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Possibilities and obstacles in recovery of genomes from elusive microbes in complex metagenomes

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Introduction

Representative genomes provide an entry point for understanding a given microbial ecosystem. The genomes give insights into the metabolic potential and possible roles of the bacteria, as well as being essential when applying other -omics based techniques. For elusive, unculturable bacteria, metagenomics can be a useful tool to extract their genomes directly from their environments. However the approach has limitations when the community complexity is high and

Conclusions

- Enrichments decreased overall complexity, changed relative abundance and facilitated better coverage binning and allowed for numerous high quality genome bins to be extracted.
- Despite the short-term enrichments microdiversity was still present, which compromised recovery of complete genomes from many species.
- The simulation underlines the devastating impact of micro-diversity on assembly and binning. •



microdiversity is present.

In this study, we explored the potentials and obstacles faced when assembling genomes from complex metagenomes using activated sludge as a model system.

Aims

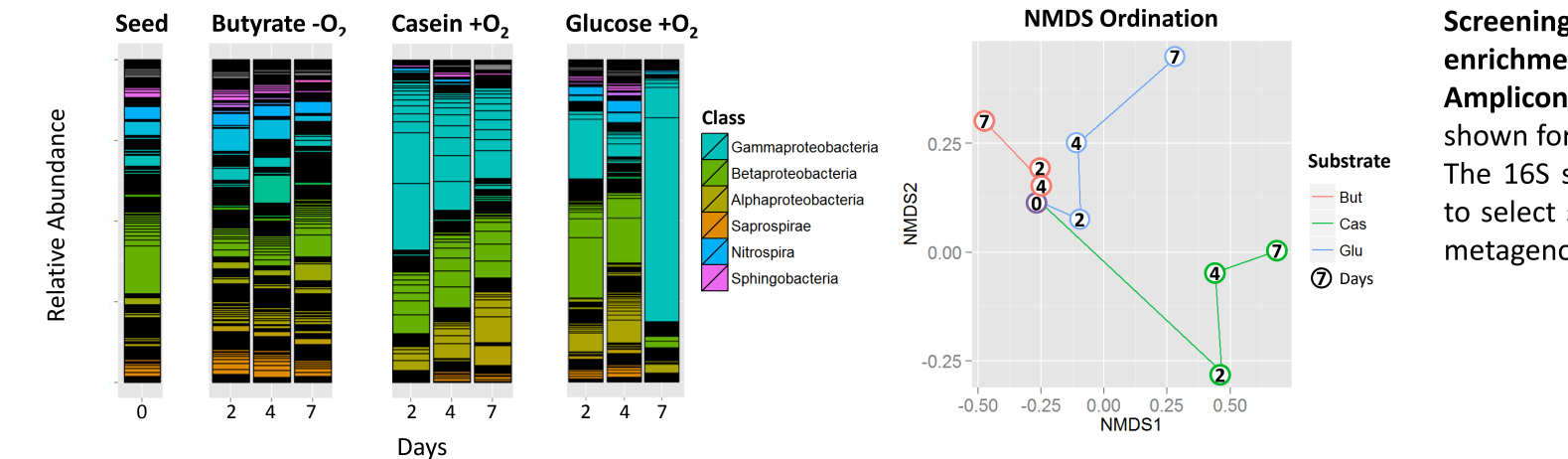
- To investigate the impact of microdiversity and community complexity on metagenomic assembly, binning and genome extraction.
- To investigate whether short-term enrichment can mitigate the adverse effects of the above

Methods

Short-term Enrichment

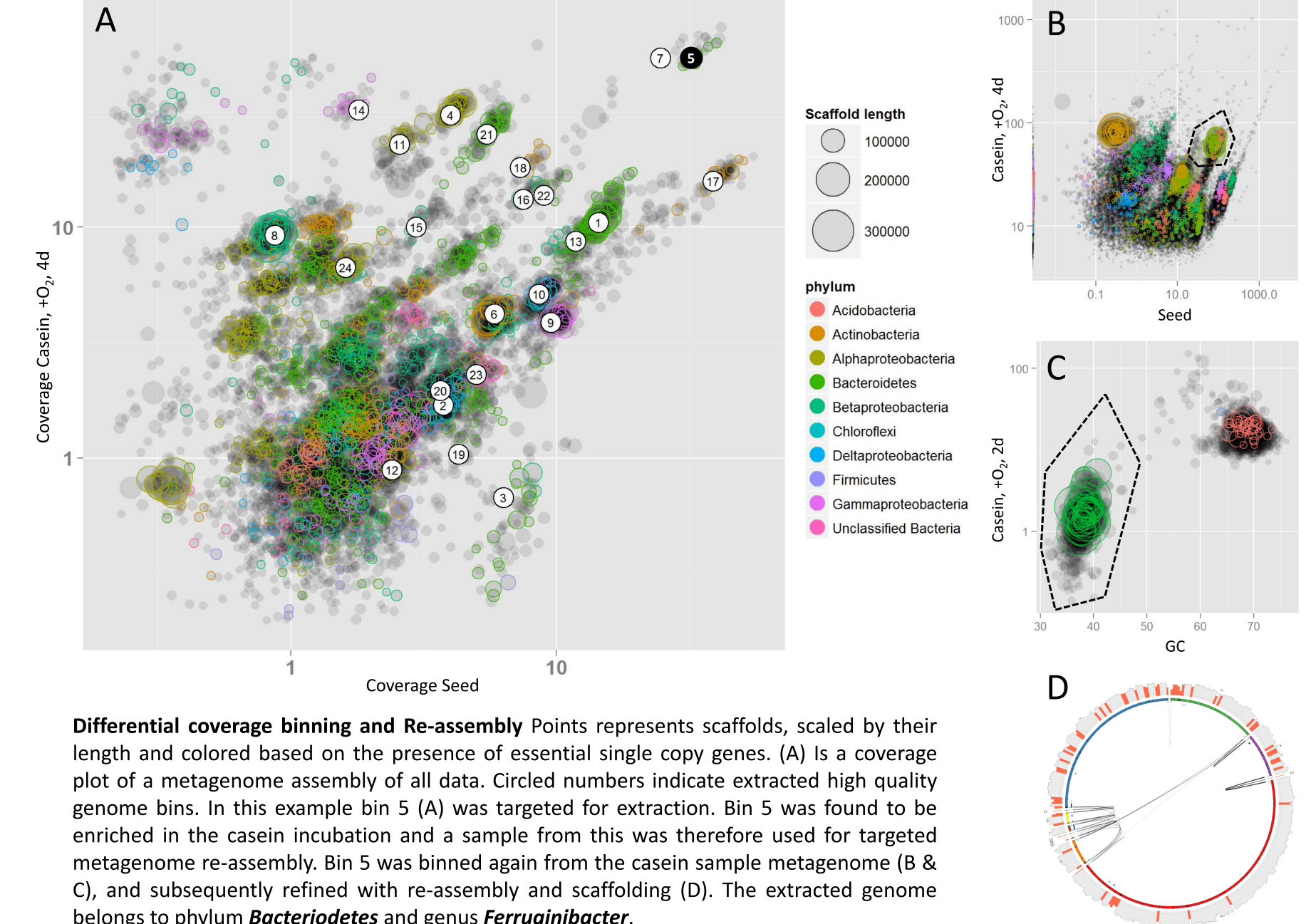
Seed sample Batch Incubations		Screen incubations		
Fullscale EBPR WWTP	Casein, Glucose or Butyrate	• V4 16S rRNA amplicons		

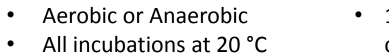
• 1 sample/d for 7 days



Screening of short-term enrichments with 16S V4 Amplicon sequencing. Data shown for 3 of 6 incubations. The 16S screening was used to select samples for further metagenome sequencing.

ΔΔΙΙ





• 14 of 50+ samples chosen for metagenome sequencing



Metagenome Binning

Sequencing and Assembly

- Illumina PE/MP 2x150bp (300 Gbp)
- De novo assembly (kmer: 64) and read mapping using CLC genomics workbench



Re-Assembly and Finishing

- Bin specific reads *de novo* re-assembled in SPAdes
- Manual finishing using SSPACE, Circos and R-biostrings



Coverage Binning

Essential genes

samples

• Read coverage profiles from 14

Paired-end read connections

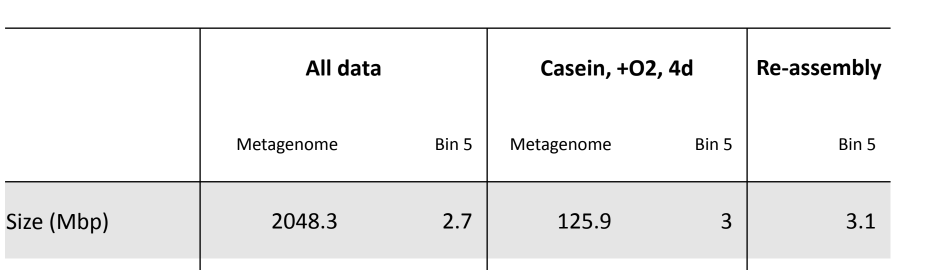
Tetranucleotide frequencies

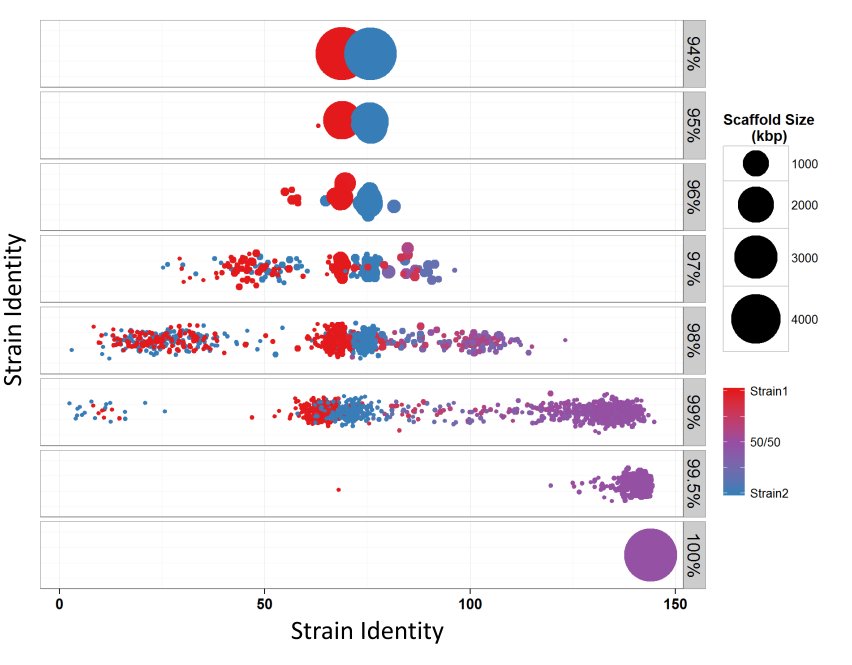
mmgenome

An R toolbox for reproducible genome extraction from metagenomes

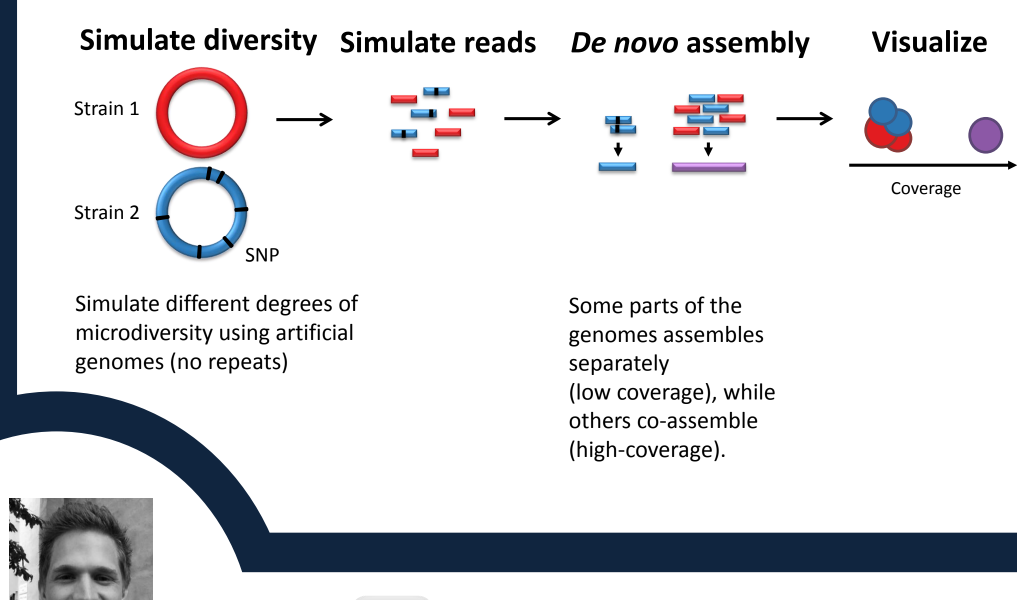
Taxonomic assignments

belongs to phylum *Bacteriodetes* and genus *Ferruginibacter*.





Microdiversity Simulation



Scaffolds	915 040	702	62422	357	21
Scaffold Max (kbp)	340	34	94	67	1382
Ess. Genes	43522	93/105	2996	104/105	105/105
Species	> 430	1	>29	1	1
Seq. data (Gbp)	300	-	15.5	-	_

Binning Statistics. The statistics show the genome extraction progression from the metagenome assemblies, through binning to the final refinement. It is clear that the assembly of the enriched casein sample with less microdiversity provides a better basis for binning, despite far less sequence data being used.

Simulation of different degrees of microdiversity. Strain identity refers to the percent bp in the two artificial genomes that are identical.

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