Genome analysis

MetaPathways v2.5: quantitative functional, taxonomic and usability improvements

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Abstract

Summary: Next-generation sequencing is producing vast amounts of sequence information from natural and engineered ecosystems. Although this data deluge has an enormous potential to transform our lives, knowledge creation and translation need software applications that scale with increasing data processing and analysis requirements. Here, we present improvements to MetaPathways, an annotation and analysis pipeline for environmental sequence information that expedites this transformation. We specifically address pathway prediction hazards through integration of a weighted taxonomic distance and enable quantitative comparison of assembled annotations through a normalized read-mapping measure. Additionally, we improve LAST homology searches through BLAST-equivalent E-values and output formats that are natively compatible with prevailing software applications. Finally, an updated graphical user interface allows for keyword annotation query and projection onto user-defined functional gene hierarchies, including the Carbohydrate-Active Enzyme database.

Availability and implementation: MetaPathways v2.5 is available on GitHub: http://github.com/hallamlab/metapathways2.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Since the publication of MetaPathways (Konwar et al., 2013), a modular annotation and analysis pipeline that enables construction of environmental pathway/genome databases using Pathway Tools (Karp et al., 2002b, 2010) and MetaCyc (Caspi et al., 2012; Karp et al., 2000, 2002a), there have been improvements to the software via the Knowledge Engine data structure, a graphical user interface (GUI) for data management and browsing and a master–worker model for task distribution on grids and clouds (Hanson et al., 2014b). Version 2.5 features faster and more accurate quantitative functional and taxonomic inference. Inspired by the pathway-centric analysis of the Hawaii-Ocean Time-series (Hanson et al., 2014a), a weighted taxonomic distance (WTD) has been integrated to detect taxonomic divergence of predicted MetaCyc pathways. Next, because it is difficult to determine relative open reading frame (ORF) abundance in assembled datasets, we adopt reads per kilobase per million mapped (RPKM) to provide a quantitative measure of sequence-coverage on a per-ORF basis (Patil et al., 2011). Additionally, the LAST code has been modified to calculate BLAST-equivalent Bit-score and E-value statistics (Altschul et al., 1990;
MetaPathways v2.5 now addresses quantitative functional and pathway prediction hazards based on WTD and RPKM calculations, providing performant LAST output format and statistics. We modified the LAST code to produce the compatible Bit-score and E-value calculations.

3 Results
We benchmarked the implemented improvements described earlier using Illumina-sequenced marine metagenomic samples. (Joint Genome Institute: ‘Marine microbial communities from Expanding Oxygen minimum zones project’ [JGI Project IDs: 4093112, 4093113, 4093125, 4093127–4093132, 4093144–4093149, 4096364–4096371, 4096373, 4096375, 4096377–4096379, 4096381–4096383, 4096385–4096387, 4096389–4096396, 4096398–4096406 and 4096409–4096453]). The WTD distribution can be used as an informative tool to place pathways into different taxonomic hazard classes based on their order statistics (Fig. 1a). Protein annotations of BLAST and LAST are highly correlated in terms of E-value (Fig. 1b), suggesting roughly equivalent results, but with LAST being significantly faster. Although there is a positive correlation between RPKM score and ORF count, variance about the regression line indicates RPKM makes a correction in many instances (Fig. 1c).

4 Conclusions
MetaPathways v2.5 now addresses quantitative functional and pathway prediction hazards based on WTD and RPKM calculations, provides performant LAST output equivalent with BLAST, and more flexible annotation subsetting and projection via GUI keyword searches. These improvements enable improved large-scale comparative analysis of next-generation environmental sequence information.
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Conflict of Interest: none declared.

References

Hanson, N.W. et al. (2014a). Metabolic pathways for the whole community. BMC Genomics, 15, 619.