New insights from 33,813 publicly available metagenome-assembled-genomes (MAGs) assembled from the rumen microbiome

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Abstract
Recent rumen microbiome metagenomics papers¹,² have published hundreds of metagenome-assembled genomes (MAGs), comparing them to 4,941 MAGs published by Stewart et al³ in order to define novelty. However, there are many more publicly available MAGs from ruminants. In this paper, for the first time, all available resources are combined, catalogued and de-replicated to define putative species-level bins. As well as providing new insights into the constitution of the rumen microbiome, including an updated estimate of the number of microbial species in the rumen, this work demonstrates that a lack of community-adopted standards for the release and annotation of MAGs hinders progress in microbial ecology and metagenomics.

Background
Recent advances in metagenomic assembly and binning have given rise to very large collections of metagenome-assembled genomes (MAGs), which often represent the only genomic information about a microbial strain or species that has not yet been cultured. These MAGs provide essential insight into the functional potential of individual strains and species, as well as the microbiomes they inhabit.

As globally important food-producing animals, ruminants are the subject of intense research, particularly as the microbiome in the rumen is primarily responsible for the breakdown of recalcitrant plant material into nutrients that the host can absorb. Peng et al¹ and Gharechahi et al² both recently published hundreds of rumen-derived MAGs, and compared them to 4941 rumen MAGs from Stewart et al³, and a previously published culture collection⁴. However, many more rumen-derived MAGs are publicly available, including 15 MAGs released by Hess et al⁵, 79 from Solden et al⁶, 99 from Svartstrom et al⁷, 251 from Parks et al⁸ (which analysed rumen data from Wallace et al⁹ and Shi et al¹⁰), 1200 from Wilkinson et al¹¹, 4199 released by Glendinning et al¹² (391 of which were the subject of the publication), and 20469 released by Stewart et al³ (4941 of which were the subject of the publication).

The 719 MAGs from Peng et al and the 538 MAGs from Gharechahi et al were compared against the full set of existing 32,557 rumen MAGs that were publicly available at the time of publication, in addition to 460 cultured genomes from the Hungate collection. Using permissive settings designed to reduce the number of false-positive species-level bins, the entire set of rumen microbial genomes was de-replicated to produce a set of putative microbial species-level bins. In addition to providing new taxonomic insights into the rumen microbiome, this work also demonstrates that the lack of a single, community-adopted repository for metagenomic bins and MAGs; the use of non-INSDC data repositories; a lack of community-adopted standards for MAG annotation; and a lack of enforcement of standards by journals, are all barriers to future research in metagenomics.
Methods

Data collection
MAGs from Peng et al, Gharechahi et al, Stewart et al, Stewart et al, Wilkinson et al, Parks et al and Glendinning et al were downloaded from the ENA or NCBI SRA databases using BioProject identifiers listed in the relevant publication. 460 isolate genomes were downloaded from Seshadri et al (those which were present in INSDC databases at the time of download). MAGs from Hess et al were downloaded from the NERSC portal. MAGs from Solden et al were provided by the authors via Google Drive. MAGs from Svartstrom et al were created by parsing the FASTA headers of metagenome assemblies released under the BioProject accession listed in the publication.

Stewart et al and Glendinning et al are noteworthy as they released large numbers of assembly bins (20,469 and 4,199 respectively) that were not described as part of the paper, instead choosing to select high quality MAGs from those bins for further analysis and description. However, those datasets are still useful and were included in the analysis.

Completeness and contamination scores for Stewart et al, Stewart et al, and Glendinning et al were retrieved from the ENA using the BioProject identifiers listed in the relevant publication. Completeness and contamination scores for Peng et al and Gharechahi et al were retrieved from the publication and linked to the published sequences using assembly statistics. Completeness and contamination scores for Seshadri et al, Solden et al, Svartstrom et al, Hess et al and Parks et al were calculated using CheckM. Finally, completeness and contamination scores for Wilkinson et al were retrieved from the authors.

Dereplication
The software dRep was used to de-replicate 33,813 publicly available FASTA files representing MAGs and isolate genomes from ruminants. The parameters used were:

- --S_algorithm fastANI
- --multiround_primary_clustering
- -comp 50
- -con 10
- -sa 0.95
- -nc 0.3

These settings delineate species-level bins if genomes have lower than 95% ANI across 30% of their length. The low length parameter (30%) recognizes that incomplete genomes may not overlap along a large proportion of their shared sequence, and using a higher value may split genomes which belong to the same species and over-estimate the number of species. A 30% coverage threshold has been used in previous large-scale MAG studies which involved 50% complete genomes.

Taxonomic classification and phylogeny
MAGpy and GTDB-Tk (with the “classify_wf” option) were used to assign a taxonomy to the “winning” MAGs from dRep. To create a representative phylogenetic tree, genomes that were at least 90% complete and less than 5% contaminated were used as input to PhyloPhlAn. Proteins were predicted for the genomes with Prodigal and PhyloPhlAn was run with options:
-d phylophlan
-t a
-f supermatrix_aa.cfg
--diversity low
--fast
--verbose
--maas phylophlan_substitution_models/phylophlan.tsv
--remove_fragmentary_entries
--fragmentary_threshold 0.85

A plot of the phylogenetic tree was created using GraphlAn\textsuperscript{21} with genomes coloured by the Phylum assigned by GTDB-Tk.

\textbf{Figure 1.} Phylogenetic tree of 2,696 species level bins chosen for high completeness (>90%) and low contamination (<5%). The tree was created using PhyloPhlAn and drawn using GraPhlAn, with taxonomic assignments from GTDB-Tk.

\textbf{Results}

After de-replication using the parameters above, there remained 7,533 putative species-level bins. Treating all data released by a single publication as a single dataset (n=10), 4,794 were singletons,
representing putative microbial species that are represented in only one dataset. Conversely, 2,739 putative microbial species were seen in more than one dataset. There were no species-level bins present in all datasets. The maximum number of datasets any species-level bins were present in was 7, and only three bins were present in these 7 datasets: two from Stewart et al, and one from Gharechahi et al. Results of the de-replication process are summarized in supplementary data 1.

Of the “winners”, 4482 came from Stewart et al, 728 from Wilkinson et al, 694 from Peng et al, 660 from Glendinning et al, 358 from Gharechahi et al, 313 from Seshadri et al, 149 from Parks et al, 78 from Svartstrom et al, 62 from Solden et al and 9 from Hess et al.

Using the Chao 1 index, we update the estimate of the number of microbial species in the rumen (previously estimated by Stewart et al) to 13,616.

Of the 7,533 species-level bins, GTDB-Tk identified 155 as Archaea. The majority of these were assigned to the Phylum Methanobacteria (119; 77%), with 31 being assigned to Thermoplasmata and 5 to Halobacteria. GTDB-Tk was unable to assign a Genus to eight (5%) of the Archaeal MAGs, and was unable to assign a species to 110 (71%). GTDB-Tk assigned 7,378 species-level bins to the Bacteria domain. These spanned 28 different Phyla, the most popular being Firmicutes_A (3339; 45%), Bacteroidota (1671; 23%), Firmicutes (807; 11%), Proteobacteria (299; 4%) and Verrucomicrobiota (248; 3%). Of particular interest as high fibre-degrading microbes, the dataset also contains 63 members assigned to the phylum Fibrobacterota. Of the 7,378 Bacterial species-level bins, GTDB-Tk was unable to assign a Class to one, unable to assign a Family to 99, unable to assign a Genus to 1,087 (15%) and unable to assign a Species to 5,796 (78%). The full results of the GTDB-Tk taxonomic assignments can be found in supplementary data 2.

A phylogenetic tree of the 2,696 highly complete MAGs can be seen in Figure 1. The tree is dominated by large clades of Firmicutes_A and Bacteroidota, though other significant clades exist with the MAGs spread across 26 different phyla.

Conclusions and discussion
Assembly of genomes from metagenomic sequencing is becoming routine, with new MAG datasets published frequently. In order for comparisons to be made between new and existing MAG catalogues, it should be easy to find and retrieve all MAGs from a particular biome, alongside metadata in a standardised format. The MIMAG standard attempts to define standards for metadata around MAGs, but these have not been universally adopted, and cannot be applied retrospectively to historical datasets. MAGs should be deposited in INSDC databases alongside metadata adhering to the MIMAG standard, and with completeness and contamination estimates as a minimum. In addition, it would be beneficial to submit all of the following: the raw reads; the raw metagenome assemblies; all metagenome bins created during the binning process; the final set of metagenome-assembled genomes. The European Nucleotide archive (ENA) provides a suitable repository and guidance for submitting all of these; and have metadata checklists for both binned genomes and metagenome-assembled genomes that allow for the storage of essential metadata needed to interpret and compare metagenomics data. EBI’s MGnify also provides added value services on these datasets.

There are approximately 4 billion food-producing ruminants on the planet at any one time (FAOSTAT), and as such, the rumen and its microbiome are a priority area for research globally.
Recent advances in metagenomics mean we can now study the structure and function of the rumen microbiome in unprecedented detail. Here, for the first time, all data resources representing binned metagenomes from ruminants were combined and de-replicated to produce 7,533 putative species-level bins. However, combining and de-replicating partial and contaminated genomes is an inexact science, and it is difficult to effectively delineate species given the amount of missing information. Relatively permissive parameters were chosen so as to avoid over-inflation of the number of species-level bins. However it is still possible that this is an over-estimate due to the inclusion of incomplete genomes.

The fact that the majority of the species-level bins for ruminants are singletons suggests that we are yet to sample the entire species-level sequencing space of global ruminants, and the lack of a species-level core microbiota suggests a large amount of variation in the constitution of the rumen microbiome. We update the estimate of the total number of microbial species in the rumen microbiome to 13,616 and provide taxonomic labels for all known species to date, which span 26 different microbial phyla. It is essential that researchers are funded to culture this microbial diversity, and study the role of these species in ruminant productivity and health, in climate change and sustainability, and in global food security.

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