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SymbAphidBase: a new database dedicated to aphid symbionts to store novel sequenced genomes and standardize their annotations.

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Complete sequences of bacterial genomes are accumulating with an unprecedented speed due to the democratization of NGS technologies. This is also true for symbiotic bacteria and genomic comparisons are key to understanding their contribution to host biology. We developed SymbAphidBase [1]: an ad hoc genome database to store and analyse aphid symbionts’ genome sequences. Aphids harbour an obligate primary endosymbiont, Buchnera aphidicola, and several facultative secondary symbionts. SymbAphidBase is designed to integrate data from all these bacteria. At present it includes the sequenced genomes of 17 strains of B. aphidicola from 8 different aphid species available in GenBank. To implement this database we used the GMOD’s tools: the chado database to store the genomic data and annotations, coupled with the JBrowse genome browser. SymbAphidBase includes an interface that gives access to data in different formats: a genome browser, a Blast server, comparative genes/proteins statistics and downloadable files. From the beginning of the project, the need to generate a unified gene annotation and identification scheme was apparent. In fact, if we were to use the original gene functional annotations and names, often a small fraction of genes would be found to be common in the different B. aphidicola genomes when performing pairwise comparisons (as low as 10%).

In light of these results, we decided to re-annotate the genomes using EuGene-P, a prokaryotic gene finder tool. The genes are later re-annotated (or annotated for the new genome sequences) using a Blastx analysis against the HAMAP [2] protein database that includes 10 highly curated B. aphidicola genomes. The final assignment of gene names is prioritized in a filtered pipeline to include the SwissProt or TrEMBL IDs when available with variable homology criteria (that are registered in the new gene ID). With this approach we are able to increase the number of common genes when performing pairwise comparisons among B. aphidicola genomes (40-99% with our method depending on the chosen parameters). For genes that do not get a name and functional annotation using this automated method, we are working on other approaches that would use phylogeny and/or expert manual annotation. Beyond this novel unified gene annotation, to facilitate the direct comparison of different genomes, we implemented a double browser interface to facilitate the contemporary visualization of two genomes at the same time. All these database generation steps are automated with specific pipelines developed using mainly Perl, PHP, and jQuery languages. In conclusion, SymbAphidBase is a companion database to AphidBase [3] (the aphid genome database) to facilitate genomic data storage and analysis to study symbiosis in the aphid model.


Keywords: genome database; genome annotation; aphid symbionts; SymbAphidBase