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Recent Work

Title
Genome Annotation: Going Beyond Human Genome

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Three human chromosomes [1-3] and a variety of eukaryotic genomes [4-9], dozens of microbial genomes and environmental communities [10-12] are being sequenced, assembled and annotated at the DOE Joint Genome Institute (http://genome.jgi-psf.org). Differences in genome organization and gene structure as well as variation in the data available for training and validation of gene predictions require custom approaches for each genome and create additional challenges for annotation. To address these challenges we have developed JGI annotation pipeline that allows us to annotate these genomes in high-throughput manner but also take into account specifics of individual genomes. The pipeline integrates a suite gene prediction and annotation methods, custom methods for validation and consolidation of various gene models, and visualization tools for genome-scale analysis. Analysis of genomes annotated using the Annotation Pipeline compared to the earlier genome releases, predictions made by other groups, and tests on data not used for gene prediction training indicate overall improvement in quality of predicted gene models.


