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Accessibility
MetaRef: a pan-genomic database for comparative and community microbial genomics

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ABSTRACT

Microbial genome sequencing is one of the longest-standing areas of biological database development, but high-throughput, low-cost technologies have increased its throughput to an unprecedented number of new genomes per year. Several thousand microbial genomes are now available, necessitating new approaches to organizing information on gene function, phylogeny and microbial taxonomy to facilitate downstream biological interpretation. MetaRef, available at http://metaref.org, is a novel online resource systematically cataloguing a comprehensive pan-genome of all microbial clades with sequenced isolates. It organizes currently available draft and finished bacterial and archaeal genomes into quality-controlled clades, reports all core and pan gene families at multiple levels in the resulting taxonomy, and it annotates families’ conservation, phylogeny and consensus functional information. MetaRef also provides a comprehensive non-redundant reference gene catalogue for metagenomic studies, including the abundance and prevalence of all gene families in the >700 shotgun metagenomic samples of the Human Microbiome Project. This constitutes a systematic mapping of clade-specific microbial functions within the healthy human microbiome across multiple body sites and can be used as reference for identifying potential functional biomarkers in disease-associate microbiomes. MetaRef provides all information both as an online browsable resource and as downloadable sequences and tabular data files that can be used for subsequent offline studies.

INTRODUCTION

High-throughput sequencing has become an invaluable tool for scientific investigation, and computational resources are still adapting to the ubiquity and scale of modern genomic resources. This has been particularly true for microbial sequencing, where new methods must be developed to organize tens of thousands of genomes, hundreds of isolates per species and a pan-genome that comprises millions of gene families. Many databases catalogue microbial sequences, but few yet place these sequences within their phylogenetic and functional context.

For example, many microbial clades have now been shown to have surprisingly large pan-genomes in contrast to the size of their core genomes (1). Any one Escherichia coli isolate typically contains ~4700 genes, but only some 2000 of these are found in all E. coli, and the pan-genome selected from all strains in the species now greatly exceeds 8000 genes (2). Yet current microbial genome resources rarely indicate the core or pan-genomes of a clade of interest, nor do they conversely assess the phylogenetic distribution of individual gene families.

No comprehensive tool is thus available for interrogation of microbial clade-specific gene sequences, families or functional annotations. Here, we describe the algorithms, database and online interface for such a framework, initially detailing over 10 million genes from 2800 microbial genomes. All information is both browsable interactively through an online web interface and downloadable for offline analysis, and the underlying data are automatically updated every 6 months.

OVERVIEW OF THE METAREF DATABASE

MetaRef is a novel online resource reporting a systematic sequence-based catalogue of the diversity and
characteristics of the gene repertoires of all microbial clades with available whole-genome sequence reference information. It identifies and reports each gene family present in at least one genome, organizing them into pan, core or marker families, reconciles their functional annotations when possible, and systematically surveys their presence in multiple body sites of the human microbiome.

MetaRef serves as a comprehensive and convenient resource for tasks in microbial genome investigations, comparative genomics, and quantitative microbial ecology and offers several features not provided by existing systems. The MetaRef microbial gene family system (defined at 80% full-length nucleotide identity) includes every annotated reading frame in currently sequenced genomes, including singleton and uncharacterized genes. The underlying gene family clustering algorithm is defined hierarchically so as to scale efficiently to the increasing number of sequenced genomes (currently >2800); many existing catalogues such as COG (3) and KEGG KOs (4) neither match this scale nor typically annotate more than a fraction of genes in any one genome. Unsupervised approaches such as KEGG OC (5), PHOG (6) and OMA (7) face scalability issues and are often only appropriate for finished genomes. The largest current databases, such as MBGD (8), eggNOG (9) and UniRef (10), extend these systems to all proteins in a larger number of input genomes, but uniformly at the cost of a loss of transferable information—all provide comprehensive definitions of microbial gene families, but none link these families to phylogenetic and functional information for downstream interpretation.

The novel resource provides information useful for at least two different classes of investigator. For microbiologists focusing on individual microbial clades (e.g. a genus or a species), MetaRef provides a pre-computed pan-genome comprising core families consistently present in the clade, marker families present uniquely in the clade, and pan families present only in some clade members. This constitutes a tree-of-life-wide resource for phylogenetic analysis (e.g. by aligning and concatenating MetaRef core families) with several additional features unique to the MetaRef system including consensus-based functional characterization and pre-computed conservation/diversity scores. Second, for investigators focusing on large-scale comparative genomics or ecology, MetaRef also provides a catalogue of pre-identified marker genes for microbiome taxonomic profiling (11), and the means to assess the relevance of any microbial gene families with respect to their symbiotic healthy relationship with the human body, as the system reports the abundance and prevalence of each family in each clade in six body sites of the healthy human microbiome integrating shotgun metagenomic data produced by the Human Microbiome Project (HMP) (12). No existing resources for microbial genomics provide this level of clade-specific detail and whole-microbiome analysis.

MetaRef thus provides a new resource based on automatic and unsupervised complete gene clustering, processing a highly scalable number of sequenced microbial genomes. Both final and draft genomes are included in this process; open reading frame (ORF) calls are the only necessary annotations. The MetaRef web interface has been developed to provide a variety of clade-centric and gene family-centric interactive views to access and explore the sequenced microbial diversity. All MetaRef information, including sequence data, annotations and metagenomic mapping, is also available for download in plain text format for offline processing. While MetaRef provides novel aspects such as clade-specificity for the gene family system and metagenomic surveys of each gene family considered, it also combines four features currently not present simultaneously in other resources: (i) it comprehensively processes all available microbial genomic data; (ii) it provides both online and downloadable sequence information; (iii) it efficiently scales to the increasing throughput of reference genomes; and (iv) it leverages functional and phylogenetic information to gain insights into each gene family.

**DATABASE GENERATION, CONTENT AND DEFINITIONS**

The MetaRef database and online resource are built upon publicly available microbial genomic and metagenomic information automatically acquired and processed on a regular basis (Figure 1). The computational pipelines we implemented process the ~3000 microbial genomes available (from IMG (13)) with the associated taxonomic and gene information and the ~3.5TB of shotgun metagenomic data of 757 human microbiome samples (from the HMP (14)). These pipelines include PhyloPhlAn (15) for taxonomic curation and a novel clustering pipeline for pan gene classification (11,16) (see Supplementary Methods). The second phase of computational processing comprises metagenome mapping using a Bowtie-based strategy (17), a novel taxonomic and phylogenetic tree visualization (GraPhlAn, publicly available at http://huttenhower.sph.harvard.edu/graphlan), and the derivation of consensus functional annotations. The MetaRef online interface is a dynamic website with a back-end MySQL relational database, developed using Django (version 1.4.3 https://www.djangoproject.com); (see Supplementary Methods for implementation details).

**The MetaRef system: definition of marker, core, and pan genes and families**

The phylogenetic diversity within each microbial taxonomic clade is captured by cataloguing the overall repertoire of distinct ORFs contained in the pan-genome (18,19). MetaRef naming conventions refer to individual microbial ORF sequences as genes, and groups of genes related by some homology criterion as families. Pan genes are defined as ORFs present at least once in at least one genome of a clade of interest. Pan genes are usually grouped into homologous groups, called pan families, which capture a combination of paralogy, orthology and horizontal gene transfer relationships through sequence clustering requiring a full gene length similarity threshold (20). Among the pan genes forming a pan family, the ORF minimizing the dissimilarity with respect to all the other pan genes is called the pan centroid.
Some pan families are conserved within all organisms of a clade and are thus defined as core families (Figure 2). When a core family for a clade (say the \textit{Staphylococcus aureus} species) is not conserved in its direct ancestor clade (in this case the genus \textit{Staphylococcus}), it is of biological interest as a potentially conserved function distinguishing the clade from sibling clades (e.g. \textit{Staphylococcus epidermidis} and other \textit{Staphylococcus} spp.) and is called a crown family (Figure 2). A crown family can still be simultaneously present in unrelated branches of the taxonomy, possibly as a consequence of evolutionarily successful horizontal gene transfer events or complex gene loss patterns. However, when a crown family appears to be uniquely present in a given clade, we define it as a marker family (11), containing genes that represent specific sequences and, possibly, specific functions universally distinguishing the clade within the full microbial phylogeny (Figure 2).

\textbf{MetaRef families consist of the union of all pan, core, crown and marker families that determine the full microbial gene repertoire. MetaRef families can in the largest cases contain hundreds or thousands of genes, in which case it is convenient to use a unique representative sequence for the whole family, referred to as the family centroid (as for the pan families introduced above). Singleton families consisting of only one gene cannot fall into any of these categories and are excluded from core, crown and marker family definitions. Overall, MetaRef comprises \~5M families, 3.6M of which are core families and \~1M are markers as reported in Table 1 together with the statistics of some representative clades.}

\textbf{Organization of the MetaRef interactive web resource}

The MetaRef website provides two basic routes for data browsing: clade-centric and gene family-centric. The clade-centric view presents data based on the...
hierarchically structured taxonomy clades. The gene family-centric view provides biological information of individual MetaRef gene families in great detail. We recognize the intrinsic limitation of online low-throughput data navigation; therefore, throughout the site, the underlying data reported on screen can always be downloaded in plain text and FASTA format for further offline analysis.

The clade-centric view is organized according to the hierarchical (parent–child) relationships between clades and is rooted at two domain levels (Bacteria and Archaea) for which MetaRef currently catalogues 2706 and 112 genomes, respectively. The clade-centric webpage for a given query clade (example in Supplementary Figure S1) indicates the full taxonomic lineage of that clade and links to pages describing all related clades. When the query clade is at the family, genus or species level, a summary table of the core/pan genes of all sequenced genomes in the clade is reported. Links provided from this table connect to the gene family-centric view for further gene-specific inquiry.

To indicate how much sequencing data are currently available in the query clade, the total number of genomes in the clade and its direct child clades are reported and hyperlinked to genome pages. A circular genome tree of the query clade is provided to help visualize the number and distribution of genomes (the leaf nodes) sequenced in the clade and its direct child clades. When the query clade is itself a genome, more information is reported, including a table that summarizes the number of genes in that genome that are core at which ancestor clade and how many genes are unique markers of the genome. Additionally, a download link gives access to a detailed tab-delimited table describing all genes in the genome. The original external source of the genome is also provided for more inquiry.

The family-centric view (Figure 3) focuses on biological information for each individual MetaRef gene family. Information reported on the gene family page includes functional annotations aggregated from individual gene members and external functional databases; summary

Table 1. MetaRef family statistics for representative phyla, genera and species

<table>
<thead>
<tr>
<th>Species</th>
<th>Genomes</th>
<th>Genes</th>
<th>MetaRef families</th>
<th>Tot. Pan families</th>
<th>Tot. core families</th>
<th>Tot. marker families</th>
</tr>
</thead>
<tbody>
<tr>
<td>All microbes</td>
<td>1184</td>
<td>2818</td>
<td>10 880 874</td>
<td>5 006 295</td>
<td>3 600 814</td>
<td>1 028 534</td>
</tr>
<tr>
<td>Proteobacteria</td>
<td>463</td>
<td>1162</td>
<td>4 560 201</td>
<td>1 802 529</td>
<td>1 078 082</td>
<td>315 490</td>
</tr>
<tr>
<td>Bacteroidetes</td>
<td>88</td>
<td>124</td>
<td>474 420</td>
<td>294 243</td>
<td>23 924</td>
<td>48 040</td>
</tr>
<tr>
<td>Actinobacteria</td>
<td>148</td>
<td>272</td>
<td>999 822</td>
<td>582 201</td>
<td>38 028</td>
<td>43 935</td>
</tr>
<tr>
<td>Firmicutes</td>
<td>302</td>
<td>838</td>
<td>2 600 128</td>
<td>890 666</td>
<td>211 005</td>
<td>452 615</td>
</tr>
<tr>
<td>Staphylococcus</td>
<td>11</td>
<td>92</td>
<td>248 936</td>
<td>26 206</td>
<td>16 204</td>
<td>91 25</td>
</tr>
<tr>
<td>Strep. pneumonia</td>
<td>42</td>
<td>93 431</td>
<td>4341</td>
<td>2733</td>
<td>1579</td>
<td>29</td>
</tr>
<tr>
<td>Staph. aureus</td>
<td>72</td>
<td>198 655</td>
<td>5301</td>
<td>3137</td>
<td>1973</td>
<td>191</td>
</tr>
</tbody>
</table>

Singleton families are not reported.
plots of the prevalence and abundance of the gene as found at the major HMP body sites; and reports on homologs of the query gene family found outside of its clade. Access to taxonomic clades and MetaRef families is also possible through keyword search (see Supplementary Methods).

Consensus functional annotation of MetaRef families

MetaRef gene families were defined solely using sequence homology with stringent criteria (80% full-length identity, see Supplementary Methods). The genes clustered into a family were therefore likely to be genes that would carry out the same biological functions. However, we observed that functional annotation of individual members within a family were, at times, inconsistent, often stemming from the fact that individual genomes were annotated by different methods at different times, with varying criteria and terminology. Consolidating annotation information from individual gene members within a family helps improve consistency of annotation interpretation across genes and provides putative annotation corrections in many cases. Additionally, MetaRef directly links a gene family with corresponding taxonomic clades, which is crucial when focusing on the property of a function encoded by a gene restricted to a specific group of microbes sharing a common ancestor (e.g. studying a given gene family in *E. coli* rather than on all microbes possessing the gene).

The current MetaRef implementation provides a simple and conservative effort towards consensus annotation by tallying identical annotations from all available annotation schemes, including free-text deflines, KO, COG and EC assignments. Each family’s resulting table is dynamically and automatically generated from the consensus of its gene members. By reviewing the tally table, functions of some unknown/hypothetical protein genes have become clearer and recognizable. For example, a 71 gene core family of *S. aureus* (reported in Supplementary Figure S2) is consistently annotated as a putative siderophore biosynthesis protein subunit, although eight individual gene members were previously only annotated as hypothetical proteins. In another case (Supplementary Figure S2), two DNA mismatch repair proteins in a core family are labelled as HexA, whereas all the other members of the family (70) are labelled as MutS, suggesting a
misannotation. This potential inconsistency has been confirmed by ad hoc sequence-based analysis, which showed that both of the former two proteins possess the MutS_1, MutS_2 and MutS_3 superfamilies and the ABC MutS1 domains and that they differ from several consistently annotated MutS proteins by only one peptide along the full-length of 872 peptides. These examples confirm the usefulness of MetaRef efforts towards large-scale sequence-based consensus annotation.

MetaRef characterization of metagenomic samples

Comprehensive studies of human-associated microbial communities are still in their first generation and have yet to be well-integrated with microbial isolate sequence resources such as the Human Microbiome Project's (HMP) Data Analysis and Coordination Center (21) (http://hmpdacc.org). MetaRef includes what we believe to be the first comprehensive gene family-centric reference generated from a large cohort of healthy human subjects: using our pre-computed database of family centroids along with the metagenomes sequenced by the HMP, we have compiled baseline data characterizing the relative abundance and prevalence of all gene families across all major body sites in healthy individuals. Investigators studying site-specific pathologies, for example, can use this data to efficiently characterize differences in functional potential (at the individual gene level) between healthy- and disease-associated microbial communities (22) by looking for significant differences in observed prevalence/abundance of detected genes (23). It has been shown (11) that marker centroids can be reliably used to estimate the presence and relative abundance of their associated clades. As a result, a single alignment pass of a collection of pathology-associated metagenomic sequence reads against the full collection of MetaRef centroids—which is then compared to analogous data compiled from a cohort of healthy subjects at the same body site—can be interrogated to detect both taxonomic and functional differences between health- and disease-associated microbiota.

To construct our baseline reference data, we first used Bowtie (17) to align all sequence reads from 757 shotgun metagenomic HMP samples—covering all five studied body areas (airways, GI tract, mouth, skin and urogenital tract, including their various subsites)—against 5 006 294 family centroid sequences (Figure 4). For each sample, we counted all matches to each centroid, then converted these raw counts to RPKMs (24) to normalize with respect both to differences in inter-sample sequencing depth and differences in individual centroid lengths. Using these normalized counts, we calculated, for each sample, the relative abundance of each centroid as a percentage of all normalized counts. For all body sites, we computed the mean relative abundance (over all non-zero samples) of each centroid detected at that site, to generate a single statistic estimating the typical relative abundance of each centroid at that site. The presence or absence of each centroid at each site was also used to compute prevalence statistics: for each centroid and each body site in which it was found, we report the percentage of samples in which that centroid was detected. Body-site-based relative abundance and prevalence statistics are summarized (as box plots and bar plots, respectively) on individual centroid pages on the MetaRef website (cf. Figure 3). Although for completeness of the downloadable resource we aligned all 757 HMP samples against the centroid collection, the prevalence and abundance data provided by MetaRef reflects the subset of 630 samples that passed HMP’s quality-control procedures for the six body sites considered. Please see the Supplementary Methods for a discussion of our alignment strategy, precise details of the alignment operations, and a summary of the results of validation tests we conducted to ensure that our results were sufficiently representative of the sampled microbial communities.

The number of centroids detected in each sample that passed QC ranged from 2563 (from a sample taken from the posterior vaginal fornix: when healthy, such samples exhibit the lowest-diversity microbe communities of all studied body sites (25)) to 4128821 (from a distal gut sample, the most complex of all human-associated microbial communities (25)), with a median of 94 135 (from an oral sample). As examples, one core gene (encoding a sigma factor subunit of RNA polymerase) for Bacteroides ovatus—a bacterium commonly found in the human gut (26,27)—was observed in 81.53% of all GI-tract samples, and in <5% of samples sequenced from all other body sites. Its average relative abundance within gut samples was more than two orders of magnitude higher than its relative abundance in all other body sites. Similarly, a core gene (encoding a cell division protein) for S. epidermidis—a common commensal colonist of the human nasal passage (28,29)—was seen in ~25% of all nasal samples, but <3% of samples from other body sites; its average relative abundance in nasal samples was again at least two orders of magnitude greater than in samples from any other body site. Comprehensive prevalence/abundance statistics for all family centroids (including markers) have been made available for direct download from the MetaRef website, and graphical reports for individual centroids can be browsed directly (Supplementary Figure S3). These can be conveniently used as a robust collection of healthy baseline data for any research involving the human microbiome, whether the focus be on taxonomic composition, functional potential or both.

DISCUSSION

With MetaRef.org, we provide the most comprehensive and up-to-date database of a non-redundant reference gene catalogue that not only delivers information on gene family conservation, but also captures phylogeny and consensus functional annotation. It is easily accessible as an online browsable resource allowing quick functional and phylogenetic explorations of gene families, and its contents are also available by direct download for large-scale bioinformatics applications. MetaRef is committed to providing a comprehensive microbial gene family resource, and thus includes all systematically
available finished and draft genomes from the time of creation, automatic detection of taxonomic inconsistencies and suggestions of plausible corrections, and is designed with regular automated updates at increasing scale in mind. We anticipate future versions including additional sources of microbial genomes and expanding rapidly as isolate sequencing becomes increasingly ubiquitous.

With microbial sequence data in all its diversity being strongly on the rise (30–32), a new approach such as MetaRef is needed to simultaneously organize information on gene function, phylogeny and microbial taxonomy throughout the tree of life to facilitate downstream biological interpretation. Going forward, we also plan to increase the use of and connectivity to other genome databases, particularly those providing extended annotation schemes in order to continue the harmonization process for annotations. We will also automatically incorporate additional phylogenetic lineages currently being sequenced (32,33) as well as additional meta’omic datasets (34–37), and based on current performance we expect the system to scale smoothly in anticipation of the rapid increase of available genomes.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online, including [38–40].

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