Shotgun metagenomics, from sampling to sequencing and analysis

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Diverse microbial communities of bacteria, archaea, viruses and single-celled eukaryotes have crucial roles in the environment and human health. However, microbes are frequently difficult to culture in the laboratory, which can confound cataloging members and understanding how communities function. Cheap, high-throughput sequencing technologies and a suite of computational pipelines have been combined into shotgun metagenomics methods that have transformed microbiology. Still, computational approaches to overcome challenges that affect both assembly-based and mapping-based metagenomic profiling, particularly of high-complexity samples, or environments containing organisms with limited similarity to sequenced genomes, are needed. Understanding the functions and characterizing specific strains of these communities offer biotechnological promise in therapeutic discovery, or innovative ways to synthesize products using microbial factories, but can also pinpoint the contributions of microorganisms to planetary, animal and human health.

Introduction

High throughput sequencing approaches enable genomic analyses of ideally all microbes in a sample, not just those that are more amenable to cultivation. One such method, shotgun metagenomics, is the untargeted (“shotgun”) sequencing of all (“meta”) of the microbial genomes (“genomics”) present in a sample. Shotgun sequencing can be used to profile taxonomic composition and functional potential of microbial communities, and to recover whole genome sequences. Approaches such as high-throughput 16S rRNA gene sequencing 1, which profile selected organisms or single marker genes are sometimes mistakenly referred to as metagenomics but are not metagenomic methods, because they do not target the entire genomic content of a sample.

In the past 15 years since it was first used, metagenomics has enabled large-scale investigations of complex microbiomes2–7. Discoveries enabled by this technology include the identification of previously unknown environmental bacterial phyla with endosymbiotic behavior 8, and species that can carry out complete nitrification of ammonia 9,10. Other striking findings include the widespread presence of antibiotic genes in commensal gut bacteria 11, tracking of human outbreak pathogens 4, the strong association of both the viral 12 and bacterial 13 fraction of the microbiome with inflammatory bowel diseases, and the ability to monitor strain-level changes in the gut microbiota after perturbations such as those induced by faecal microbiome transplantation 14.

In this Review we discuss best-practice for shotgun metagenomics studies, including identifying and tackling limitations, and provide an outlook for metagenomics in the future.
Figure 1. Summary of a metagenomics workflow. **Step 1: Study design and experimental protocol.** The importance of this step is often underestimated in metagenomics. **Step 2: Computational pre-processing.** Computational quality control steps minimize fundamental sequence biases or artefacts e.g. removal of sequencing adaptors, quality trimming, removal of sequencing duplicates (using e.g. fastqc, trimmomatic, and Picard tools). Foreign or non-target DNA sequences are also filtered and samples are sub-sampled to normalize read numbers, if the diversity of taxa or functions is compared. **Step 3: Sequence analysis.** This should comprise a combination of ‘read-based’ and ‘assembly-based’ approaches depending on the experimental objectives. Both approaches have advantages and limitations (See Table 4 for a detailed discussion). **Step 4: Post-processing.** Various multivariate statistical techniques can be used to interpret the data. **Step 5: Validation.** Conclusions from high dimensional biological data are susceptible to study driven biases so follow-up analyses are vital.

**Shotgun metagenomics study design**

A typical shotgun metagenomics study comprises five steps following the initial study design; (i) the collection, processing, and sequencing of the samples, (ii) the preprocessing of the sequencing reads, (iii) the sequence analysis to profile taxonomic, functional, and genomic features of the microbiome, (iv) the postprocessing statistical and biological analysis, and (v) the validation (Figure 1). Numerous experimental and computational approaches are available to carry out each step, which means that researchers are faced with a daunting choice. And, despite its apparent simplicity, shotgun metagenomics has limitations, owing to potential experimental biases and the complexity of computational analysis and their interpretation. We assess the choices that need to be made at each step and how to overcome common problems.
The steps involved in the design of hypothesis-based studies are outlined in Supplementary Figure 1 with specific recommendations summarized in Supplementary Box 1. Individual samples from the same environment can be variable in microbial content, which makes it challenging to detect statistically significant, and biologically meaningful, differences among small sets of samples. It is therefore important to establish that studies are sufficiently powered to detect differences, especially if the effect size is small. One useful strategy may be to generate pilot data to inform power calculations. Alternatively, a two-tier approach in which shotgun metagenomics is carried out on a subset of samples that have been prescreened with less expensive microbial surveys such as 16S rRNA gene sequencing, may be adopted.

Controls are also important but it can be difficult to obtain representative samples from a suitable control group, particularly when studying environments such as humans, in which the resident microbial communities are influenced, to a different extent, by factors such as host genotype, age, diet and environmental surroundings. Where feasible, we recommend longitudinal studies that incorporate samples from the same habitat over time rather than simple cross-sectional studies that compare “snapshots” of two sample sets. Importantly, longitudinal studies do not rely on results from a single sample that might be a non-representative outlier. Exclusion of samples that may be confounded by an unwanted variable is also prudent. For example, in studies of human subjects, exclusion criteria might include exposure to drugs that are known to impact the microbiome, e.g. antibiotics. If this is not feasible, then potential confounders should be factored into comparative analyses (see Supplementary Box 1).

If samples originate in animal models, particularly those involving co-housed rodents, the roles of animal age and housing environment, and the sex of the person handling the animals, may have on microbial community profiles should be taken into account. It is usually possible to mitigate against potential confounders in the study design by housing animals individually to prevent the spread of microbes between cage mates (although this may introduce behavioural changes, potentially resulting in different biases), mixing animals derived from different experimental cohorts together within the same cage, or repeating experiments with mouse lines obtained from different vendors or with different genetic backgrounds.

Finally, regardless of the type of sample being studied, it is crucial to collect detailed and accurate metadata. MIMARKS and MiXs standards were set out to provide guidance for required metadata, but metagenomics is now applied on such disparate kinds of environments that it is difficult to choose parameters that are suitable and feasible to obtain for every sample type. We recommend associating as much descriptive and detailed metadata as possible with each sample, in order to make it more likely that comparisons between study cohorts or sample types can be correlated with a particular environmental variable.

Sample collection and DNA extraction

Sample collection and preservation protocols can affect both quality and accuracy of metagenomics data. Importantly, the effect size of these steps, in some circumstances, can be greater than the effect size of the biological variables of interest. Indeed variations in sample processing protocols can also be important confounders in meta-analyses of datasets from different studies. Collection and storage methods that have been validated for one type of sample type cannot be assumed to be optimal.
for different sample types. As such, careful preliminary work to optimize processing conditions for sample types is often necessary (Supplementary Figure 1).

<table>
<thead>
<tr>
<th>Enrichment technique</th>
<th>Advantages</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whole genome amplification</td>
<td>Highly sensitive - can generate sufficient DNA for sequencing from even tiny amounts of starting material. Cost effective - can be applied directly to extracted environmental DNA, no need to isolate cells. Non-specific and untargeted - can amplify DNA from the whole range of species present within a given sample.</td>
<td>Amplification step can introduce significant biases, which skew resulting metagenomics profiles. Chimeric molecules can be formed during amplification, which can confound the assembly step. Non-specific – unlikely to improve proportional abundance of DNA from a species of interest.</td>
</tr>
<tr>
<td>Single-cell genomics</td>
<td>Can generate genomes from uncultured organisms. Can be combined with targeting approaches such as fluorescence in situ hybridization to select specific taxa, including those that might be rare members of the microbial community. Places genomic data within its correct phylogenetic context. Reference genomes can aid metagenomics assemblies.</td>
<td>Can be expensive to isolate single cells, requires specialist equipment. Requires whole genome amplification step – see limitations above. Biases introduced during genome amplification mean that it is usually only possible to recover partial genomes. Prone to contamination.</td>
</tr>
<tr>
<td>Flow-sorting</td>
<td>High throughput means to sort cells of interest. Targeted approach - can select specific taxa, including those that might be rare members of the microbial community.</td>
<td>Expensive equipment, requiring specialist operators. Requires intact cells. Any cells in the sample that are attached to surfaces or fixed in structures e.g. biofilms may not be recovered. Flow rates and sort volumes limit the number of cells that can be collected.</td>
</tr>
<tr>
<td>In situ enrichment</td>
<td>Simplifies microbial community structure - can make it easier to assemble genomes from metagenomics data. Presence of particular taxa within enriched samples can give clues as to their functional roles within the microbial community.</td>
<td>Requires that cells of interest can be maintained stably in a microcosm over the entire enrichment period. Simplifies microbial community structure - biases results in favour of organisms that were able to thrive within the microcosm.</td>
</tr>
<tr>
<td>Culture/microculture</td>
<td>Cultured isolates can be extensively tested for phenotypic features. Reference genomes can aid metagenomics assemblies. Functional data can improve metagenomics annotations. Places genomic data within its correct phylogenetic context.</td>
<td>Low throughput, can be highly labor intensive. Extremely biased - many microbes are inherently difficult to culture in the laboratory. Unlikely to recover rarer members of a microbial community, as cultured isolate collections will be dominated by the most abundant organisms.</td>
</tr>
<tr>
<td>Sequence capture technologies</td>
<td>Oligonucleotide probes can be used to identify species of interest as recently demonstrated for culture-independent viral diagnostics By focusing only on species of interest, higher sensitivity can be achieved particularly when large amounts of host contamination are present.</td>
<td>Capture kits can be expensive. Like PCR, capture fails when target organisms vary compared to the reference sequences used to design the probes Genome coverage of targeted organisms can be uneven, affecting assemblies.</td>
</tr>
<tr>
<td>Immunomagnetic separation</td>
<td>Targeted approach - can enrich specific taxa, including those that might be comparatively rare members of the microbial community. Far less expensive than many other targeted enrichment techniques such as single cell genomics or flow sorting. Less technically challenging and time consuming than other targeted enrichment techniques.</td>
<td>Requires intact cells. Requires a specific antibody for the target cells of interest. If target cell numbers are low, whole genome amplification may be needed following cell separation – see limitations above.</td>
</tr>
<tr>
<td>Background (e.g. human / eukaryotic) depletion techniques</td>
<td>Particularly useful for samples where microbial cell numbers are much lower than eukaryotic cells (e.g. biopsies) Improves sensitivity - enhanced detection of microbial genomic data. Lower sequence depth required to obtain good coverage of microbial genomes, reduced sequencing costs. Relatively inexpensive, not technically challenging.</td>
<td>Concomitant loss of bacterial DNA of interest can occur during processing steps, can bias subsequent microbiome profiling. May introduce contamination.</td>
</tr>
</tbody>
</table>

Table 1: Summary of the advantages and limitations of methods to enrich for microbial cells/DNA before sequencing.

Key objectives are to collect sufficient microbial biomass for sequencing, and to minimize contamination of samples. Enrichment methods can be used for those environments in which microbes are scarce (see Table 1). However, enrichment procedures can introduce bias into sequencing data. Since several studies have shown that factors such as length of time between sample collection and freezing or the number of times samples go through freeze-thaw cycles can affect the microbial community profiles that are detected, both collection and storage protocols/conditions should be recorded (Supplementary Box 1).

The choice of DNA extraction method can affect the composition of downstream sequence data. The extraction method must be able to lyse diverse microbial taxa, otherwise sequencing results may be dominated by DNA derived from easy-to-lyse microbes. DNA extraction methods that include mechanical
lysis (or bead-beating) are often considered superior to those that rely on chemical lysis. However, bead-beating based approaches do vary in their efficiency. Vigorous extraction techniques such as bead-beating can result in shortened DNA fragments, which can contribute to DNA loss during library preparation methods that use fragment size selection techniques.

Contamination can be during sample processing stages. Kit/laboratory reagents may contain variable amounts of microbial contaminants. Metagenomics datasets from low biomass samples (e.g. skin swabs) are particularly vulnerable to this problem, because there is less “real” signal to compete with low-levels of contamination. We advise those working with low biomass samples to use ultraclean reagents, and to incorporate “blank” sequencing controls, in which reagents are sequenced without adding sample template. Other types of contamination are cross-over from previous sequencing runs, presence of PhiX control DNA that is typically used as part of Illumina-based sequencing protocols, and human or host DNA.

**Library preparation and sequencing**

Choosing a library preparation and sequencing method hinges on availability of materials and services, cost, ease of automation, and DNA sample quantification. The Illumina platform has become dominant as a choice for shotgun metagenomics due to its wide availability, very high outputs (up to 1.5 Tb per run) and high accuracy (with a typical error rate of between 0.1-1%), although the competing Ion Torrent S5/S5 XL instrument is an alternative choice. Recently, long read sequencing technologies such as the Oxford Nanopore MiniON and Pacific Biosciences Sequel have scaled up output and can reliably generate up to 10 gigabases per run and may therefore soon start to see adoption for metagenomics studies.

Given the very high outputs achievable on a single instrument run, multiple metagenomic samples are usually sequenced on the same sequencing run, by multiplexing up to 96 or 384 samples typically using dual indexing barcode sets available for all library preparation protocols. The Illumina platforms are known to suffer from issues of carry-over (between runs) and carry-between (within runs). Recently, concern has been raised that newer Illumina instruments using isothermal cluster generation (ExAmp) suffer from high rates of ‘index hopping’ where incorrect barcode identifiers are incorporated into growing clusters, although the extent of this problem on typical metagenomics projects has not been evaluated and approaches to mitigate it have been suggested. To help evaluate the extent of such issues, randomly chosen control wells containing known spiked-in organisms as positive controls, and template negative controls should be used to assess the impact of these issues. Such controls are particularly critical for diagnostic metagenomics projects where small numbers of pathogen reads may be a signal of infection against a background of high host contamination. Although still uncommon in the field, performing technical replicates would be useful to assess variability, and even subjecting a subset of samples to replication may give enough information to disentangle technical from true variability.

Multiple methods are available for the generation of Illumina sequencing libraries: these are usually distinguished by the method of fragmentation used. Transposase-based “tagmentation”, for example in the Illumina Nextera and Nextera XT products, are popular owing to their low cost (list prices of $25-40 per sample, with dilution methods potentially able to reduce these costs even further). Tagmentation approaches only require small DNA inputs (1 ng of DNA recommended, but lower amounts can be used). Such low inputs are achieved due to a subsequent PCR amplification step. However, as tagmentation targets
specific sequence motifs it may introduce amplification biases along with the well-known GC content biases associated with PCR. One way of reducing these biases is to use a PCR-free method relying on physical fragmentation (e.g. PCR-free TruSeq) to produce a sequencing library that may be more representative of the underlying species composition in a sample.

There are no published guidelines for the “correct” amount of coverage for a given environment or study type, and it is unlikely that such a figure exists. As a rule of thumb, we therefore often recommend choosing a system that maximizes output in order to retrieve sequences from as many low-abundance members of the microbiome as possible. Illumina HiSeq 2500/4000, NextSeq, and NovaSeq all produce high volumes of sequence data (between 120 gigabases and 1.5 terabases per run) and are well suited for metagenomics studies (with the caveat of index hopping). The throughput per run of these instruments is known and, by deciding the level of multiplexing, the investigator can set the desired per-sample sequencing depth. Typical experiments in 2017 aim to generate between 1 and 10 gigabases, but these depths may be either excessive or woefully little depending on the sensitivity required to detect rare members of a sample.

The Illumina platforms mainly differ by their total output and maximum read length. The Illumina HiSeq 2500, although now two generations old, is a popular choice for shotgun metagenomics as it is able to generate 2x250 nt in rapid run mode (generating up to 180 Gb per flowcell), or up to 1Tb in high output mode with 2x125 nt reads. The newer HiSeq 3000 and 4000 systems further increase the overall throughput of a run (up to 1.5 terabases for the 4000) but are limited to read lengths of 150 nt. The NextSeq benchtop instrument has similar output to the HiSeq 2500’s rapid run mode, but are limited to 150 nt reads. However the NextSeq is less than half the price of the HiSeq and so may be attractive to research groups wishing to operate their own instrument. The recently released NovaSeq platform promises up to 3 terabases per run in the near future. The Illumina MiSeq is limited by output (up to 15Gb in 2x300 mode) but remains the de facto standard for single marker gene microbiome studies. The MiSeq (or MiniSeq) may still be useful for metagenomics for sequencing a limited number of samples or to assess library concentrations and barcode pool balancing, providing confidence of good results, before running on the higher-throughput (and much more expensive) instruments where individual runs may cost >$10,000.

**Metagenome assembly**

Numerous approaches to computationally reconstruct the composition of the microbial community from the pool of sequence reads have been published. Choosing the “best” approach is a daunting task but largely depends on the aims of the study.

Metagenome *de novo* assembly, is conceptually similar to whole genome assembly. The de Bruijn graph approach is currently the most popular metagenome assembly method. For single draft genome assemblies a de Bruijn graph is constructed by breaking each sequencing read into overlapping subsequences of a fixed length k. This set of overlapping “k-mers” defines the vertices and edges of the de Bruijn graph. The assembler’s task is to find a path through the graph that reconstructs the genome(s). This task is complicated by sequencing errors, which generate non-genomic sequences that must be avoided, and repetitive sequence, which can cause misassemblies and fragmentation of the assembly.

Metagenome assembly presents challenges not faced in single genome assembly. First, when assembling a single genome it is typically assumed that sequence coverage along the genome will be approximately
uniform. An assembler can use sequence coverage to identify repeat copies, distinguish true sequence from sequencing errors 42 and identify allelic variation 43. Metagenome assembly is more difficult because the coverage of each constituent genome depends on the abundance of each genome in the community. Low abundance genomes may end up fragmented if overall sequencing depth is insufficient to form connections in the graph. Using a short k-mer size in graph formation can assist in recovering lower abundance genomes, but this comes at the expense of increasing the frequency of repetitive k-mers in the graph, obscuring the correct reconstruction of the genomes. The assembler must strike a balance between recovering low-abundance genomes and obtaining long, accurate contigs for high abundance genomes. A second problem is that a sample can contain different strains of the same bacterial species. These closely related genomes can cause branches in the assembly graph where they differ by a single nucleotide variant, or by the presence/absence of an entire gene or operon. The assembler will often stop at these branch points, resulting in fragmented reconstructions.

Metagenome-specific assemblers try to overcome these challenges. Meta-IDBA 44 uses a multiple k-mer approach to avoid the difficult task of choosing a k-mer length that works well for both low and high abundance species. Meta-IDBA has extensions to partition the de Bruijn graph (as does MetaVelvet 45) and the latest version, IDBA-UD, optimizes the reconstruction for uneven sequence depth distributions 46. The SPAdes assembler 47 has been extended for metagenome assembly and can be used for assembling libraries sequenced with different technologies (hybrid assembly).

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Metagenomic assembly method</th>
<th>Assembly statistics for contigs longer than 1kb (values in parenthesis refers to perfect contigs)</th>
<th>Reconstruction %</th>
<th>N50</th>
<th>% identity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td># contigs</td>
<td>Total assembly size</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Env. Mock community 55</td>
<td>MetaSPAdes</td>
<td>16.22k (11.26k)</td>
<td>150.47M (108.39M)</td>
<td>80.93% (58.30%)</td>
<td>26.46k (25.88k)</td>
</tr>
<tr>
<td></td>
<td>MegaHit</td>
<td>21.82k (16.67k)</td>
<td>146.72M (124.67M)</td>
<td>78.91% (67.05%)</td>
<td>16.94k (17.94k)</td>
</tr>
<tr>
<td>HMP Mock community 2</td>
<td>MetaSPAdes</td>
<td>0.72k (0.42k)</td>
<td>62.67M (31.95M)</td>
<td>95.15% (48.50%)</td>
<td>260.45k (178.28k)</td>
</tr>
<tr>
<td></td>
<td>MegaHit</td>
<td>1.43k (1.14k)</td>
<td>62.09M (54.56M)</td>
<td>94.27% (82.84%)</td>
<td>124.02k (113.11k)</td>
</tr>
</tbody>
</table>

Table 2: Comparative evaluation of metagenomic assembly on mock microbial communities with known composition.

For complex samples that are likely to contain hundreds of strains, the sequencing depth must be increased as much as possible. Computational time and memory may be insufficient to complete such assemblies. Distributed assemblers 48 such as Ray, which spread memory load over a cluster of computers, have been used to assemble metagenomes from human faecal samples 49. To help assemble very complex samples Pell et al. developed a lightweight method to partition a metagenome assembly graph into connected components that can be assembled independently 50. Another method, named Latent Strain Analysis, partitions reads using k-mer abundance patterns which enables assemblies of individual low-abundance

1 'perfect contigs' are those contigs reconstructed by metagenomic assembly that have a match with >99% identity with the reference genome over the full length of the contig. Notably, 'perfect contigs' excludes chimeric contigs.

2 The N50 value corresponds to the size of the contig for which longer contigs represent at least half of the total assembly.
genomes using a limited amount of memory. MegaHIT uses succinct data structures to reduce the memory requirements of assembling complex metagenomes and achieves very quick run times.

There is little community consensus on how well different assemblers perform with respect to key metrics such as completeness, continuity and propensity to generate chimeric contigs. Despite metagenomic analysis “bake-offs” aimed at making concrete recommendations for analysis software, it is likely that software performance will depend on biological factors such as underlying microbial community structure, and technical factors, such as sequencing platform characteristics and coverage. This effect was observed at an Assemblathon, where no single assembler came out “best”.

<table>
<thead>
<tr>
<th>Sample 3</th>
<th>Assembler</th>
<th>#genes 4</th>
<th>#matches against nr (95% identity)</th>
<th># of species observed (nr at 95% identity)</th>
<th>Median # of single core genes</th>
<th># of annotated COGs</th>
<th># of annotated KEGG orthologues</th>
</tr>
</thead>
<tbody>
<tr>
<td>Env Mock community 55</td>
<td>MetaSPAdes</td>
<td>164750</td>
<td>154403</td>
<td>103</td>
<td>49.5</td>
<td>100681</td>
<td>91376</td>
</tr>
<tr>
<td>Env Mock community 55</td>
<td>MegaHIT</td>
<td>164146</td>
<td>154185</td>
<td>105</td>
<td>49</td>
<td>97119</td>
<td>91035</td>
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<tr>
<td>HMP Mock community 2</td>
<td>MetaSPAdes</td>
<td>62850</td>
<td>61362</td>
<td>30</td>
<td>20</td>
<td>44625</td>
<td>36082</td>
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<tr>
<td>HMP Mock community 2</td>
<td>MegaHIT</td>
<td>63304</td>
<td>61617</td>
<td>38</td>
<td>20</td>
<td>44289</td>
<td>36394</td>
</tr>
<tr>
<td>Gut sample 2</td>
<td>MetaSPAdes</td>
<td>169399</td>
<td>111119</td>
<td>365</td>
<td>44.5</td>
<td>79414</td>
<td>76500</td>
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<tr>
<td>Gut sample 2</td>
<td>MegaHIT</td>
<td>166289</td>
<td>109777</td>
<td>381</td>
<td>41.5</td>
<td>77666</td>
<td>75020</td>
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<td>Ocean sample 6</td>
<td>MetaSPAdes</td>
<td>124251</td>
<td>7397</td>
<td>118</td>
<td>42</td>
<td>51138</td>
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<td>Ocean sample 6</td>
<td>MegaHIT</td>
<td>151627</td>
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<td>60.5</td>
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<td>87344</td>
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<td>MetaSPAdes</td>
<td>34118</td>
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<td>10448</td>
<td>15312</td>
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<tr>
<td>Soil sample 129</td>
<td>MegaHIT</td>
<td>44396</td>
<td>11008</td>
<td>132</td>
<td>11.5</td>
<td>17671</td>
<td>22524</td>
</tr>
</tbody>
</table>

Table 3: Comparative evaluation of metagenomic assembly of a set of metagenomes from diverse environments. Functional annotations performed as previously described.

We analysed assembly results from mock synthetic and real communities (Table 2 and Table 3). We evaluated two assemblers, MegaHIT and MetaSPAdes for their ability to reconstruct known genomes from the mock communities, and capture taxonomic and gene diversity in the real datasets. They both successfully reconstructed more than 75% of the mock communities (one comprising 20 organisms, the other 49 bacterial and 10 archaeal species). MetaSPAdes generated longer contigs, but these appeared to be less accurate. When restricted to contigs that exactly matched the references in the mock community then MegaHIT succeeded in reconstructing more of the true genomes. Choice of assembler in this case would therefore depend on the relative importance of contig size versus accuracy. Across the true datasets (Table 3), consistent patterns were hard to discern. However, examining median single-copy core gene number (which will estimate the number of genomes in the assembly) suggests that for the more complex soil and ocean communities, MegaHIT succeeded in assembling more genes that could then be functionally annotated. However, the key message here is that different state-of-the-art programs will be optimal on

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3 All samples have been subsampled to 50 million reads for inter sample comparability
4 total number of genes identified from the assembled contigs using Prodigal
different datasets while requiring similar run times (about 48 hours using 16 threads on the largest sample) and main memory usage (not exceeding 125GB). It is prudent, therefore, to attempt more than one assembly approach. The CAMI challenge reported that MegaHIT was in the top three best metagenomics assemblers across their benchmark data sets and together with MetaSPAdes (not evaluated in CAMI) these are probably the best current choices. Whatever assembler is used the result will not be genomes but rather potentially millions of contigs, and this motivates the need for binners that attempt to link those contigs back into the genomes they derived from.

Binning contigs

Metagenome assemblies are highly fragmented, comprising thousands of contigs (Table 2), and the challenge is that we do not know a priori which contig derives from which genome. We do not even know how many genomes are present. The aim of contig “binning” is to group contigs into species. Supervised binning methods use databases of already sequenced genomes to label contigs into taxonomic classes. Unsupervised methods, or clustering, look for natural groups in the data.

Both supervised and unsupervised methods have two main elements: a metric to define the similarity between a given contig and a bin, and an algorithm to convert those similarities into assignments. For taxonomic classification, contig homology against known genomes is a potentially useful approach, but most microbial species have not been sequenced so a large fraction of reconstructed genomic fragments cannot be mapped to reference genomes. This has motivated the use of contig sequence composition for binning. Different microbial species’ genomes contain particular combinations of bases, and this results in different k-mer frequencies. Metrics based on these k-mer frequencies can be used to bin contigs, with tetramers considered the most informative for binning of metagenomics data. Many different software choices are available that are based on these frequencies such as Naïve Bayes classifiers or support vector machines, but sequence composition often lacks the specificity necessary to resolve complex datasets to the species level in complex communities.

Clustering of contigs is appealing because it does not require reference genomes. Until recently, most contig clustering algorithms such as MetaWatt and SCIMM used various species composition metrics, sometimes coupled with total coverage. Recently, as multi-sample metagenome datasets have been produced it has been realized that contig coverage across multiple samples provides a much more powerful signal to group contigs together. The principle is that contigs from the same genome will have similar coverage values within each metagenome, although intra genome GC content variation, and increased read depth around bacterial origins of replication, can challenge this assumption. The first algorithms, e.g. extended self-organising maps, required human input to perform the clustering, which is based on coverage information and composition that could be visualized in 2D. Completely automated approaches such as CONCOCT, GroopM and MetaBAT are now available and they are convenient, particularly for large datasets, but better results may still be obtained when combined with human refinement, for instance using a visualization tool named Anvio.
Methods for reconstructing metagenomic assembled genomes (MAGs) are indispensable to uncover the hitherto inaccessible diversity of bacteria. The recovery of nearly a thousand MAGs from candidate phyla, with no cultured representatives, from acetate enriched and filtered groundwater samples showcased the potential of this approach. Recovered genomes were all small, with minimal metabolism, and formed a monophyletic clade, separate from the previously cultured diversity of bacteria. These have been proposed as a new bacterial sub-division, the candidate phyla radiation, revealed through metagenomics.

Completeness of MAGs is usually evaluated by examining single-copy core genes, which are found in most microbial genomes, for example tRNA synthetases or ribosomal proteins. A pure MAG will have all these genes present in single copies. Once constructed, the MAGs provide a rich dataset for comparative genomics, including the construction of phylogenetic trees, functional profiles and comparisons of MAG abundance across samples (see left panel in Figure 2 and the step-by-step tutorial we provide at https://github.com/chrisquince/metag-rev-sup).

Assembly-free metagenomic profiling

Taxonomic profiling of metagenomes identifies which microbial species are present in a metagenome, and estimates their abundance. This can be carried out without assembly using external sequence data resources, such as publicly available reference genomes. This approach can mitigate assembly problems,
speed up computation, and make it possible to profile low-abundance organisms that cannot be assembled de novo (Supplementary Box 1). The main limitation is that previously uncharacterized microbes are very difficult to profile (Supplementary Box 1). However, the number of reference genomes available is increasing rapidly, with thousands of genomes being produced each year, including some derived from difficult-to-grow species targeted by new cultivation methods \(^{71}\), single-cell sequencing approaches \(^{72}\), or metagenomic assembly itself. The diversity of reference genomes available for some sample types, such as from the human gut \(^{73}\), is now extensive enough to make assembly-free taxonomic profiling efficient and successful, including for comparatively low abundance microbes that lack sufficient sequence coverage and depth to enable the assembly of their genome. Analysis of more diverse environments including soil and oceans is hampered by a lack of representative reference genomes. As a result, it is generally inadvisable to avoid assembly when analyzing metagenomes from these environments.

### Limitations of shotgun metagenomics

**“Entry-level access” issues.** It is still expensive to sequence and analyze large numbers of metagenomes without access to sequencing and computational facilities. Improved sequencing platforms and cloud computing facilities should decrease these entry-level costs.

**Comprehensiveness of genome catalogs.** The set of >50,000 microbial genomes available is biased toward model organisms, pathogens, and easily cultivable bacteria. All metagenomic computational tools, to some extent, rely on available genomes and they are thus affected by the biases in the reference sequence resources.

**Biases in functional profiling.** Profiling of the functional classes present in a metagenome is hindered by the lack of validated annotations for most genes, an issue that can be mitigated only by expensive and low-throughput gene-specific functional studies. Moreover, intrinsic microbiome properties such as its average genome size can critically impact the quantitative profiling \(^{102}\).

**Microbial dark matter.** Several members of a microbiome might have not been characterized before with culture-based methods or with metagenomics. This is regarded as microbial dark matter, and assembly-based approaches can recover part of this unseen diversity. A fraction of reads may still remain unused after assembly, and the size of this fraction is highly dependent on community structure and complexity (e.g. see the analysis reported in Table 2 and 3). It is also impacted by features such as sequencing noise, contaminant DNA, and microbes and plasmids that remain taxonomically obscure even after assembling part of their genome.

**“Live or dead” dilemma.** DNA persists in the environment after the death of the host cell, so the sequencing results may not be representative of the active microbial population. Compounds such as propidium monazide, which binds to free DNA, as well as DNA within dead or damaged cells, or techniques such as metatranscriptomics, may be used if the aim is to study the active microbes.

**“Curse of compositionality”.** Quantitative metagenomic features are reported as fractional values without links to the real absolute concentration. Variations in the true concentration of organisms across samples can thus produce false correlations. For example, if a highly abundant organism doubles its concentration in two otherwise identical samples, all the other organisms in the sample will appear to be differentially abundant after normalization.

**Mucosa-associated microbiome sequencing.** Human mucosal tissues are crucial interfaces between microbes and the immune system, but sequencing the mucosal microbiome with shotgun metagenomics is very challenging due to the extremely high fraction of human DNA and the low microbial biomass.

### Challenges in shotgun metagenomics

**Integrative meta-omics.** Complementing DNA sequencing with RNA, protein, and metabolomic high-throughput assays is possible with shotgun metatranscriptomics, mass-spectrometry-based metaproteomics and metabolomics \(^{74}\). Despite the potential of these technologies, it is unclear how to integrate and analyze meta-omic data within a common framework.

**Virome shotgun sequencing.** Viral organisms can be detected by shotgun metagenomics, but virome enrichment techniques are usually needed to access a broader set of viruses. Virome analysis is also computationally challenging because of limited availability of viral genomes and a lack of inter-family phylogenetic signals.

**Strain-level profiling.** The genomic resolution of single isolate sequencing is still higher than what can be achieved for single organisms in a metagenomic context. Increasing the profiling resolution to the level of single strains would be crucial for in-depth population genomics and microbial epidemiology.

**Longitudinal study designs.** Many shotgun metagenomic studies are cross-sectional and thus unpowered for assessing inter versus intra subject variability and microbiome temporal evolution. Tools for longitudinal settings have been developed \(^{41}\) but more methods and data are needed to investigate the temporal dimension \(^{113}\).

**Disentangling cause from effect.** Hypotheses from metagenomic studies should be followed up with experimental work to validate correlations and associations. Longitudinal and prospective settings can potentially provide direct insights into the causative dynamics of conditions of interest.

**Validation of microbiome biomarkers.** Microbiome biomarkers of a given condition are often strongly study-dependent. It is thus crucial to validate biomarkers across technologies and cohorts to enhance reproducibility and minimize batch effects.

**Data sharing, open data, open source, and analysis reproducibility.** Data and metadata sharing is strongly encouraged, raw data deposition is usually requested prior to publication, and open source software is desirable. However, metagenomics has still to reach the level of standardization that is characteristic of other more established high-throughput techniques.

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**Box 1. Limitations and opportunities in metagenomics.**
Assembly-free taxonomic profilers with species-level resolution utilize information available in reference genomes and in environment-specific assemblies, and have been used in the largest human-associated metagenomics investigations performed so far. The simple brute force mapping of reads to genomes can result in profiles with many false positives but, nonetheless, this approach has been proven to be effective when the output is post-processed based on lowest common ancestor (LCA) strategies or coupled with compositional interpolated Markov models. However, the run times of these approaches do not improve on assembly-based methods. Kraken also exploits LCA but dramatically speeds up the computation by substituting sequence matching with k-mer matching.

Taxonomic profiling by selecting representative or discriminative genes (markers) from available reference sequences is another fast and accurate assembly-free approach that has been implemented with several variations. By looking at co-abundant markers from pre-assembled environment specific gene catalogs, for example, the MetaHIT consortium was able to characterize known and novel organisms in the human gut. Similarly, mOTU focuses on universally conserved but phylogenetically informative markers (e.g., genes coding for ribosomal proteins), whereas MetaPhlAn (right panel of Figure 2) adopts several thousands of clade-specific markers with high discriminatory power, and proved effective to quantitatively profile the microbiome from multiple body areas for the Human Microbiome Project with a very low false positive discovery rate. These methods are scalable and can be used for large metagenomics meta-analyses. Marker-based approaches can also be used for strain-level comparative microbial genomics using thousands of metagenomes. Importantly, the accuracy of these methods will improve as more reference genomes and high-quality metagenomic assemblies become available. For large datasets with hundreds of samples on which performing or interpreting metagenomics assembly is impractical, marker-based approaches are currently the method of choice especially for environments with a substantial fraction of microbial diversity covered by well-characterized sequenced species.

**Genes and metabolic pathways from metagenomes**

With a fragmented but high-quality metagenome assembly, the gene repertoire of a microbial community can be identified using adaptations of single-genome characterization tools. These include a gene identification step, usually with a metagenomic-specific parameter setting, followed by homology-based annotation pipelines commonly used for characterizing pure isolate genome assemblies. Indeed, some of the largest shotgun sequencing efforts performed so far used metagenomic assemblies to compile the microbial gene catalog of the human and mouse gut metagenomes, although this approach is often limited by the large fraction of uncharacterized genes in the reference database catalogs.

Other large metagenomic datasets were interpreted by translated sequence searches against functionally characterized protein families. Databases, that include combinations of manually annotated and computationally predicted proteins families such as KEGG or UniProt, can be used for this task and enable characterization of the functional potential of the microbiome (Figure 2, right-hand panel). Single protein families are aggregated into higher-level metabolic pathways and functional modules providing either graphical reports or comprehensive metabolic presence/absence and abundance tables, as in the HUMAN pipeline. Regardless of whether an assembly-free or assembly-based approach is adopted, the main limiting factor in profiling the metabolic potential of a community is the lack of annotations for accessory genes in most microbial species (with the exception of selected model organisms, Box 1). This
means that highly conserved pathways and housekeeping functions are more consistently detected and quantified in metagenomes, which might explain why functional traits are often reported to be surprisingly consistent across different samples and environments, even when taxonomic composition is highly variable \(^2\). Experimental characterization of microbial proteins, coding genes, and other genomic features (tRNAs, non-coding RNAs, CRISPRs) to more thoroughly assess functions of individual loci is a bottleneck that currently has a crucial impact on our ability to profile the functions of metagenomes \(^85\).

A complementary approach to metabolic function profiling of metagenomes is an in-depth characterization of specific functions of interest. For example, identifying genes involved in antibiotic resistance (the “resistome”) in a microbial community can inform on the spread of antibiotic resistance \(^97\). Ad-hoc methods \(^98\) and manually curated databases of antibiotic resistance genes have been crucial to this approach; ARDB \(^99\) was the first widely adopted resistance database and is now complemented by additional resources such as Resfams \(^100\). Comparably large efforts are also devoted to reporting the virulence repertoire of a metagenome; targeted analyses of metagenomes for specific gene families of interest can also be used to validate findings from single, cultivation-based isolate experiments.

**Post-processing analysis**

Regardless of the methods used for primary metagenomic sequence analyses, the outputs will comprise data matrices of samples versus microbial features (species, taxa, genes, pathways). Post-processing analysis uses statistical tools to interpret these matrices, and decipher how the findings correlate with the sample meta-data. Many of these statistical approaches are not specific for metagenomics. Specific challenges of metagenome-derived quantitative values include the proportional nature of the taxonomic and functional profiles, and the log-normal long-tailed distribution of abundances. These issues are also problematic in high-throughput 16S rRNA gene amplicon sequencing datasets, and several popular R packages such as DESeq2 \(^101\), vegan \(^102\), and metagenomeSeq \(^103\) that were originally developed for amplicon sequencing can be used for metagenomics.

Post-processing tools include traditional multivariate statistics and machine learning. Unsupervised methods include simple clustering and correlation of samples, and visualization techniques such as heatmaps, ordination (e.g. PCA and PCoA), or networks, which allow the patterns in the data to be revealed graphically. Some unsupervised statistical tools aim to specifically address the problems introduced by the proportional nature of metagenome profiles (compositionality issue \(^104\), Box 1) and try to infer ecological relationships within the community \(^105\). Supervised methods include both statistical methods such as multivariate analysis of variance ANOVAs for direct hypothesis testing of differences between groups, or machine learning classifiers that train models to label groups of samples, such as Random Forests or Support Vector Machines \(^106\). A classic machine learning example would be to diagnose disease (e.g. for type 2 diabetes \(^76\)) on the basis of community dysbiosis, although developing cross-study predictive signatures is challenging \(^106\).

Unsupervised and supervised methods consider the community as a whole. A complementary strategy is to ask which specific taxa or functional genes are statistically different between sample types or patient groups. Given the complexity of metagenomics datasets, and the huge numbers of comparisons that can typically be made, correction for multiple comparisons \(^107\) or effect size estimation \(^108\) are vital for this task.
Robust statistical testing is key to determining the validity of results, but compact graphical representations can intuitively reveal patterns. In many cases visualization of post-processing results requires ad-hoc graphical tools, and carefully adopted general visualization approaches.

Outlook

Metagenomics still faces roadblocks to applicability, usefulness, and standardization (Box 1). The lack of reference genome sequence data for large portions of the microbial tree of life, or functional annotation for many microbial genes, substantially reduce the potential for success of the computational approaches used to analyze the vast amounts of sequences produced. Metagenomes from environments such as soil or water are particularly affected by this problem owing to both their high microbial diversity, and the proportion of uncharacterized taxa in these communities. Shotgun sequencing also fails to discriminate between live and dead organisms. However, the outlook is bright, because year on year a large community of wet-lab and computational researchers are finding solutions to these problems.

<table>
<thead>
<tr>
<th></th>
<th>Assembly-based analysis</th>
<th>Read-based analysis (“Mapping”)</th>
</tr>
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<tbody>
<tr>
<td>Comprehensiveness</td>
<td>Can construct multiple whole genomes but only for organisms with enough coverage to be assembled and binned</td>
<td>Can provide an aggregate picture of community function or structure, but is only based upon the fraction of reads that map effectively to reference databases</td>
</tr>
<tr>
<td>Community complexity</td>
<td>In complex communities only a fraction of the genomes can be resolved by assembly</td>
<td>Can deal with communities of arbitrary complexity given sufficient sequencing depth and satisfactory reference database coverage</td>
</tr>
<tr>
<td>Novelty</td>
<td>Can resolve genomes of entirely novel organisms with no sequenced relatives</td>
<td>Cannot resolve organisms for which genomes of close relatives are unknown</td>
</tr>
<tr>
<td>Computational burden</td>
<td>Assembly, mapping and binning are all computationally costly steps</td>
<td>Can be performed efficiently, enabling large meta-analyses</td>
</tr>
<tr>
<td>Genome resolved</td>
<td>Can link metabolism to phylogeny through completely assembled genomes, even for novel diversity</td>
<td>Can only typically resolve the aggregate metabolism of the community, links with phylogeny are only possible in the context of known reference genomes</td>
</tr>
<tr>
<td>metabolism</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Expert manual</td>
<td>Manual curation required for accurate binning/scaffolding, and for misassembly detection</td>
<td>Manual curation usually not needed, although the selection of reference genomes to use could involve human supervision.</td>
</tr>
<tr>
<td>Supervision</td>
<td></td>
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<tr>
<td>Integration with</td>
<td>Assemblies can be fed into microbial genomic pipelines designed for analysis of genomes from pure cultured isolates</td>
<td>Obtained profiles cannot be directly put into the context of genomes derived from pure cultured isolates</td>
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<tr>
<td>Microbial genomics</td>
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*Table 4. Strengths and weaknesses of assembly-based and read-based analyses for primary analysis of metagenomics data.*

Metagenome bioinformatics tools, especially for translating raw reads into meaningful microbial features (genomes, species abundances, functional potential profiles) (Figure 1), are continually improving. For example, strain-level analyses are now possible. There remains an active debate about which sequence analysis approach is best (see Table 4). Metagenomic assembly is the preferred theoretical solution if there is sufficient genome coverage (i.e. >20x), but this level of coverage is difficult to obtain for most of the members of the microbiome (Table 4) and assembly-free methods have other advantages.
including the possibility to perform large-scale strain-level analyses. The success of either approach depends on the microbial community composition and complexity, sequencing depth, size of the dataset, and available computational resources (Table 4). We recommend that researchers use both approaches for sequence analysis whenever possible, as they complement and validate each other.

As for the technological improvements in the sequencing of community DNA, long-read sequencing platforms have matured and are likely to become useful for metagenomics assembly strategies, although publications are few at present. The Pacific Biosciences instruments can deliver complete or nearly complete isolated microbial genomes with low base error rates if sufficient coverage is achieved (typically 30-100X). The Oxford Nanopore MinION single molecule, long read instrument holds appeal because of its size and portability (smartphone size) and early analysis of reads from this platform indicates it has an error rate akin to Pacific Biosciences reads \(^{114}\). Assembly of isolate genomes is possible into single contigs \(^{115}\) so the portability of the MinION raises the tantalizing possibility of performing metagenomic sequencing in the field.

An alternative experimental approach to improve genome reconstruction from metagenomes couples Illumina sequencing chemistry with a multiplexed pooling library preparation protocol. This so-called Synthetic Long Reads technology relies on the dilution of genomic DNA into fragmented and barcoded pools consisting of hundreds to thousands of individual molecules. These pools are sequenced and assembled de novo to produce synthetic long reads. One benefit of synthetic long reads is that because they are built from a consensus of Illumina sequences, the base error rate is extremely low. However, the protocol is rather laborious and requires high DNA input (between 1 and 10 µg of DNA), plus, problems persist with local repetitive sequences. Reports suggest that this approach is useful for metagenomics, especially when coupled with standard shotgun sequencing, as it can reconstruct genomes from closely related strains, as well as those from rare microorganisms \(^{116,117}\).

Another outstanding problem in shotgun metagenomics is the accurate reconstruction of strain-level variation from mixtures of genetically related organisms \(^{118}\), with several solutions proposed \(^{14,90,111-113,119,120}\) that are based on assembly, mapping, or a combination of the two. Mapping to genes that are unique to a species \(^{88}\) can resolve the dominant haplotype in a sample, and this method has been applied to thousands of unrelated metagenomes, providing strain-level phylogenies that enable microbial population genomics for hundreds of largely uncharacterized species \(^{111}\). Mixtures of strains from the same species in a single sample cannot be resolved by consensus approaches, but if the same strains are present in multiple samples there will be characteristic signatures in single nucleotide variations. These nucleotide variations can be linked together to deduce haplotypes and their frequencies \(^{90,113,119}\). This methodology was initially only applied after mapping to reference genes \(^{90}\), and optionally with simultaneous strain phylogeny reconstruction \(^{119}\), but it has now been applied directly to contig bins with inference of strain gene complement in an entirely reference free method \(^{113}\). One limitation of this approach is that in some environments, including the human gut, it has been shown that one strain usually dominates over other strains from the same species \(^{111}\). It is therefore challenging to detect non dominant strains of low-abundance species, and the user has to weight the increased robustness of profiling only the dominant strains \(^{111}\) with the potential additional information that can be garnered from characterizing mixtures of strains \(^{113}\). Strain-level metagenomics is an active area of research \(^{118}\) and has the potential to empower metagenomics with similar resolution to that which can be derived from sequencing of pure culture single
isolates. Although long read technologies can aid these efforts in the future, solving the computational challenges of strain-level profiling from metagenomics is arguably the biggest challenge in the field at the moment.

Conclusions

Since the pioneering application of whole DNA sequencing to environmental samples by teams led by Jillian Banfield \textsuperscript{121} and Craig Venter \textsuperscript{7} in 2004, shotgun metagenomics has become an important tool for the study of microbial communities. Widespread adoption of metagenomics has been enabled by the falling cost of sequencing and the development of tractable computational methods. The main limitations facing researchers now are the costs of training computational scientists for analyzing the complex metagenomic datasets, and of sequencing enough samples for properly powered study designs. Initiatives such as the Critical Assessment of Metagenomic Interpretation \textsuperscript{56} are vital for an unbiased assessment of computational tools to improve reproducibility and standardization.

Shotgun metagenomics will play an increasingly important part in diverse biomedical and environmental investigations and applications. We hope that this Review will provide an understanding of the basic concepts of shotgun metagenomics including both its limitations and its immense potential.

Acknowledgments

AWW and The Rowett Institute, University of Aberdeen, receive core funding support from the Scottish Government’s Rural and Environmental Science and Analysis Service (RESAS). NS is supported by the European Research Council (ERC-STG project MetaPG), European Union FP7 Marie-Curie grant (PCIG13-618833), MIUR grant FIR RBFR13EWWI, Fondazione Caritro grant Rif.Int.2013.0239, and Terme di Comano grant. CQ and NL are funded through a MRC bioinformatics fellowship (MR/M50161X/1) as part of the MRC Cloud Infrastructure for Microbial Bioinformatics (CLIMB) consortium (MR/L015080/1). JTS is supported by the Ontario Institute for Cancer Research through funding provided by the Government of Ontario.

Author Contributions

C.Q., A.W.W., J.T.S., N.J.L, and N.S. drafted the sections, revised the text, and designed figures, tables, and boxes. C.Q. and N.S. performed the metagenomic analyses described in the manuscript.

Competing Financial Interest

The authors declare no competing financial interests.
Supplementary Figure 1. Example workflow for planning a metagenomics study. The advice presented here is targeted towards entry-level researchers in this area, with a particular focus on hypothesis-driven experiments, which of course may be designed very differently compared to exploratory/hypothesis-generating studies. Key considerations for study design (blue box), sample collection (green box) and experimental procedures (yellow box) are highlighted. Understanding the potential for confounding factors, and optimization of design, can substantially improve the quality of both metagenomic sequence data, and interpretation. Supplementary Table 1 contains further specific recommendations.
Supplementary Box 1. Common difficulties in Study design: problems and some recommended solutions.

**Powering the study / Read depth requirements.** The number of samples and sequencing depth required to be able to detect significant differences will depend on factors such as consistency of microbiome composition between different samples, the inherent microbial diversity of the samples, and effect size of the phenomenon being studied. **Solution:** These decisions can often be guided by results from previous studies in the same type of environment. In cases where this information is lacking it may be prudent to carry out preliminary marker gene-based studies to gauge the relative impact of each of the factors listed opposite.

**Confounding variables and control groups.** It is often very difficult to select a control group to compare against the samples of interest that is free from other confounding variables. An example of this is rodent microbiome research, where cage and animal batch effects can result in dramatic differences in microbiome composition, independent of the variable being studied. Another example is the cross-sectional study of the microbiome associated with a disease for cases in which the patients cannot be sampled in the absence of active treatment. **Solution:** Current best practice is to collect as much metadata about each of the study groups as possible and factor these into the subsequent analyses when comparing groups. For clinical samples this typically includes features such as gender, age, antibiotic/medication use, location, dietary habits, and Bristol stool chart scores. For environmental samples this commonly includes associated parameters such as geographic location, season, pH, temperature etc. Further extensive advice for planning rodent microbiome studies is available. Longitudinal sampling from the same patient/location can also act as an additional control, especially when longitudinal changes can be correlated with associated metadata.

**Sample collection/preservation.** It may be difficult to process and store all samples in exactly the same way (for example when samples are provided from a number of locations by different research groups). With longitudinal studies, samples collected at the final time point may spend less time in frozen storage prior to DNA extraction than samples collected at other time points. Such changes in sampling and preservation procedures may introduce systematic biases. **Solution:** Where possible, collection and preservation methodologies should be standardized throughout for all samples within a given study. All procedures used should also be recorded and included as pertinent metadata when carrying out subsequent data analyses. This should ideally include factors such as time between collection and DNA extraction, length of time in frozen storage, and number of freeze-thaw cycles. For mammalian gut samples there is some evidence that storage in glycerol may result in more representative compositional results following long term frozen storage. Similarly, freeze drying prior to long-term frozen storage may be a prudent approach.

**Biomass/Contamination.** Modern sequence based technologies are highly sensitive, meaning very small amounts of DNA are sufficient for sequencing. However, common laboratory kits and reagents are not sterile, meaning that any contamination that is present in these can potentially overwhelm the “real” signal in samples containing only a very low microbial biomass. **Solution.** It is prudent to gauge the level of biomass present in samples before sequencing using a quantitative approach such as qPCR. Samples containing fewer than $10^4$ microbial cells appear to be most impacted by background contamination. Table 1 offers some approaches that may be tried in order to enrich cell numbers/DNA yields from samples prior to sequencing. Negative control samples, that have been processed using the same kits/reagents as the actual samples, should be sequenced in order to determine the types of contaminating microbes present. Sequence data derived from these contaminants might then be removed bioinformatically from the final sequence datasets. Note that the sensitivity of these negative controls can be enhanced by the use of carrier DNA.

**Choice of DNA extraction methodology.** This step can hugely impact the results of a metagenomics study. If the approach selected is not stringent enough to extract DNA from some cell types they will not be represented accurately in the subsequent sequence data. Fundamentally, the optimal type of DNA extraction approach will depend on the underlying composition of the cell types that are present within a given sample. Unfortunately this can vary greatly, even within the same type of sample (e.g. the faeces of some humans are dominated by Gram negative species with cell walls that are relatively easy to disrupt, while those of others are dominated by relatively recalcitrant Gram positive species). As a result, no one DNA extraction approach will work optimally for all sample types. **Solution:** The use of defined mock community controls consisting of cultures derived from a mixture of the types of species that are common within a given environment can be a useful starting point to test the efficiency and accuracy of different DNA extraction methods. Mock communities can be optimized by including a phylogenetically diverse collection of species that are known to be commonly abundant in the sample type being studied. However, it is difficult to mimic the complexity of real microbial communities using simplified mocks, and impossible to test for the efficiency of the extraction step for unknown/uncultured organisms. Much evidence suggests that incorporating a bead-beating step into the DNA extraction process improves yield and representativeness of resulting species profiles compared to chemical-only lysis (ref#133 C.O.,N.J.L.). However, this type of approach does typically result in more sheared DNA, potentially limiting the power of burgeoning long read sequencing technologies. DNA extraction methodology should also be included as crucial metadata when uploading sequence data to public repositories. This allows variance in methodology choices to be factored into subsequent meta-analyses that incorporate metagenomic datasets from different laboratories.


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