

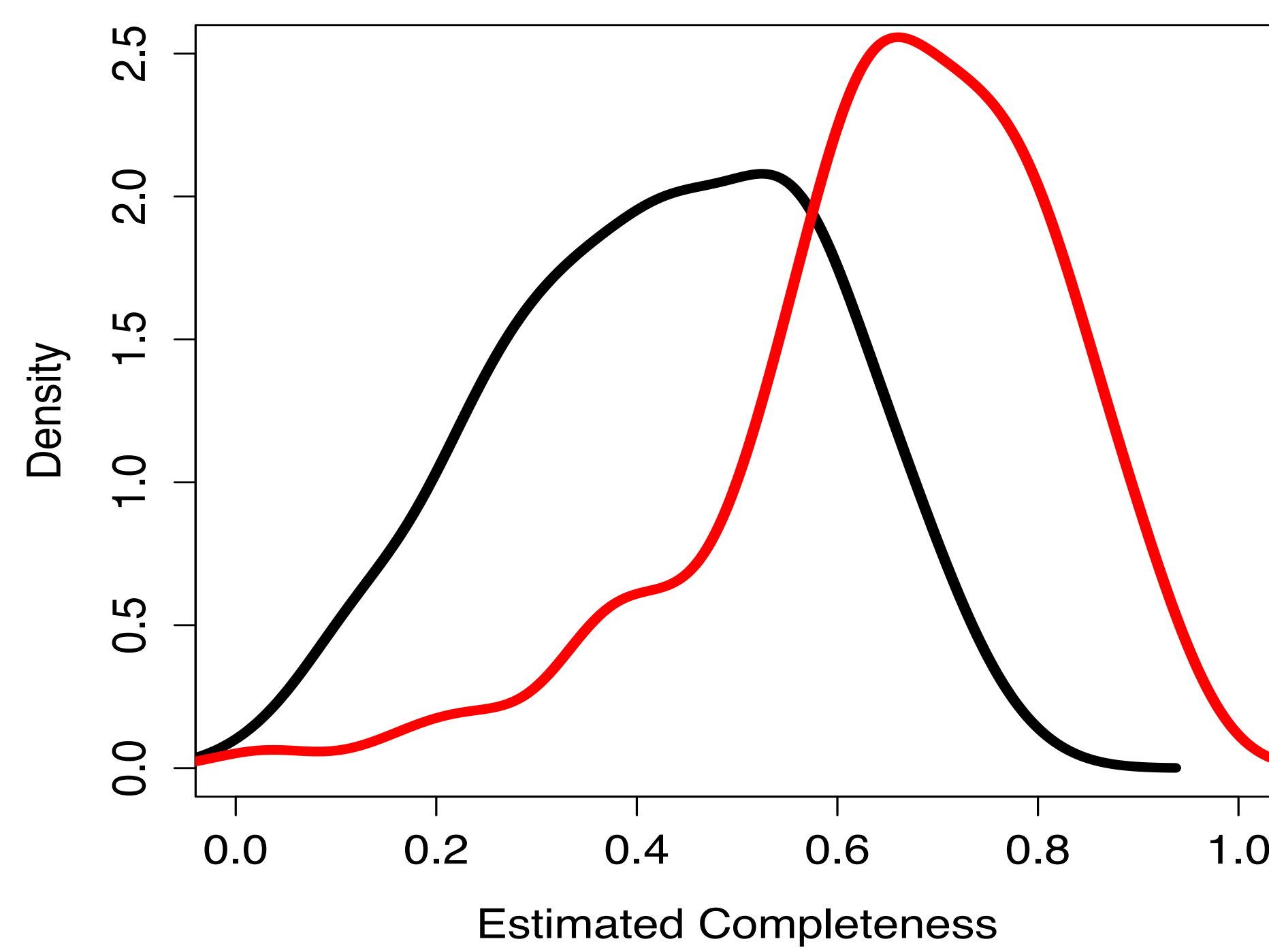
# SAGEX: The Single-cell Amplified Genome EXtrapolator

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## OVERVIEW

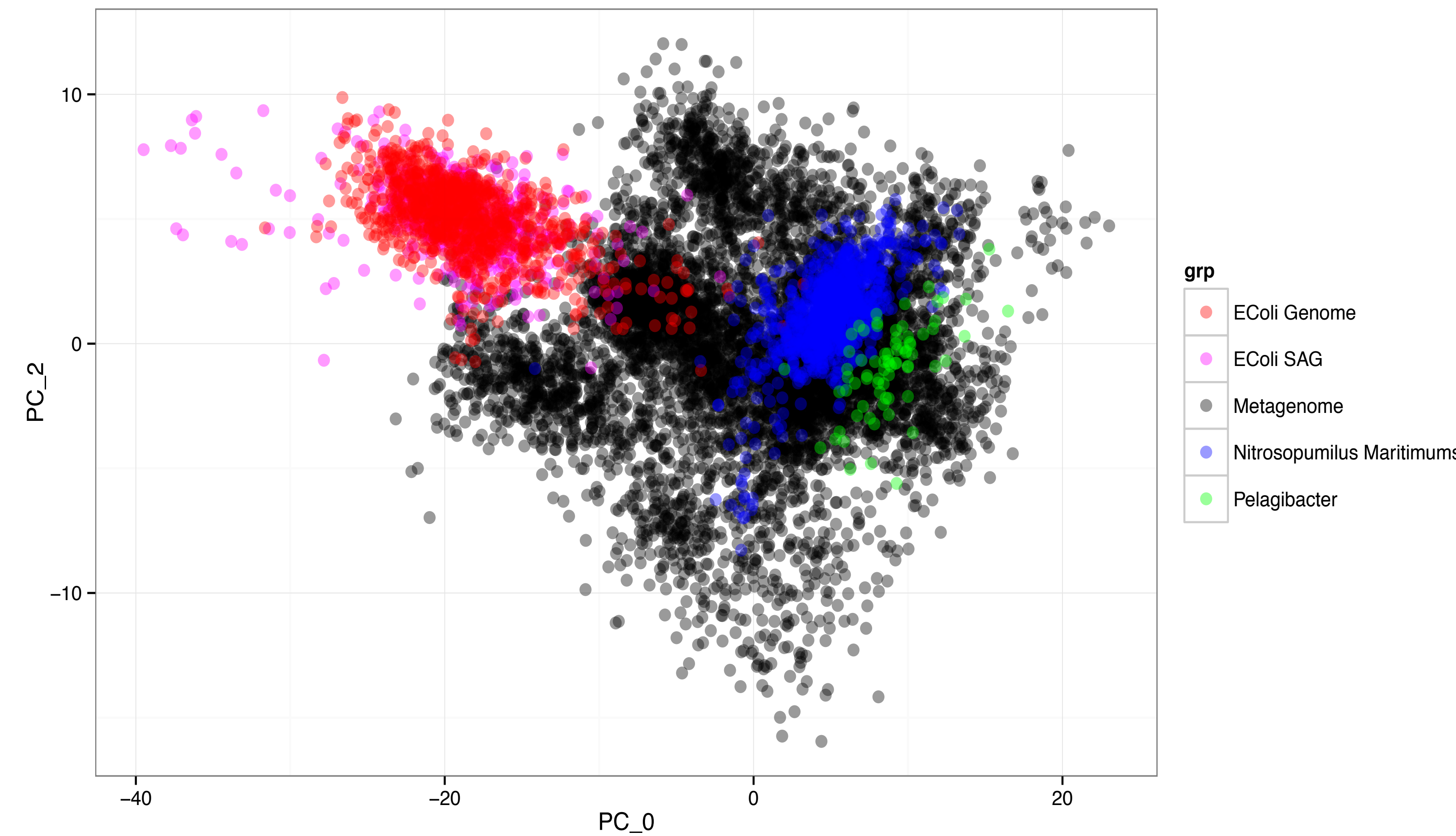
Single-cell amplified genome (SAG) sequencing promises to provide a powerful adjunct to shotgun sequencing of environmental DNA (e.g. metagenome) in determining uncultivated microbial community structure and function. Resulting SAG sequences can be unambiguously assigned to cognate taxa providing robust fragment recruitment platforms for determining gene frequency distribution patterns and population structure in natural and human engineered ecosystems. Unfortunately the multiple displacement amplification process used to generate SAG sequences results in substantial gaps in genome coverage. For example, by counting Clusters of Orthologous Groups (COGs) [1], we show that for a collection of 129 SAGs genome completeness ranges between 5-80%. Here, we propose a bioinformatic remedy to incompleteness inspired by [2], the single-cell amplified genome extrapolator (SAGEX) that leverages available SAG sequence information to extrapolate genome coverage using assembled metagenome sequences generating a population genome bin. SAGEX is capable of high Positive Predictive Value (PPV) while maintaining sensitivity making it an excellent predictor of related donor genotypes. SAGEX is implemented in C/C++ and requires no additional libraries beyond the compiler. The pipeline is fast (usually running in under 5 minutes) and can be used to optionally write kmer counts and a kmer PCA readable into R for visualization.



**FIGURE 1. SAGEX confidently extrapolates SAGs.**

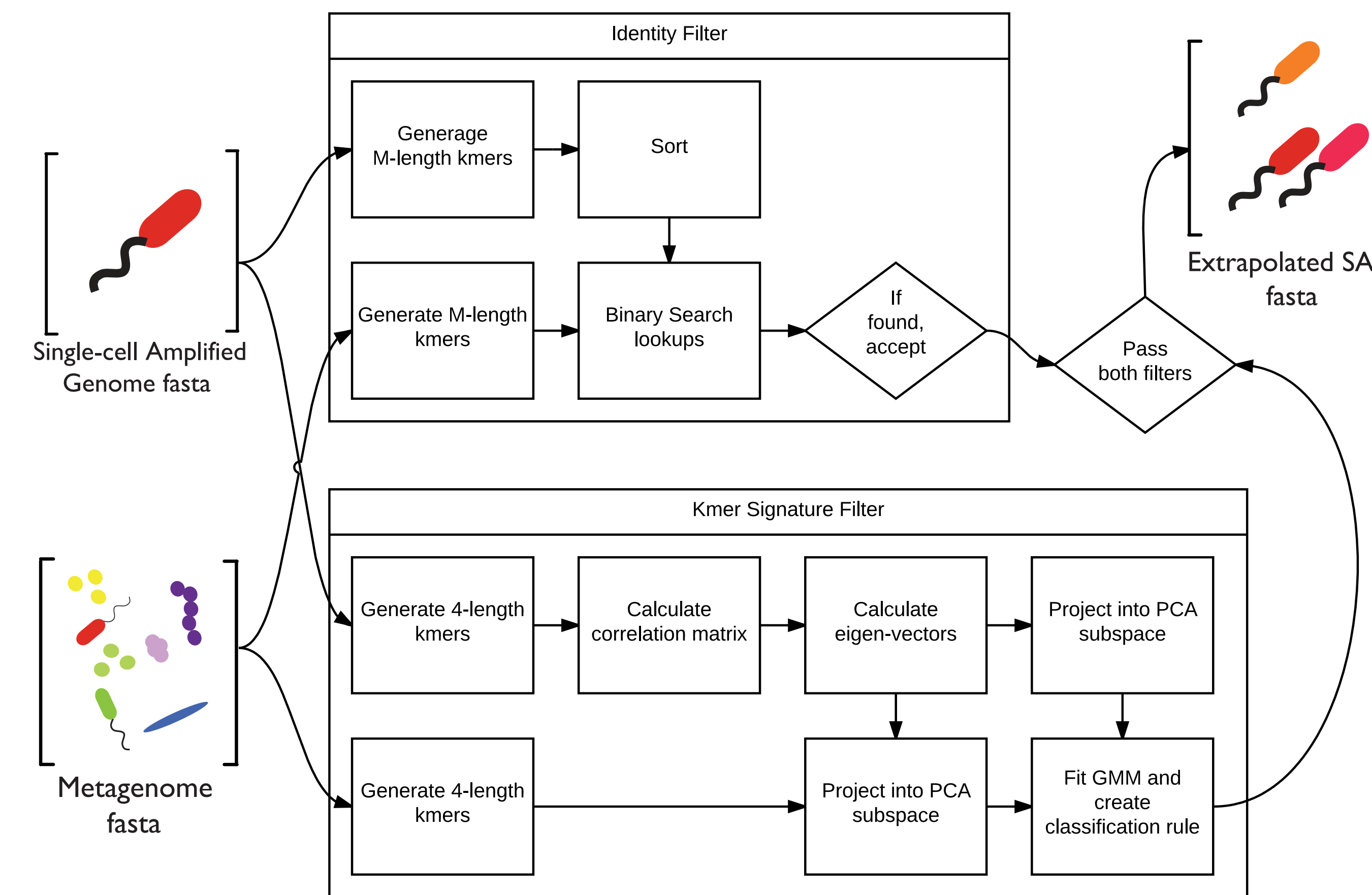
Prior to running SAGEX on 129 SUP05 SAGs sourced from the Saanich Inlet water column (black) [3], estimated completeness (by counting COGs) prior to extrapolation averages 40% (red). Using high PPV settings, SAGEX increases average completeness to 70%. Relaxing PPV settings can increase completeness of the extrapolated genome.

## HOW IT WORKS



**FIGURE 2. Genome's kmer signatures.**

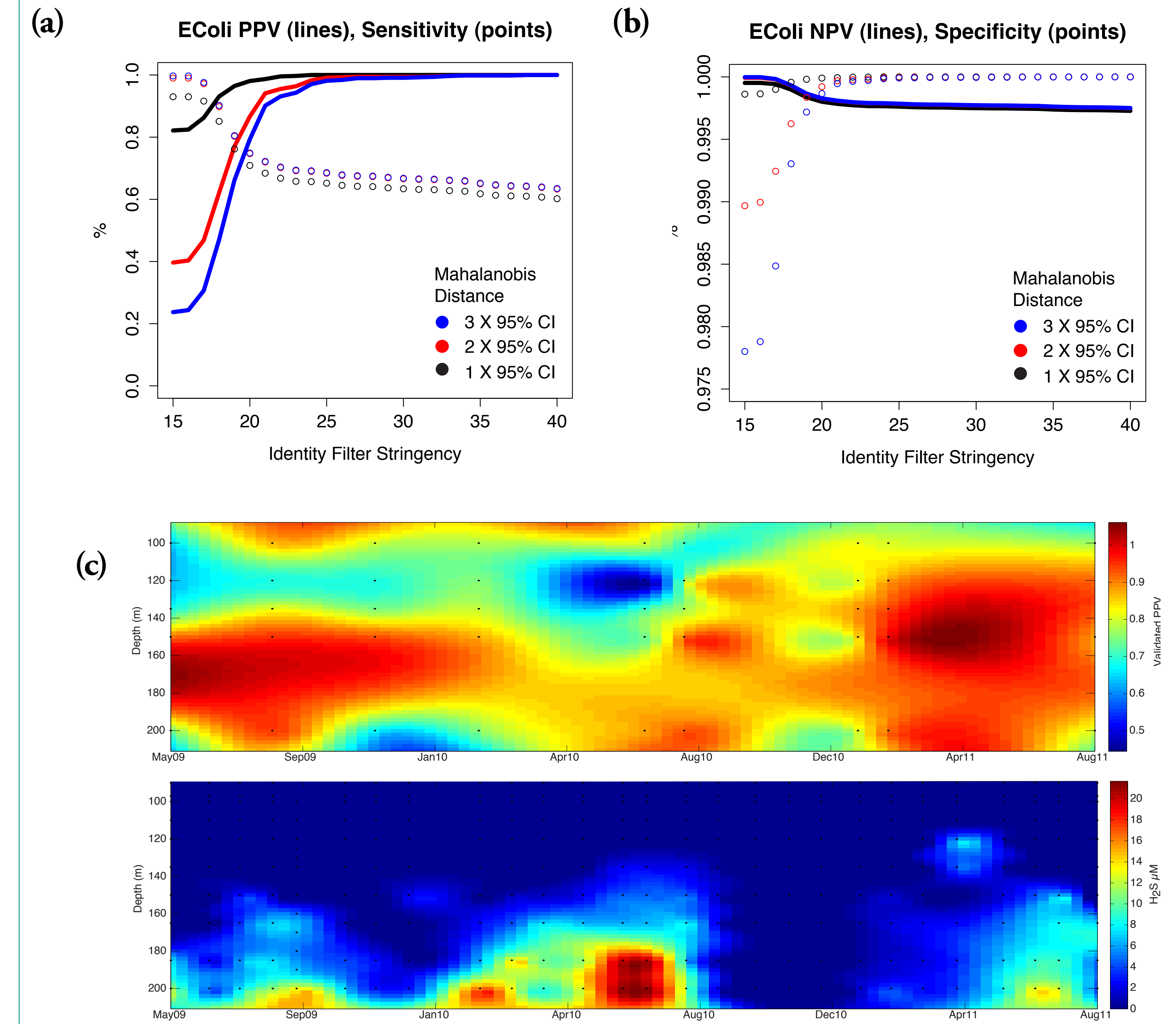
SAGEX requires contigs to pass two tests before calling them hits. First is an identity filter. Second is a kmer signature filter. Various genome kmer signatures are pictured above demonstrating the intuition behind the second test. Notice how the EColi's genome and SAG share very similar kmer signatures. The image is generated by calculating the tetranucleotide frequencies of each contig, converting counts to proportions, transforming all data to the principal component basis of the standardized metagenome, and plotting only two principal components.



**FIGURE 3. The SAGEX pipeline.**

SAGEX accepts two input FASTAs, an assembled SAG and assembled metagenome, and provides a single output FASTA, a subset of the metagenome contigs recruited by the SAG. The two tests which every hit must pass are expanded here. The identity filter test requires every hit to share at least one M-length substring with any SAG contig. The kmer signature filter requires each every hit to fall within the kmer signature region as estimated by a Gaussian Mixture Model in a kmer-proportion principal component subspace.

## ERROR ANALYSIS



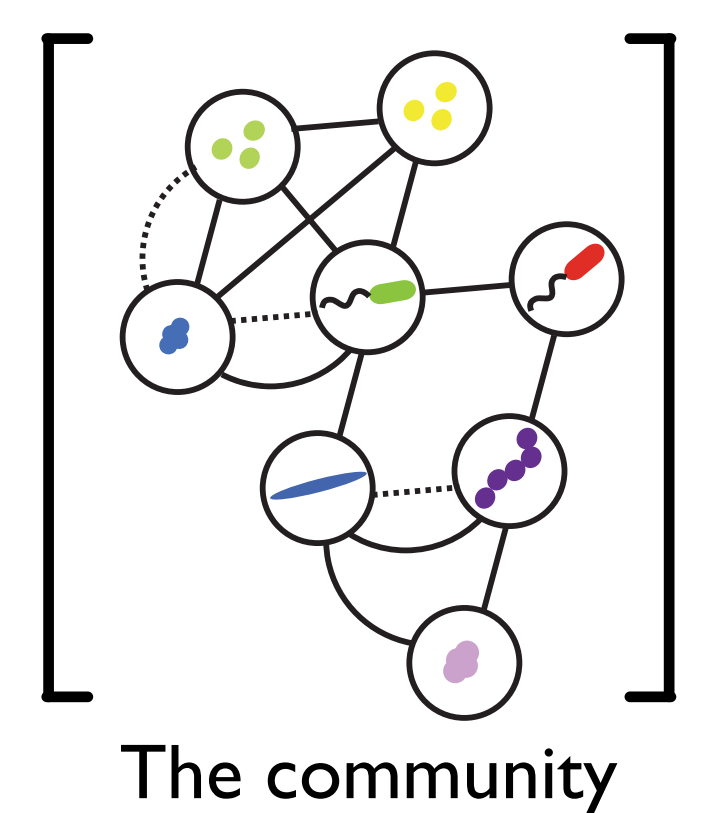
**FIGURE 4. Error analyses of SAGEX.**

(a,b) Classification errors were estimated for SAGEX using an E. Coli MG1655 SAG [4] as training data, a Saanich Inlet metagenome [5] as false hits, and an E. Coli MG1655 isolate genome [6] as true hits. E. Coli MG1655 was ideal because it is not expected to occur naturally in the metagenome, thus not confusing true and false hits. (c) SAGEX PPV was validated because E. Coli is too ideal a case. We conclude that SAGEX should be used with a SAG and metagenome from the same sample (or similar). To validate SAGEX PPV, the 129 SUP05 SAGs were run against 48 metagenomes at various depths and times. BLAST was used to query SAGEX identified ORFs against a database containing all SUP05 SAG sequences, requiring at least one successful hit for validation. (c,top), The average validated PPV per metagenome. Dots represent sample points in time and depth. Interpolation was used to create the contour plot in Matlab. (c,bottom), H<sub>2</sub>S concentrations. SUP05 is a sulfur-oxidizing gamma proteobacterial group known to inhabit anoxic sulfidic waters up to 4μM H<sub>2</sub>S. High validated PPV is associated with increased SUP05 abundance and water column H<sub>2</sub>S concentrations. The higher validated PPV in upper waters appears to be associated with a second group of putative sulfur-oxidizing bacteria, Arctic96BD-19 closely related to SUP05.

## FUTURE DIRECTIONS

**FIGURE 5. Understanding the community.**

Microbial ecology is all about relationships. Statistical methods for inferring interactions between uncultivated microbial community members promise to shed new light on distributed metabolic networks in natural and engineered ecosystems. Extrapolated SAGs can provide additional metabolic insights with increased confidence where such methods fall short. Looking forward, combining statistical methods with SAGEX will provide even more powerful "optics" for exploring the microcosmos on relationship at a time.



## REFERENCES

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## ACKNOWLEDGMENTS

