The evolution of microbial species – a view through the genomic lens

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ABSTRACT

For a long time prokaryotic species definition has been under debate and a constant source of turmoil in microbiology. This has recently prompted the ASM to call for a scalable and reproducible technique, which uses meaningful commonalities to cluster microorganisms into groups corresponding to prokaryotic species. Whole-genome Average Nucleotide Identity (gANI) was previously suggested as a measure of genetic distance that generally agrees with prokaryotic species assignments based on the accepted best practices (DNA-DNA hybridization and 16S rDNA similarity). In this work, we prove that gANI is indeed the meaningful commonality based on which microorganisms can be grouped into the aforementioned clusters. By analyzing 1.76 million pairs of genomes we find that identification of the closest relatives of an organism via gANI is precise, scalable, reproducible, and reflects the evolutionary dynamics of microbes. We model the previously unexplored statistical properties of gANI using 6,000 microbial genomes and apply species-specific gANI cutoffs to reveal anomalies in the current taxonomic species definitions for almost 50% of the species with multiple genome sequences. We also provide evidence of speciation events and genetic continuums in 17.8% of those species. We consider disagreements between gANI-based groupings and "named" species and demonstrate that the former have all the desired features to serve as the much-needed "natural groups" for moving forward with taxonomy. Further, the groupings identified are presented in detail at http://ani.igi-psf.org to facilitate comprehensive downstream analysis for researchers across different disciplines

OBJECTIVES

Polyphasic

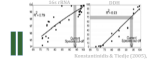
tedious +

Microbial

Whole-genome sequencing (WGS) launched microbial taxonomy into the new era of genomic microbial taxonomy. with the possibility of establishing systematics on the basis of information retrieved from complete genomes.

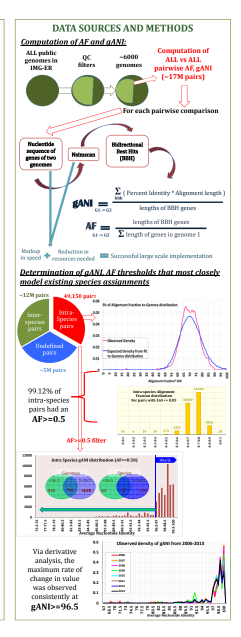
It is plausible to consider that microbial taxonomy will be steadily more dependent on genome sequences than relying on the classic polyphasic taxonomy. including phenotypic characterization using time-consuming laborious laboratory tests.

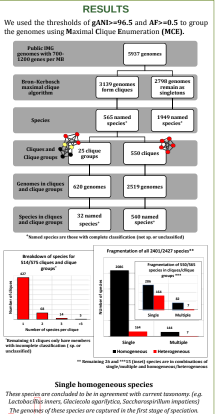
ALIGNMENT FRACTION (AF) The fraction of genes between two genomes AVERAGE NUCLEOTIDE IDENTITY (gANI) Sequence level identity across all the conserved genes



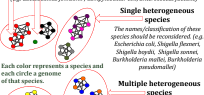
reflects the degree of evolutionary distance

In this work, we propose whole-genome Average Nucleotide Identity (gANI) together with Alignment Fraction (AF) as a robust and reliable method for grouping of microbial genomes towards the goal of species classification.





(e.g. Lactobacillus johnsonii , Campylobacter coli, Enterococcus faecalis)



Multiple homogeneous species

Such species are considered to be captured in the second stage of speciation (e.g. Bacillus amyloliquefaciens, Bacillus subtilis, Citrobacter freundii, Enterococcus casseliflavus, Enterococcus faecium, Lactobacillus jensenii)

The names/classification of these

species should be reconsidered (e.a.

Bacillus mycoides, Salinispora

arenicola. Rickettsia rickettsii)

CONCLUSIONS

Several reports have already illustrated that microbial taxonomic assignments are inconsistent with emerging genetic. systematic, and phenotypic information for a large number of species. According to a recent ASM report, "in moving forward with microbial taxonomy, it is critical to determine whether microorganisms cluster in groups with meaningful commonalities or to determine what commonalities may be best used to cluster microorganisms into meaninaful groups".

For the first time, gANI was applied across all available sequenced prokaryotic genomes and its potential to cluster microorganisms into such "meaningful groups" was explored. We demonstrate that gANI, which maximally utilizes the commonalities between microbial genomes, is a robust measure of genetic relatedness for establishing accurate evolutionary relationships. The gANI-based cliques were validated by comparisons with "named" species, similarity of 16S rDNA, and similarity of conserved core pMGs. They were then used to address central questions such as whether microorganisms form a continuum of genetic diversity, or distinct species represented by distinct genetic signatures. Thus gANI-based cliques not only provide insights into the evolutionary dynamics of prokaryotes, but also significantly assist in the refinement of the current taxonomy.

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