio, Oxford Nanopore) platforms. All metagenomics analyses start with quality control of the sequence reads, which aims to minimize sequence bias and artifacts by removing adapter sequences, low quality bases calls, and contaminant sequences that are not from the source environment (Table 1). Earlier applications of metagenomics relied largely on a gene-centric approach to quantify the relative abundance of genes of interest and their function within a metagenome [2], which requires gene detection [3] and annotation of short reads. These efforts immediately increased the number of gene clusters in databases [2,4] and spurred greater interest in the applications of metagenomics. Various tools enable this analysis (Tables 1 and 2); however, almost 50% of genes in environmental microbiomes lack annotated functions. This parallels the fact that one-third of protein-coding genes in microbial isolate genomes are unannotated [5]. Hence, our ability to identify functional genes is closely tied to the completeness of gene databases and improvements to our collective knowledge of gene functions [5].

Gene-centric metagenome analysis can be performed via stand-alone tools or web-based applications. Read based annotations require aligning predicted gene sequences to known genes to infer functional gene abundances and

Table 1**Bioinformatic programs used in sequence read quality control, assembly, binning and metagenome assembled genome (MAG) refinement**

Tools	Features	Website
<i>Sequence Read Quality Control</i>		
FastQC	Provides several graphic QC statistics information	[link]
MultiQC	Aggregates results from multiple samples into one single report	[link]
FastQ Screen	Screens sequences against a set of reference database	[link]
BBDuk	Decontaminates sequences using Kmer-based operations	[link]
Khmer	Trims and normalizes sequences for Kmer-based analysis	[link]
<i>Read Assembly</i>		
CLC Assembler	A De-bruijn graph-based assembly tool that integrates in commercial CLC workbench developed by QIAGEN	[link]
Meta-IDBA	Attempts to cover for both high and low abundant genomes by iterating with multiple k-mer size	[link]
MetaVelvet-SL	An extension of Velvet assembler hat integrating a Support Vector Machine (SVM) – is trained by a similar population of samples – to increase the performance	[link]
MEGAHIT	Uses increasing k-mer strategy with succinct de Bruijn data structure to reduce computational cost	[link]
Metaspades	A mode of the assembly software SPAdes for metagenomic assembly, using a heuristic method to distinguish interspecies repeats. Single cell mode is recommended for viromes.	[link]
<i>Assembly Quality Check</i>		
Quast	Evaluates genome/metagenome assemblies by computing various metrics such as contig length, N50, GC content	[link]
dnAQET	A Java package designed to evaluate scaffolds/contigs against a reference genome	[link]
GenomeQC	A toolkit that integrates multiple metrics to characterize both assembly and gene annotation quality across multiple data	[link]
<i>Binning and Metagenome Assembled Genome Refinement</i>		
MetaBAT2	Uses a k-medoid clustering method to bin contigs by calculating pairwise distance based on tetranucleotide frequency	[link]
Maxbin2	Employs an Expectation-Maximization (EM) algorithm to cluster contigs after co-assembly of multiple metagenomic datasets	[link]
CONCOCT	An unsupervised binning approach for metagenomic contigs by using nucleotide composition - kmer frequencies - and coverage data	[link]
GroopM	An automated binning tool that uses differential coverage (spatio-temporal model) to obtain high quality bins genomes from multi-sample metagenomes.	[link]
DASTool	Integrates results from various binning algorithms to calculate an optimized, non-redundant set of bins	[link]
CheckM	A common tool used to evaluate the quality of recovered MAGs, like completeness and contamination, based on the frequency of single-copy marker genes	[link]
<i>Gene prediction</i>		
FragGeneScan	Predicts genes from short reads incorporating a sequence error model and codon usage statistics.	[link]
Glimmer-MG	Uses interpolated Markov models (IMMs) to identify the coding regions and distinguish them from noncoding DNA.	[link]
Prodigal	Provides fast gene prediction from prokaryotic genomes, includes normal mode (reference-based) and anonymous mode (metagenomes).	[link]
MetaGeneMark	Predicts protein coding genes in metagenomic data using ab initio approaches.	[link]
Prokka	Identifies genes against series external databases that can annotate bacterial, archaeal and viral genomes.	[link]

distribution. Developing a stand-alone analysis capability requires local computational resources and proficiency in bioinformatics, but in return, users can rapidly assess the sensitivity of different parameters and analytical approaches to improve annotations. Web servers (Table 3) provide a user-friendly analysis platform that is accessible to researchers from all experience levels, but they are limited to small data sizes, provide results from a select list of analysis tools and analysis completion can take weeks to months (depending on the server load). Despite these limitations, services such as JGI IMG/M [6], MG-RAST [7], CyVerse [8] and KBase [9] serve a wide research community and support development and

deployment of new analytical tools. Metagenomics now is a well-established technology, where a growing number of datasets [10] and analysis tools are accessible to many researchers [11].

Information gleaned from gene-centric metagenomics, at times in combination with RNA sequencing (metatranscriptomics) and protein identification (metaproteomics), provides greater understanding of microbial processes governing biogeochemical cycles in ecosystems. Metagenomic analysis of taxonomic and functional diversity in prairie soil microbiomes, for example, showed how long-term agricultural practices can result in the loss of

Table 2**Commonly used software for discovery of phylogeny and functional potential based on search against database**

Database	Tools	Key features	Website
Nucleotide	Kraken2	Exact k-mer search in memory	[link]
	Bracken	Computes relative abundance of species using Bayesian estimation	[link]
	CLARK	Supervised sequence classification using discriminative k-mers	[link]
	k-SLAM	K-mer search with additional validation using pseudo-assembly	[link]
	Centrifuge	Fast and memory-efficient tools for taxonomic profiling using BWT	[link]
Protein	DIAMOND	Protein homology search using spaced seeds with a reduced amino acid alphabet	[link]
	Kaiju	Fast for large-scale profiling in protein database	[link]

Table 3**Metagenomic analysis platforms enabling gene- and genome-centric analysis**

Platforms	Key features	Website
<i>Web-based</i>		
EBI metagenomics	A comprehensive platform for the assembly, analysis and archive microbiome data	[link]
MG-RAST	An open source web application for gene-centric analysis that offer automated quality control, annotation, comparative analysis and archiving services.	[link]
KBase	A suite of microbiome analysis apps for gene- and genome-centric analysis with a graphic interface. User friendly import, export, and data edits and metabolic modelling capability	[link]
IMG/M	A platform for comparative analysis and functional annotation for public available genomes, metagenomes and metatranscriptomes.	[link]
<i>Local installation</i>		
MetAMOS	A modular framework for metagenomic assembly, taxonomic and functional annotations, and integrated HTML report.	[link]
MOCAT2	A toolkit to generate assembly, gene predictions, gene catalogs, gene catalog annotations, functional or taxonomic profiles for metagenomics.	[link]
Anvi'o	Provides integrated analysis strategies for genomics, metagenomics, metatranscriptomics, pangenomics, metapangenomics, phylogenomics, and microbial population genetics in an integrated and has extensive interactive visualization capabilities.	[link]
Metawrap	Metagenomic wrapper suite that accomplishes the core tasks of metagenomic analysis from start to finish: read quality control, assembly, visualization, taxonomic profiling, extracting and refining draft genomes (binning), and functional annotation.	[link]
METABOLIC	This software enables the prediction of metabolic and biogeochemical functional trait profiles to any given genome datasets. These genome datasets can either be MAGs, single-cell amplified genomes or pure culture genomes.	[link]
MetaSanity	Provides a unified workflow for genome assessment and functional annotation that combines all outputs into a single queryable database	[link]
DRAM/ DRAM-v	A tool for annotating MAGs and VirSorter identified viral contigs. DRAM annotates MAGs and viral contigs using KEGG (if provided by the user), UniRef90, PFAM, dbCAN, RefSeq viral, VOGDB and the MEROPS peptidase database as well as custom user databases.	[link]

keystone species leading to loss of functional diversity [12]. Combined use of metagenomics, metatranscriptomics and metaproteomics allows elucidation of microbial functions regulating greenhouse gas emissions [13] and their distribution [14] in climate sensitive arctic tundra soils. Furthermore, biochemical and environmental factors impacting microbial functions in sandy sediments were assessed via metagenomics identifying the importance of temporal processes resulting in frequent shifts between H_2 -fermentation and H_2 -respiration processes [15*]. Soil metagenomes can be further analyzed to find new biologically and environmentally important enzymes. In combination with empirical testing, this approach can be used to extend the categorization of known enzymes in databases [16].

Beyond the highly complex and diverse soil and sediment microbiomes, gene-centric analyses of rarely accessed natural and engineered environments expand our knowledge of fundamental microbial processes. Metagenomics of dust microbiomes showed that genes encoding proteins involved in repairing UV-induced DNA damage along with chemotaxis, germination, and heat-shock proteins were ubiquitous across different sampling locations in North Africa [17]. Patterns in zoonotic protist diversity in raw sewage were studied to understand their distribution in urban environments showing that functionally similar but phylogenetically diverse protist community were inhabiting New York City sewers [18]. Microbes inside or on the surface of plant tissues (roots, stems, and leaves) can impact plant productivity and health [19]. Plant-

associated microorganisms shown to contain extensive collection of carbohydrate metabolism functions and fewer motility genes, suggesting endophytic microorganisms have access to varied and widely-dispersed carbon substrates [20**]. As with other microbe-host systems, host DNA contamination can reduce the ability to sequence microbial reads in plant tissues. However, this can be leveraged to quantify microbial concentrations [21,22]. Interrogating Earth's ecosystems with metagenomic analyses is an arduous task, but such analyses may illuminate the full breadth of microbial function and ecology.

Back to the cell: genome recovery from metagenomes

The major assumption of gene-centric metagenomics is that the genes exist in a well homogenized and cellfree environment where all potential reactions can interactively occur [23]. This assumption, while fundamentally flawed, was a result of our inability to solve short read sequence puzzles into genomes. In earlier attempts, short reads could be organized to infer genome content in low diversity environments like acid mine drainage biofilms [24], but, until recently, characterizing genomes in complex communities, like soils, remained a challenge. Genome recovery from metagenomes in complex communities became possible as sequence read depth per sample increased. Sequencing deep enough to have high coverage, development of methods to reconstruct the long consensus region of DNA (contigs) from a pool of short sequence reads, and coverage-based binning of assembled contigs into population genomes gave rise to genome-centric metagenomics and metagenome assembled genomes (MAGs) [25]. Short-read assembly has unique challenges, notably due to varying abundances of bacteria and archaea within a community, high diversity, and strain-level variance. New generations of assemblers were designed to account for and leverage these distinctive aspects of the data (Table 1). Currently, there is no universally accepted assembler, thus, assembly quality is often evaluated by comparing different methods through summary statistics from single-genome assembly methods like size, contig N50 (i.e. the sequence length of the shortest contig at 50% of the total genome length), and maximum contig length. Contigs derived from assembly are still highly fragmented and redundant, and they cannot be directly grouped into genomes. Binning algorithms use a variety of genome properties such as DNA composition, GC content, tetranucleotide frequency in combination with depth of sequencing coverage, and abundance to group contigs into MAGs.

The direct output of genome binning often contains false-assignment contigs. Thus, it is common to refine and evaluate MAGs after the binning. Completeness and contamination [26] are two common metrics that are used to assess MAG quality. Completeness seeks to identify

sets of single-copy marker genes. Similarly, contamination reports on misbinings based on multiple detection of single-copy marker gene sets. Both metrics are prone to certain errors, such as insensitivity to strain heterogeneity, transposases, and RNA operons. As studies reporting MAGs are increasing, a set of standards, called the minimum information about a single MAG (MIMAG) [27], has been proposed to standardize reporting, emphasizing manual curation and review. Such standards are important in assuring that published MAGs are of high quality.

Genome-resolved metagenomics has transformed our ability to study uncultured microbes and has led to discoveries in taxonomy, microbial ecology, biogeochemistry, and evolutionary biology. Incorporation of MAGs into the tree of life has increased the number of known microbial phyla, dramatically altering our understanding of microbial phylogeny. For instance, MAG analyses contributed to the discovery of the Candidate Phyla Radiation, which includes over 70 phyla and two superphyla (*Parcubacteria* and *Microgenomates*) [28–30]. Furthermore, such analyses have contributed to the discovery of phylum '*Candidatus* Kryptonia' [31], exclusive to high-temperature pH-neutral geothermal springs. This lineage represented a taxonomic 'blind spot' because of mismatches in the primers commonly used for ribosomal gene surveys. MAGs showed a heterotrophic lifestyle and strong need for symbiosis with other microbes. More recently 52 515 MAGs were generated from over 10 000 metagenomes collected from various habitats covering all of Earth's terrestrial and aquatic environments [32]. This massive effort expanded the known phylogenetic diversity of bacteria and archaea by 44% by generating 12 556 novel candidate species-level operational taxonomic units spanning 135 phyla [32]. For archaea, the discovery of the Asgard superphylum, which also includes the *Lokiarchaeota*, was a major achievement in enhancing our understanding of archaeal taxonomy [33**]. The cellular structure of these archaea contains many eukaryotic features and provides support for the emergence of eukaryotes from within the archaeal domain of life. In addition, new CRISPR–Cas systems were identified through analysis of MAGs where Cas9, previously found only in bacterial genomes, were also detected in the archaeal domain of life [34], providing new opportunities for testing and applications in biological and clinical research.

The analysis of MAGs has also revealed new insights into microbial metabolic diversity and niche differentiation. For example, *Planctomycetes* MAGs that are abundant in water systems were discovered to be able to perform nitrogen fixation in both the Pacific and the Atlantic Oceans [35*]. This showed the importance of heterotrophic bacteria in the fixation of nitrogen in the surface ocean. A new addition to the nitrogen cycle was the discovery of a complete nitrifying organism from an

engineered system, a process referred to as comammox [36]. Our knowledge of methane metabolisms was advanced via genome-resolved metagenomics where the detection of the key enzyme for methanogenesis (Methyl coenzyme M reductase) in newly discovered *Bathyarchaeota* and *Verstraetearchaeota* MAGs overturned the long-held paradigm that this functional capacity was restricted to *Euryarchaeota* [37,38]. Furthermore, some lineages of *Bathyarchaeota* are suggested to perform homoacetogenesis, the ability to solely use CO₂ and H₂ to generate acetate [39], a metabolic process that was thought to be restricted to the bacterial domain. Genome-resolved metagenomics can unravel complex community potential and interactions involved in organic matter decomposition [40]. Large scale analysis of 1529 MAGs from a permafrost thaw gradient showed previously undescribed fungal pathways for xylose degradation in bacteria. Further pairing of specific microbial populations and biogeochemistry revealed key populations that drive the mineralization of organic matter from plant-derived organic material to simple the greenhouse gases [40]. Explicitly linking microbial function to taxonomy is a major benefit of genome-resolved metagenomics, which will continue to pave the way for new discoveries in microbial ecology. Machine learning and artificial intelligence methods may help to unravel hidden patterns and metabolic capabilities of complex microbial communities and reveal ecological implications.

Viral metagenomics: the new kid on the block

The advent of meta-omic approaches has enabled the study of uncultivated viruses and entirely reshaped our understanding of viruses as major players in many of Earth's biogeochemical cycles. For example, viruses can affect microbial metabolism via lysing their hosts, which stops the host's metabolism while releasing nutrients that may drive other metabolisms (i.e. viral shunt), and during infection, viruses redirect and potentially augment (via auxiliary metabolic genes) host metabolism, changing the function of the host and its metabolic outputs [41]. Viruses can be mined from metagenomes (DNA viruses) and metatranscriptomes (RNA viruses) along with microbes. This allows for characterization of proviruses, viral episomal elements (outside of the genome, but within the host), and virions, as well as revealing viral expression levels and virus-host dynamics [42]. Shotgun metagenomic approaches can characterize viruses in the context of a microbial community, but to obtain rarer viral genomes, a targeted metagenomic approach is needed. Targeted approaches include cell-sorting, where viruses identified within the cells or close-proximity can be sequenced [43], and viral metagenomes (viromes), where viruses (and other entities of virus size) are physically separated from larger organisms via a filter before their nucleic acid is extracted and sequenced [41].

The rate of viral discovery from omic approaches is unparalleled and is complemented by the rise of virus-specific bioinformatics and contemporary technologies. New bioinformatics tools have allowed characterization of viral ecosystem impacts [44,45**], detection of obscure viruses [46**,47**], virus taxonomy for uncultured viruses [48], and global comparisons of viruses [45**,46**,49**]. The development of long-read technology allows detection of whole virions [50,51] and when combined with short-reads, allows increased detection and characterization of viral genomes [52,53]. Powerful tools, such as stable isotope probing and nano scale secondary ion mass spectroscopy, are being leveraged to describe virus activity and quantify virus-host interactions [54,55].

There are limitations in characterizing viruses via meta-omic approaches that need attention. Viruses don't have a universal marker gene and most detected viral genes have unknown function, some of which are host-derived. These limitations create challenges for prediction of genome completeness, a complete taxonomic framework, and whether the virus is virulent or temperate [44], all of which impede a complete understanding of viral impacts in an ecosystem. Over the next decade, advancements in methodology and bioinformatics along with increased utilization of tools will push viral metagenomics to move beyond 'stamp collecting' of viral genomes to the quantification of viruses in an ecosystem and evaluation of their ecology particularly as it changes in space and time.

Conclusions and outlook

While microbial ecologists dig deep into the new information metagenomes provide, the metagenomic analyses of microbiomes will continue to evolve by technological and accessibility improvements in DNA and RNA sequencing. Long-read (>10 kb) sequencing technologies hold a great potential to improve genome assemblies and assignment of taxonomy and function. However, these advantages are constrained by a high error rate (10–15%) [56,57]. Because the error rate may be greater than the genetic difference between organisms, especially for low-abundance organisms, the use of long-read data for metagenomics is currently in its developmental stages. Ever growing use of new platforms (e.g. Hi-C [58*] and Tn-seq [59*]) with metagenomics will add to current data generation efforts and create new bottlenecks for data storage and standardization. As long-read sequencing becomes cheaper and more accurate, currently used elaborate methods for MAG discovery will be challenged. Future metagenomics will be closely tied to data analysis solutions that can facilitate, high-speed search and memory-efficient assembly methods that are compatible with terra- to petabytes of data. However, a key companion to these analytic methods is expanding high-quality annotation databases that are pivotal to understand the mechanisms underlying microbiome functions. Moreover, improvements in sample preparation and sequencing

for low DNA and RNA inputs will allow us to sample on smaller scales and enable accessing genomic information from larger spatial scales [60]. Further attempts to adapt current sequencing technologies for absolute quantification [61] of all molecules within a microbial cell can aid in the scaling of core and dynamic functionality complex microbiomes to larger biogeochemical and ecosystem level interactions that drive the Earth's material cycles. Overcoming methodological challenges will continue to increase our understanding of microbial taxonomy, function, ecology, and evolution.

Conflict of interest statement

Nothing declared.

CRedit authorship contribution statement

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