

1 Recovery of high-quality Genomes from a deep-inland Salt Lake Using 2 BASALT

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38 **Abstract**

39 Metagenomic binning enables the in-depth characterization of microorganisms. To improve the
40 resolution and efficiency of metagenomic binning, BASALT (Binning Across a Series of AssembLies
41 Toolkit), a novel binning toolkit was present in this study, which recovers, compares and optimizes
42 metagenomic assembled genomes (MAGs) across a series of assemblies from short-read, long-read or
43 hybrid strategies. BASALT incorporates self-designed algorithms which automates the separation of
44 redundant bins, elongate and refine best bins and improve contiguity. Evaluation using Critical
45 Assessment of Metagenome Interpretation (CAMI) datasets at low (132 genomes) and medium (596
46 genomes) complexities revealed that BASALT auto-binning gained up to 10 times of shared MAGs
47 with better quality when pairwise comparing to DASTool, VAMB and metaWRAP, resulting in up to
48 51% more MAGs obtained than other binners. Using BASALT, a case-study analysis of a Salt Lake
49 sediment microbial community from northwest arid region of China was performed, resulting in 426
50 non-redundant MAGs, including 352 bacterial and 69 archaeal MAGs which could not be assigned to
51 any known species from GTDB (ANI < 95%). In addition, two Lokiarchaeotal MAGs that belong to
52 superphylum Asgardarchaeota were observed from Salt Lake sediment samples. This is the first time
53 that candidate species from phylum Lokiarchaeota was found in the arid and deep-inland environment,
54 filling the current knowledge gap of earth microbiome. Overall, BASALT is proven to be a robust
55 toolkit for metagenomic binning, and more importantly, expand the Tree of Life.

56 **Keywords:** Microbiome; Metagenome; Binning refinement; Tree of Life; Salt Lake;
57 Asgardarchaeota/Lokiarchaeota

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69 **Introduction**

70 Metagenomics analyses accommodated numerous of approaches exploring microbial diversities,
71 biosynthetic potentials and evolutionary relationships of earth's microbiome (Tyson et al. 2004,
72 Temperton and Giovannoni 2012). Specifically, the development of sequencing technologies,
73 computational capacities and bioinformatic tools enabled genome-scale analyses, which freed our
74 cognition of microorganisms from only cultivated isolates and significantly boosted our
75 understandings on uncultivable microorganisms (Parks et al. 2017, Pasolli et al. 2019). Genome-
76 resolved metagenomics was firstly applied in 2004 from low microbial diversity environment (Tyson
77 et al. 2004), followed by a series of initiations such as Earth Microbiome Project (EMP) and the
78 European Nucleotide Archive (ENA) which enabled us to unravel the microorganisms on a global
79 scale (Thompson et al. 2017). For example, the latest report from EMP projects revealed more than
80 52,000 Metagenomic Assembled Genomes (MAGs) on species level (Amid et al. 2020, Nayfach et al.
81 2020), consisting a wide range of samples from environments with medium to high level of microbial
82 complexities, such as human (Pasolli et al. 2019, Almeida et al. 2021), freshwater (Ali et al. 2020),
83 marine (Tully et al. 2018, Reji et al. 2020), engineered environment (Ransom-Jones et al. 2017, Liang
84 et al. 2020) and soil (Kroeger et al. 2018, Nascimento Lemos et al. 2020), etc. Such studies
85 implementing genome-resolved metagenomic approaches have largely expanded branches of
86 microorganisms on tree of life (Hug et al. 2016). However, despite these findings, a vast majority of
87 microorganisms remain obscured due to 1) limitation of bioinformatic tools such as assembly and
88 binning; 2) large unexplored area/regions with specific environmental conditions; and 3) advanced
89 cultivation methods to be developed.

90 Aside from the developing innovations of culturing new microorganisms (Lewis et al. 2020), major
91 discovery of novel species was based on sequencing-based analyses. However, major impediments
92 that hampered us from obtaining comprehensive and high-quality MAGs from existing sequencing
93 datasets are assembly and binning steps (Nayfach et al. 2020). Due to the nature of next-generation

94 sequencing, a series of errors may occur in binning sourced from assembled short-read sequences, such
95 as mis-clustering contigs into bins, mis-separating contigs from one genome into multiple bins, and
96 mis-separating multiple genomes into bins sharing partial genomic sequences (Rinke et al. 2013, Yu
97 et al. 2018, Wang et al. 2019), resulting in redundant and artificial bins that interfere the actual binning
98 result. While the development of third generation sequencing can significantly increase the length of
99 sequencing reads to mitigate binning errors, higher costs (e.g. Pacific Biosciences sequencing, PacBio)
100 or error reads (e.g. Oxford Nanopore Technology, ONT) are still hindrances from acquiring intact
101 microbial genomes with higher completeness and lower contamination (Laver et al. 2015, Wang et al.
102 2015). To date, the development of assembly (Bankevich et al. 2012, Peng et al. 2012, Li et al. 2015),
103 binning (Alneberg et al. 2013, Wu et al. 2016, Kang et al. 2019, Nissen et al. 2021) and refinement
104 tools (Song and Thomas 2017, Urtskiy et al. 2018) have allowed us recovering MAGs from relatively
105 high diversity environments (Parks et al. 2017, Sczyrba et al. 2017), but an average of 70% of
106 sequencing reads were still unable to be exploited on genome-resolved analyses, which proportion is
107 even higher in more complexed environment such as soil (Howe et al. 2014, Nayfach et al. 2020).
108 Although a growing number of bioinformatic tools have implemented algorithms on third generation
109 sequencing datasets, such as hybrid assemblers (e.g. Unicycler, OPERA-MS, Wick et al. 2017,
110 Bertrand et al. 2019), a systematic estimation of these tools on complexed environmental samples is
111 crucially needed. Moreover, no study has utilized third generation sequences on post-binning
112 approaches calibrating assembled bins and complementing genome gaps, which can maximize the
113 exploitation of long-reads data to elevate recovered genome qualities. Therefore, to further expand tree
114 of life, more robust assembly and binning workflows, as well as post-binning refinement methods are
115 yet to be developed.

116 In addition to the demands of advanced bioinformatic tools, more investigations on remote
117 areas/regions, especially the ones from unique environmental conditions that absent from global
118 projects (Nayfach et al. 2020, Almeida et al. 2021) can largely expand our knowledge on earth genomic

119 pools. As such environmental parameters (physical, chemical and biological) have direct and indirect
120 effect on microbial communities (Berdjeb et al. 2011, Nishiyama et al. 2018, Easson and Lopez 2019,
121 Glasl et al. 2019), exploring uncommon environment can expand the breadth of our understanding on
122 microbial diversity and potential functions. A few studies have already launched targeting rarely
123 explored environments on deserts (Finstad et al. 2017), oil fields (Eze et al. 2020), hydrothermal vents
124 (Anderson et al. 2017) and non-marine soda lakes (Vavourakis et al. 2018), but such number of
125 investigations were still scarce to support the expansion of microbial diversity, compare to a vast
126 majority of studies on marine, soil and human-associated microbiomes (Hoshino et al. 2020,
127 Nascimento Lemos et al. 2020, Almeida et al. 2021).

128 In this study, we introduce a highly robust toolkit BASALT (Binning Across a Series of AssembLies
129 Toolkit) to robustly recover microbial genomes from sequencing datasets using a series of innovative
130 methodologies for assembly, binning and refinements. Firstly, BASALT uses high-throughput
131 assembly methods to automatically assemble/co-assemble multiple files in parallel to reduce the
132 manual input; Next, BASALT incorporates self-designed algorithms which automates the separation
133 of redundant bins to elongate and refine best bins and improve contiguity; Further, BASALT facilitates
134 state-of-art refinement tools using third-generation sequencing data to calibrate assembled bins and
135 complement genome gaps that unable to be recalled from bins; Lastly, BASALT is an open frame
136 toolkit that allows multiple integration of bioinformatic tools, which can optimize a wide range of
137 datasets from various of assembly and binning software. Using BASALT, we performed a case study
138 on sediment samples of Aiding Lake, Xinjiang China, a deep-inland hypersaline lake with high aridity
139 in the surrounding area (Figure S1, Guan et al. 2020), which is different from common chlorite saline
140 aquatic system. Based on the superior number and quality of MAGs obtained via BASALT, we present
141 taxonomic and genetic profiles on prokaryotic microbial communities of lake sediment samples,
142 including two Lokiarchaeota species from a recently discovered archaeal superphylum
143 Asgardarchaeota (Zaremba-Niedzwiedzka et al. 2017) that also found in our samples. Here, we

144 highlighted that BASALT can efficiently enhance the assembly and binning processes on metagenomic
145 sequences, which allows in-depth investigations on the existing dataset by increasing overall and
146 individual MAG quality. By acquiring more high-quality MAGs, we could potentially unfold much
147 more knowledges on known or unknown microbial taxa, potential functions and host-microbial
148 interactions, which further expand the tree of life.

149

150 **Results**

151 **Environment, pipeline and availability**

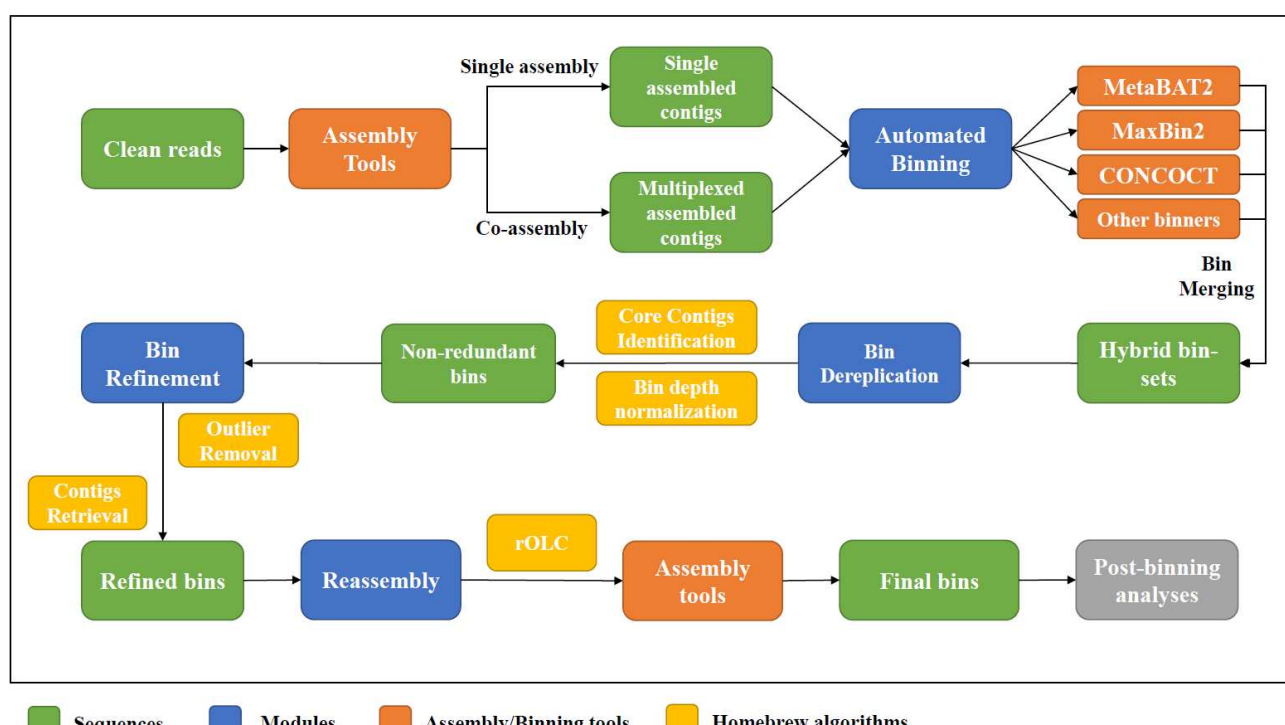
152 BASALT is command line software based on Python scripts with a series of modules, each containing
153 one or more algorithms/programs addressing data processing or analysis. Overall, BASALT is an
154 automated program running with one command line, while a few checkpoints are set in each module
155 to accommodate users to customize their preference to start at any checkpoint as needed. Further details
156 of outlines and algorithms are available at (<https://github.com/EMBL-PKU/BASALT>).

157

158 **BASALT workflow.**

159 BASALT is a versatile toolkit that provides comprehensive pipeline from mapping, automated binning
160 to post-binning refinement that enable users to retrieve non-redundant and high-quality MAGs from
161 metagenome samples. Overall, BASALT contains four major modules including Automated Binning,
162 Bin Dereplication, Bin Refinement and Reassembly, where five core algorithms were implemented in
163 these modules including Core Contigs Identification, Bin depth normalization, Outlier Removal,
164 Contigs Retrieval and Restrained Overlap-Layout-Consensus (rOLC, Figure 1). As BASALT enables
165 multiple assembly or co-assembly datasets from short-read sequences (next-generation sequencing,
166 NGS) or long-read sequences (third-generation sequencing, TGS) as input files, it is expected that a
167 potential increase of reads utilization is available (Stewart et al. 2018). By importing multiple

168 sequences with coverage information, BASALT conducted automated binning using prominent tools
169 such as MetaBAT2, Maxbin2 and CONCOCT (Alneberg et al. 2013, Wu et al. 2016, Kang et al. 2019)
170 that generated hybrid bin-sets. Raw hybrid bin-sets were firstly filtered with a homebrew Bin
171 Dereplication module to remove replicated bins obtained from the same assembly. The non-redundant
172 bins of each assembly were then merged and categorized into different groups at a customized average
173 nucleotide identity (ANI) cutoff. Each group of bins were further filtered by the homebrew algorithm
174 which classifies core contigs that further enables identification of redundant bins before selecting into
175 a single, hybrid bin-set obtained from all samples. The selected bin-set was further filtered using a
176 critical Outlier Removal algorithm, which integrated coverage and tetranucleotide frequency (TNF) to
177 remove outliers by using an interquartile ranges (IQR) method with multiple thresholds. Next,
178 sequences from assembly files were retrieved by connecting with existing contigs in the OR-filtered
179 bins to create an expanded sequences pool with potential connected contigs, while BASALT compared
180 and selected the refined bins with higher quality value. Further, a restrained Overlap-Layout-
181 Consensus (rOLC) step was conducted to overlap the replicated bins obtained from different
182 assemblies into OLC-merged bins, followed by the reassembly step to generate the finalized bin-set.
183 Notably, the sequence retrieval and reassembly step allowed utilization of long-read sequences
184 obtained from TGS to complement gaps and join overlapped regions on the bins to increase
185 completeness and reduce contamination.



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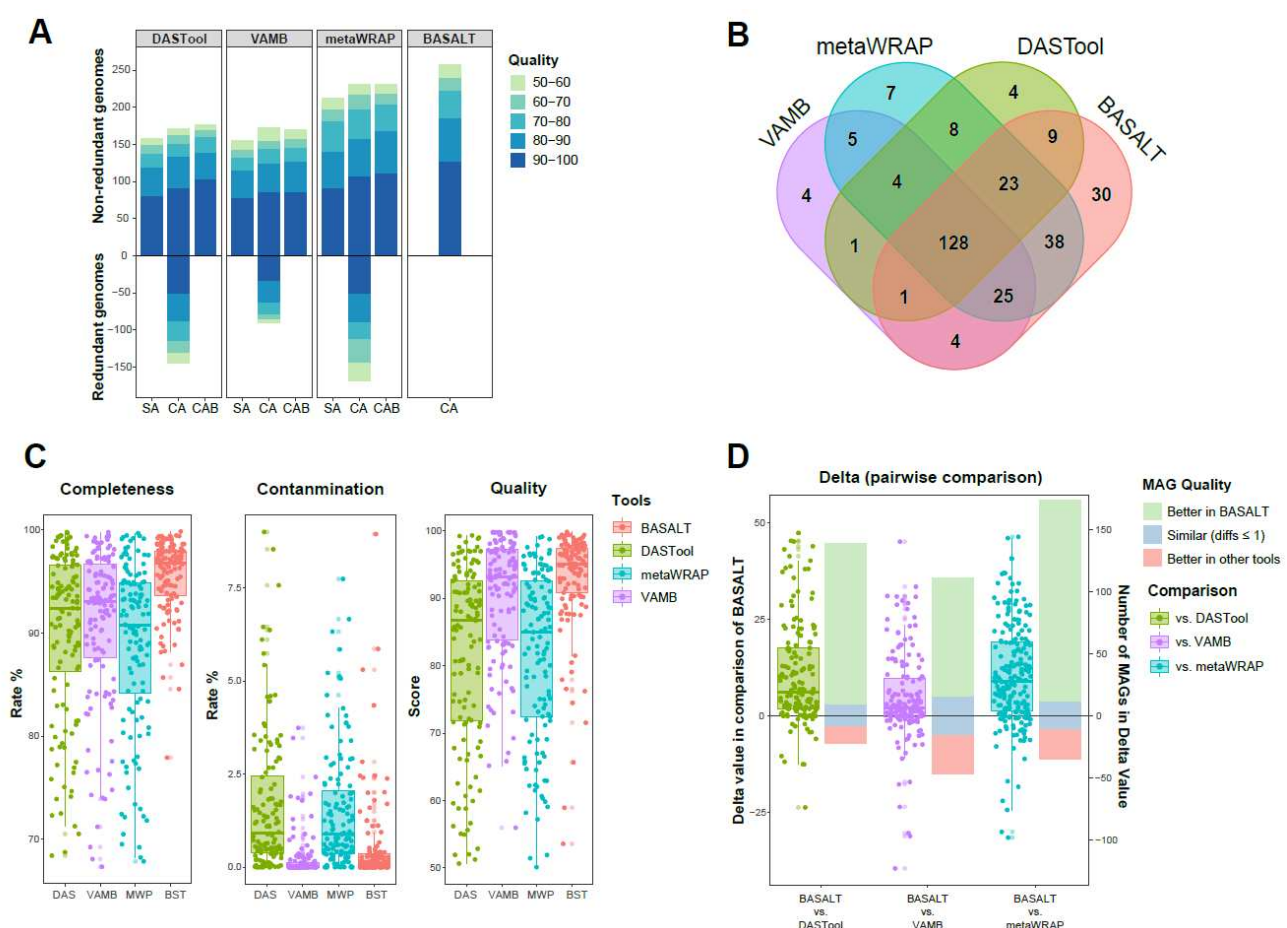
187 **Figure 1.** BASALT workflow for assembly, binning and refinement of metagenomic sequencing data (blocks
188 in green). BASALT contains four major modules (blocks in blue) including Automated Binning, Bin
189 DerePLICATION, Bin Refinement and Reassembly, where five core algorithms (blocks in yellow) were
190 implemented in these modules including Core Contigs Identification, Bin depth normalization, Outlier Removal,
191 Contigs Retrieval and Restrained Overlap-Layout-Consensus (rOLC). In addition to core workflow, assembly
192 and binning tools (blocks in orange) were also flexibly embedded in BASALT.

193

194 **BASALT improves recognition of non-redundant bins.**

195 BASALT pipeline enables multiple input files for assembly, including long-reads files for hybrid
196 assembly. The advantage of input with multiple files is not limited to the reduction of computational
197 time but could also generate more bins than individual assembled samples. For example, binning using
198 multiplexed samples generated 16.3%, 14.2% and 11.1% more non-redundant MAGs when using DAS
199 Tool (MCM), VAMB and metaWRAP (MCM), respectively on CAMI-medium dataset (Table S1,
200 Figure S2), while the increasing rate on CAMI-high dataset were 8.2%, 11.0% and 9.0%, respectively
201 (Table S1, Figure 2A). Despite the advantage above, a major drawback using multiplexed samples was
202 the byproducts of replicated bins, which were considered as redundant or pseudo- genomes. To address

203 this issue, BASALT pipeline accommodates a Bin Dereplication module with homebrew algorithms
204 which can remove redundant bins generated from the multiplexed assemblies as well as hybrid bin-
205 merging from automated binning, resulting in optimized, non-redundant bin-sets (Figure 1).
206 Comparing with standard CAMI-medium and -high genomes, DAS Tool, VAMB and metaWRAP
207 generated 85.9%, 77.1%, 95.0% (CAMI-medium) and 84.8%, 52.3%, 72.7% (CAMI-high) redundant
208 MAGs, respectively (Figure 2A, Figure S2), while no redundancy was observed in BASALT MAGs,
209 suggesting high redundancy rate were existed using the first three binning tools/toolkits. Remarkably,
210 BASALT Bin Dereplication, Refinement and Reassembly modules can not only eliminate redundant
211 MAGs generated from DAS Tool, VAMB and metaWRAP pipelines, but also increase the overall
212 quality of non-redundant MAGs (Co-assembled data refined with BASALT, CAB, Figure 2A),
213 suggesting good efficiency of redundancy removal and quality improvement using BASALT from co-
214 assembled bins.



216 **Figure 2.** Comparison of BASALT with other binning tools/pipelines on CAMI-high dataset. **A)** Number of
217 MAGs recovered from CAMI-high dataset using DASTool (MaxBin2, CONCOCT and MetaBAT2, MCM),
218 VAMB, metaWRAP (MCM) and BASALT. In the first three tools, Co-assembly (CA) resulted in higher number
219 of non-redundant MAGs compare to single assembly (SA) approach, while BASALT refinement module (Co-
220 assembly refined with BASALT, CAB) removed redundant MAGs and generated higher quality of MAGs in
221 the co-assembly approach. Color of bars indicated the quality of MAGs (50-100, from light to dark). **B)** Venn
222 diagram showing number of MAGs recovered using different tools. There were 128 MAGs found shared across
223 all tools, while 4, 4, 7 and 30 MAGs were uniquely recovered using DAS Tool (green), VAMB (purple),
224 metaWRAP (cyan) and BASALT (red) pipelines, respectively. **C)** Completeness, contamination and quality of
225 128 shared MAGs recovered using DAS Tool (DAS, green), VAMB (purple), metaWRAP (MWP, cyan) and
226 BASALT (BST, red). MAGs recovered using BASALT had lower contamination rate compared to DAS Tool
227 and metaWRAP, and higher completeness and quality compared to all other tools. **D)** Pairwise comparison of
228 MAGs shared by corresponding pairs of tools. Overall, BASALT was superior in obtaining higher qualitied
229 MAGs compared to other tools. The number of MAGs that BASALT gained higher quality value (bars in light
230 green) was much more than the number of MAGs that other tools gained higher quality value (bars in light red)
231 or had similar quality value (difference of value ≤ 1 , bars in light blue).

232

233 **BASALT generates higher number and qualitied MAGs in synthetic microbial communities.**

234 The BASALT refinement module not only removes redundant bins generated in other pipelines, but
235 also improves the quality and number of MAGs. In comparison of MAGs (Quality score ≥ 50) obtained
236 via different toolkits, BASALT resulted in 27.1%, 7.0%, 1.7% (CAMI-medium) and 50.9%, 50%, 11.7%
237 (CAMI-high) more non-redundant MAGs than DAS Tool, VAMB and metaWRAP, respectively
238 (Figure 2A, Figure S2, Table S1). Moreover, in top-qualitied CAMI-high MAGs (Quality score ≥ 90),
239 BASALT obtained 38%, 47.7% and 17.6% more non-redundant MAGs than DAS Tool, VAMB and
240 metaWRAP, respectively (Figure 2A). This result suggested BASALT is more robust retrieving high-
241 qualitied and non-redundant MAGs from metagenome samples, especially on those samples with
242 higher complexity.

243 To individually evaluate the BASALT refinement module, a further refinement step was performed on
244 CAMI-high datasets processed with DAS Tool, VAMB and metaWRAP. Although less MAGs were

245 recovered compare to the result using comprehensive BASALT pipeline, 3.7%, 2.4% and 7.0% more
246 MAGs (Quality score ≥ 80) were retrieved, respectively, compared to the default approaches of other
247 toolkits (Table S1, Figure 2A). This result suggested that BASALT can also optimize number and
248 quality of MAGs even based on the datasets processed with other pipelines.

249 Comparing MAGs recovered from the abovementioned toolkits with reference genomes in CAMI-
250 high dataset, 128 MAGs were found universally presented across all pipelines, while 4, 4, 7 and 30
251 MAGs were uniquely recovered using DAS Tool, VAMB, metaWRAP and BASALT pipelines,
252 respectively (Figure 2B). In comparison of the 128 shared MAGs in their completeness, contamination
253 and quality score (completeness – 5*contamination) across different toolkits, BASALT has the overall
254 advantages in acquiring high-quality and low contamination MAGs in comparison with others (Figure
255 2C). Further, we performed pairwise comparison between BASALT and the other tools of shared
256 MAGs and calculated delta value (difference of MAG quality between two tools) on the same genome.
257 The delta value showed BASALT could retrieve 10, 3.1 and 6.8 times of high-quality MAGs compared
258 to DAS Tool, VAMB and metaWRAP, respectively (Figure 2D), indicating that BASALT can
259 substantially obtain better quality of MAGs comparing with other tools. In summary, BASALT can
260 retrieve a greater number of non-redundant MAGs from medium-high complexed samples with higher
261 qualities.

262

263 **BASALT efficiently retrieves genomes from high complexity samples.**

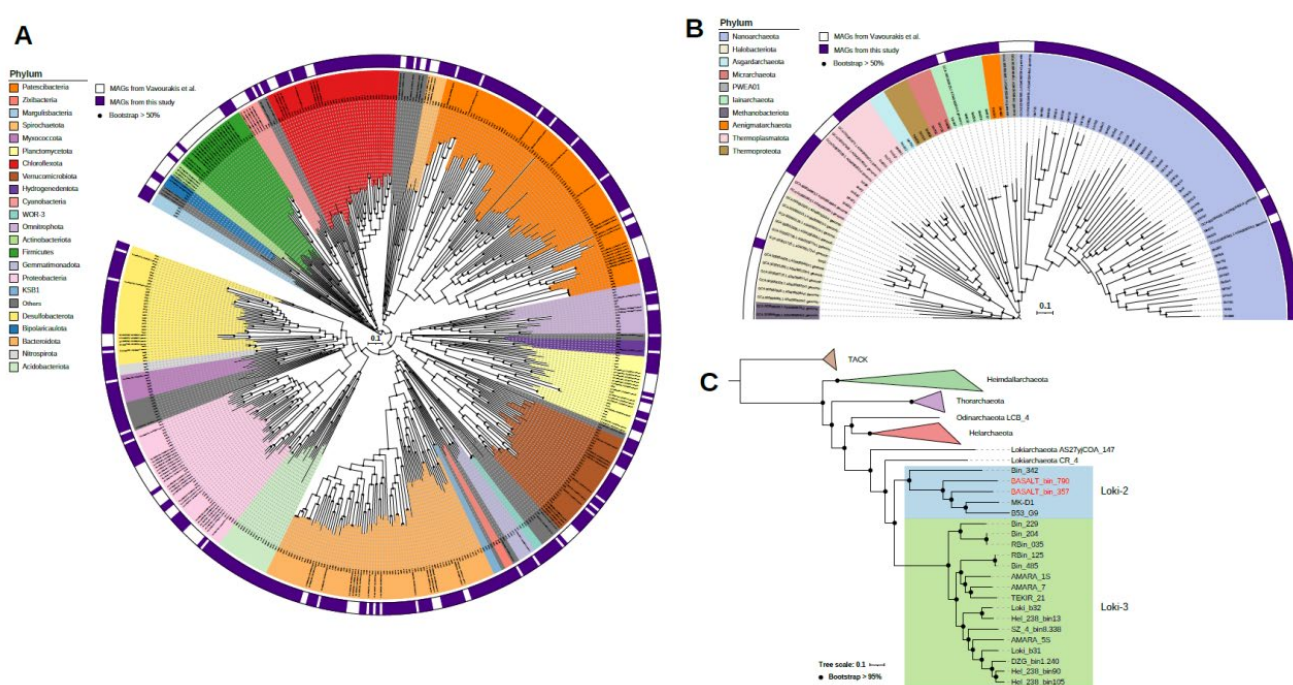
264 We performed four metagenome samples from Aiding Lake sediments using both BASALT to evaluate
265 the capacity of BASALT on improving genome qualities recovered from complexed environmental
266 samples. Using BASALT, assembled bins were quality-checked with CheckM (Parks et al. 2015),
267 resulting in 426 non-redundant MAGs (completeness – 5*contamination ≥ 50 , mean completeness =
268 79%, mean contamination = 1.7%, mean quality value = 70.4), including 113 MAGs above high-

269 quality level (quality ≥ 80). As a majority of metagenomic sequences could not be utilized to recover
270 high-quality genomes (Nayfach et al. 2016, Nayfach et al. 2020), we estimated the efficiency of
271 sequences utilized in the lake sediment samples, resulting in 26.9% of reads mapped to the MAGs,
272 which largely improved the utilization rate of metagenomic sequences in recovered MAGs compared
273 to other samples with high complexity (Howe et al. 2014).

274 In reference to Genome Taxonomy Database (GTDB, Parks et al. 2018), all 426 MAGs from BASALT
275 pipelines were annotated spanning 39 bacterial and 8 archaeal phyla. MAGs classified into bacterial
276 phyla were mainly focused in Patescibacteria, Chloroflexota, Verrucomicrobiota, Bacteroidota,
277 Proteobacteria and Desulfobacterota, while MAGs classified into archaeal phyla were Halobacteriota,
278 Thermoplasmatota, Asgardarchaeota, Thermoproteota, Iainarchaeota and Micrarchaeota in archaeal
279 domain (Figure 3A), representing a unique characteristic of microbial communities in the Salt Lake
280 sediment. Among these 426 MAGs, 98.9% bacterial MAGs and 100% archaeal MAGs could not be
281 assigned to any known species from GTDB (ANI $< 95\%$). At genus level, 57.6% bacterial MAGs and
282 66.7% archaeal MAGs could not be assigned to any known genus from GTDB. This result suggested
283 that there was a large repository of genetic pool to be uncovered in Salt Lake sediments.

284 To further validate the MAGs recovered from Aiding Lake sediments, we compared these assembled
285 genomes with another study by Vavourakis et al. (2018) where samples were collected from several
286 soda lakes in the Kulunda Steppe (south-western Siberia, Altai, Russia). Generally, the two datasets
287 shared a vast majority of phyla (Figure 3A), indicating that despite the different geographical location,
288 bacterial assemblages of the Salt Lakes might be similar. Specifically, diverse bacterial taxa from
289 phylum Patescibacteria were observed in our study, which paralleled with the study by Vavourakis et
290 al. (2018). On the other hand, in the unique of phyla observed in two datasets, Acidobacteriota,
291 Nitrospirota, Zixibacteria, WOR-3 and KSB31 were only present in this study (Figure 3A). In archaeal
292 communities, MAGs from phyla Halobacteriota, Thermoplasmatota, Iainarchaeota and
293 Nanoarchaeota were observed in both datasets, while phyla Methanobacteriota and PWEA01 were

294 only present in Vavourakis et al. (2018) and phyla/superphyla Asgardarchaeota, Thermoproteota,
295 Micrarchaeota and Aenigmataarchaeota were only present in our study (Figure 3B). In the shared phyla,
296 a large number of MAGs from phylum Nanoarchaeota found, but only one MAG from phylum
297 Halobacteriota observed in this study, while a large community of Halobacteriota was observed in
298 Vavourakis et al. (2018). The difference of the two datasets might be due to the physiochemical
299 parameters between two contrasting sites of lakes (Sorokin et al. 2014), but such differences could also
300 be related to the different strategies of assembly, as well as binning pipelines.



301

302 **Figure 3.** Phylogenetic trees of **A)** bacterial MAGs from Vavourakis et al. (2018) and this study, **B)** archaeal
303 MAGs from Vavourakis et al. (2018) and this study, and **C)** Maximum-likelihood tree of Asgardarchaeotal
304 MAGs based on protein encoded genes from the whole genome. The unrooted phylogenetic trees in A) and B)
305 were constructed with Fasttree, while the Maximum-likelihood tree in C) was constructed with PHYML and re-
306 rooted with superphylum TACK. The two Lokiarchaeota MAGs found in this study were highlighted with red
307 in C), and all genomes used to create trees that not obtained from this study were listed in Table S2.

308

309 **Newly discovered Lokiarchaeota species from a deep-inland non-marine hypersaline lake.**

310 Using four metagenome samples, BASALT pipeline expanded 421 of known species on prokaryotic

311 phylogenetic tree with samples from one Salt Lake. Remarkably, two MAGs classified as
312 Lokiarchaeota belongs to the superphylum Asgardarchaeota, were also observed in our results.
313 Although previous study has reported Asgardarchaeota phyla found in hypersaline lakes (Bulzu et al.
314 2019), to the best of our knowledge, this is the first time that archaeal phylum Lokiarchaeota was found
315 from the deep inland non-marine samples. Comparing with other MAGs/isolates obtained in previous
316 studies, the two MAGs found in our study were grouped with Loki-2 MAGs/isolates (Figure 3C), next
317 to *Ca. Prometheoarchaeum syntrophicum*, a strain isolated from a deep-sea sediment sample at Nankai
318 Trough, Japan (Imachi et al. 2020). Interestingly, other Loki-2 MAGs were found globally, including
319 Bin_342 from Shark Bay, Australia (Wong et al. 2020) and B53_G9 from Guaymas Basin, US (Seitz
320 et al. 2019), suggesting that Loki-2 species were widely distributed not only in marine sediments but
321 also in the deep-inland terrestrial environment.

322

323 **Discussion**

324 BASALT is superior in quality and number of MAGs on low (132 genomes) to medium (596 genomes)
325 complexity samples. In regards of MAG quality assessment, CheckM (Parks et al. 2015) is widely
326 used in a vast majority of studies. However, in the presence of standard CAMI datasets, we calculated
327 the MAG quality against the corresponding genomes that can result in more accurate evaluations.
328 Notably, our dereplication module implemented in the integration step can efficiently remove
329 redundant bins that generated in co-assembly and bin selection steps, which was evidenced in the test
330 analysis using both CAMI-medium and CAMI-high datasets (Figure 2A). Although there are other
331 redundancy removal methods such as dRep (Olm et al. 2017), result suggested that redundant genomes
332 cannot be efficiently identified and removed using dRep on CAMI-medium or CAMI-high datasets
333 (Table S1), suggesting that BASALT Bin Dereplication gains more advantages in removing redundant
334 bins under higher complexed samples. Due to the scarcity of standard synthesized community with

335 high complexity to date, further works testing the efficiency of Bin Dereplication module on the
336 standardized samples with high complexity are needed.

337 BASALT-integrated Bin Dereplication module can efficiently remove redundant bins generated by
338 binning tools. However, due to the major impediments of short-fragmented technology of next-
339 generation sequencing (NGS) and post sequencing algorithm, recovered MAGs cannot specify
340 differences of genomes at strain level with high similarity. While long read sequencing such as ONT
341 has become more popular in the current metagenomic studies (Jain et al. 2016), high error rates still
342 required to be rectified by NGS. In this study, third-generation sequences were innovatively integrated
343 into our toolkit where long reads can be used to amend sequences and fill gaps on assembled genomes,
344 which can improve the overall quality of bins and increase the number of high-qualitied MAGs.
345 Although standard dataset with high complexity is currently not available in the database, our case
346 study revealed that Salt Lake sediments can be considered as samples with relatively high complexity
347 (6,993 ZOTUs from 16S rRNA gene amplicons), which was comparable with other studies on soil
348 samples regardless the primer selection (Fulthorpe et al. 2008, Xiong et al. 2021). In the context of
349 high complexed samples of Salt Lake sediments, BASALT could also efficiently conduct reads
350 utilization in metagenomic binning with 26.9% mapped sequences, which to some extent helped to
351 resolve the difficulty raised in the EMP project (Nayfach et al. 2020). Prospectively, the trend in the
352 development of third-generation sequencing has inspired that further exploitation of long-read
353 sequences can increase the resolution of MAGs at strain or single nucleotide polymorphism (SNP),
354 which may boost the outcome of recovered genomes in high complexity samples. Therefore, future
355 development should focus on the combination of NGS and TGS to improve the efficiency of binning,
356 which consequently expand our knowledge on the tree of life.

357 In addition to the BASALT toolkits introduced in this study, one major finding in the case study was
358 the discovery of two novel Lokiarchaeota genomes from the sediment samples of Aiding Lake.
359 Lokiarchaeota belongs to a recently discovered superphylum Asgardarchaeota, which is a hot topic

360 linked to the origin of eukaryotes (Spang et al. 2018). To date, candidate species of Lokiarchaeota were
361 universally found near deep sea hydrothermal vents and marine sediments (Spang et al. 2015, Spang
362 et al. 2018, Hoshino et al. 2020, Wong et al. 2020, Yin et al. 2020). A recent study has found candidate
363 Lokiarchaeota present in hypersaline lakes near Black Sea (Bulzu et al. 2019), whereas our study
364 highlighted the first time that Loki-2 species were found from deep-inland hypersaline lake sediments.
365 Given the genetic analysis on Lokiarchaeota along with other candidate Asgardarchaeota species
366 (Zaremba-Niedzwiedzka et al. 2017, Seitz et al. 2019, Imachi et al. 2020, Wong et al. 2020, Yin et al.
367 2020) have suggested that candidate Lokiarchaeota species were adaptive in marine environment with
368 distinct metabolic pathway, such as lignin or protein degradations. Thus, it was unlikely that candidate
369 Loki-2 species were newly emerged in the deep-inland hypersaline lake. Therefore, the discovery in
370 this study might have provided a landmark that candidate Lokiarchaeota species might exist in the
371 ancient age before the plate movement event occurred. However, insufficient MAGs/isolates revealed
372 to date hampered us to make rigid conclusions, that more investigation on this group of archaea is
373 critically required in the future studies.

374

375 **Methods**

376 **Overview of BASALT**

377 BASALT is a versatile toolkit that recovers, compares and optimizes MAGs across a series of
378 assemblies assembled from short-read, long-read or hybrid strategies. We established five homebrew
379 algorithms to carry out Core Contigs Identification (CCI), Bin Depth Normalization, Outlier Removal
380 (OR), Contigs Retrieval, and restrained overlap-layout-consensus (rOLC). These algorithms consist
381 three core modules of BASALT, such as Bin Dereplication, Bin Refinement and Bin Reassembly
382 modules. Besides, BASALT contains an autobinning module that uses mapping tools (e.g. bowtie2,
383 Langdon 2015) and binners (e.g. MetaBAT2, Maxbin2, and CONCOCT) to generate a raw hybrid bin-

384 set after input of assemblies (Figure 1).

385

386 **Dereplication of redundant bins**

387 Incompleteness and contamination of MAGs would hinder the dereplication of bins from the hybrid
388 bin-set. We developed an effectively strategy that can remove most of the potential contaminated
389 sequences while large number of sequences are still kept to carrying out precise redundancy
390 identification. Different from previously reported genome-wide ANI-based and marker gene- based
391 de-replicating methods such as dRep (Olm et al. 2017), the present method firstly generated a core
392 contig pool by filtering out potentially contaminated contigs of target bins. The depth and
393 tetranucleotide frequency (TNF) value of the selected contigs ranked from a range of 25 to 75
394 percentile of all the contigs of target bins. Then, selected bins were grouped into different raw bin-sets
395 based on overall similarity of core contigs. Secondly, we evaluated the sequencing depth discrepancy
396 among bins across different bin-sets to ensure the correct identification of redundant bins with relative
397 low similarity. The average depth of one bin should be equivalent to the average depth of the core
398 contigs of this bin. For neutralizing the sequencing depth discrepancy yielded by mapping the same
399 reads to different assemblies, we designed a method to calculate the normalization ratio between two
400 possibly redundant bins, which identifies near-identical sequences (99.8% similarity across longer than
401 50% of the whole length of sequence) between two potential redundant bins as candidate bins. A depth
402 normalization ratio between candidate bins was then calculated by using the depth of these contigs,
403 which was then used to neutralize the average depth of candidate bins. Those candidate bins with
404 similar average sequencing depth ($\delta \leq 10\%$) were considered as redundant bins to be removed.
405 Finally, the bin with better quality value (completeness – 5* contamination) estimated by CheckM
406 (Parks et al. 2015) was kept being the better bin for further refinement and reassembly.

407

408 **Refinement**

409 BASALT refinement module contains two adversary processes: Outlier Removal (OR) and Contig
410 Retrieval. To effectively remove contaminated contigs from a certain bin, we designed an outlier
411 removal algorithm that removes contigs with an outlier value of sequencing depth or TNF based on an
412 interquartile range (Formula 1), while different thresholds (k) were set (e.g. 1, 1.5, 3) to determine
413 these contigs. In the context of bin quality, OR keeps bins with higher quality value, while bins with
414 lower quality were discarded. If no refined bin with higher quality than the original bin was generated,
415 OR would acquiescently eliminate sequences marked as depth or TNF outliers under the threshold of
416 3. Notably, the default setting of OR mainly removes contaminated sequences, but it may cause an
417 unnecessary removal of contigs due to restricted threshold.

418 ***Formula 1:***

$$419 \quad x_i > Q3 + k(IQR) \vee x_i < Q1 - k(IQR)$$

420 In this formula, Q1 and Q3 stands for 25 and 75 percentiles of all contigs, where IQR (interquartile
421 ranges) was calculated by $Q3 - Q1$. $k \geq 0$.

422 Contig Retrieval algorithm was designed to retrieve sequences that have not been clustered into the
423 target bin in the binning process, especially multicopy sequences or unnecessarily removed sequences
424 by Outlier Removal. Contig Retrieval identifies contigs that potentially connected to existing
425 sequences in the target bin. These candidate contigs were further assessed by an interquartile range
426 method to remove depth or TNF outliers as described above and connected to the target bin by paired-
427 end tracking (Albertsen et al. 2013) or long-read mapped method, forming a refined bin. Refined bins
428 were expected to have higher quality value than target bins which were further selected to form a
429 refined bin-set for further reassembly.

430

431

432 **Reassembly**

433 BASALT reassembly module includes a restrained overlap-layout-consensus (rOLC) process and a
434 reassembly process. The rOLC algorithm was designed to retrieve sequence which was not included
435 in the BASALT binning and refinement processes. Specifically, this process re-utilized sequences from
436 redundant bins identified and removed from the dereplication module to overlap the sequences from
437 the target bin. Using a loose threshold (overlap length: 300 bp; similarity: 99%), rOLC algorithm
438 aggressively recorded the redundant candidate sequences removed from the target bin in the previous
439 steps. As rOLC process may increase the contamination of sequences, a reassembly step was
440 implemented after rOLC to amend the contamination caused by rOLC, and more importantly, to
441 precisely elongate the length of sequences in the target bin. In the reassembly process, short-read or
442 long-read sequences were extracted from datasets by Bowtie2 and Minimap2, respectively (Langdon
443 2015, Li 2018). Three state-of-art assemblers were implemented in the reassembly step: SPAdes, FLYe,
444 and Unicycler (Bankevich et al. 2012, Wick et al. 2017, Kolmogorov et al. 2019) to carry out short-
445 read reassembly, long-read reassembly and hybrid reassembly, respectively. A final bin selecting
446 process was exploited to select the best bins from the reassembly step to form the final bins for post-
447 binning analysis.

448

449 **Sample collection**

450 Salt Lake sediment samples were collected in July 2018 from Aiding Lake, an arid region in Turpan
451 City, Xinjiang Uygur Autonomous Region (42°52'9" N, 89°03'5" E). Briefly, about 50 grams of
452 sediment samples (n = 4) were randomly collected at 0-10 cm depth in the lake into sterile 50 ml falcon
453 tubes. Samples were immediately placed on dry ice before brought to laboratory and transferred to -
454 80 °C freezer until further DNA extraction was performed.

455

456 **DNA extraction and sequencing**

457 Frozen stored sediment samples (~250 mg dry weight per sample) was used to extract genomic DNA
458 using DNeasy PowerSoil Kit (Qiagen, Hilden, Germany), following the manufacturer's instructions.

459 Extracted DNA was quality checked by NanoDrop 2000 (Thermo Fisher Scientific, Waltham,
460 Massachusetts, US), quantity checked by Qubit Fluorometer (Thermo Fisher Scientific) and PCR
461 checked to confirm the amplifiability.

462 For 16S rRNA gene amplicon sequencing, barcode as a marker for each sample DNA was added at the
463 5' end of primers targeting V4 region for bacterial and archaeal communities (515F-806R, Caporaso
464 et al. 2012). Sequencing was performed at MAGIGENE, Guangzhou, China on an Illumina NovaSeq
465 6000 platform (2 × 250 bp paired end chemistry).

466 For shotgun metagenomics sequencing, quantity and quality checked genomic DNA was sent to
467 Novegene Co., Ltd, Nanjing, China on an Illumina NovaSeq (2 × 150 bp paired end chemistry).

468

469 **Sequencing processing**

470 To evaluate the efficiency of BASALT, standard Critical Assessment of Metagenome Interpretation
471 (CAMI) datasets including a simple-complexity (132 genomes, CAMI-medium) and a medium-
472 complexity (596 genomes, CAMI-high) synthesized communities were downloaded from
473 (<https://data.cami-challenge.org/participate>) (Sczyrba et al. 2017). Raw paired-end reads were initially
474 filtered using fastp (Chen et al. 2018). Fifty percent of bases were filtered based on a minimum quality
475 score of 5 and sequence length of 150 bp, allowing no ambiguous bases. Clean reads were individually
476 and co-assembled using SPAdes (version3.14.1, Bankevich et al. 2012) into contigs specifying k-mer
477 sizes of 21, 33, 55, 77 and finally reserved contigs > 1,000 bps. To compare with other binners/toolkits,
478 filtered contigs were processed with DASTool, VAMB, metaWRAP and BASALT, respectively. The
479 redundancy, completeness and contamination of the MAGs were calculated against standard CAMI

480 datasets using a homebrew script *Bin_quality_evaluation.py* available on github
481 (<https://github.com/EMBL-PKU/BASALT>) to ensure high accuracy of results obtained from the four
482 binners/toolkits. High-quality MAGs (completeness - 5*contamination \geq 50%) were kept for further
483 statistical analysis, whereas bins did not meet the quality were discarded.

484 For Aiding Lake sediment samples, raw 16S rRNA gene amplicon sequences were processed using
485 Quantitative Insights Into Microbial Ecology (QIIME2) pipeline (<http://qiime.org>) (Caporaso et al.
486 2010). DADA2 was used to filter low-quality sequences with lengths < 230 bp, remove chimeric
487 sequences, singletons, and join the quality-filtered paired-end reads. Unique sequences (100%
488 similarity) were taxonomically assigned using Naive Bayes classifier against the SILVA 16S rRNA
489 gene reference alignment database (release 123) (Pruesse et al. 2007). To compare the complexity of
490 Salt Lake sediment samples with other high-complexed samples such as soil, sequences were rarefied
491 to an even sampling depth at 10,000 reads per sample, before singleton was removed from the
492 generated OTU table. Overall, a total number of 6,993 ZOTUs (Zero-radius OTUs) were obtained.

493 For shotgun metagenomic sequences of Aiding Lake sediment samples, sequences were processed
494 following the same procedure on CAMI-medium and CAMI-high datasets using BASALT. The
495 completeness and contamination of the MAGs were then estimated using CheckM version 1.1.3 (Parks
496 et al. 2015) with lineage-specific marker genes and default parameters, with only high-quality MAGs
497 (completeness - 5*contamination \geq 50%) were kept for further analyses.

498

499 **Phylogenetic analysis**

500 The GTDB-Tk version (version1.4.1, Chaumeil et al. 2020) program was used to assign taxonomic
501 classifications to the MAGs (release r95). To make comparison with another study of soda lake samples
502 (Vavourakis et al. 2018), dereplicated MAGs were downloaded from NCBI Assembly database and
503 phylogenetic analyses were conducted based on MAGs from both studies using Fasttree (Price et al.

504 2010). For phylogenetic analysis of Asgardarchaeota, MAGs/isolates were downloaded from other
505 studies listed in Table S2, and Maximum-likelihood tree was constructed using PHYML version 3.0
506 (Guindon et al. 2010) with 1000 bootstrap iterations. Phylogenetic trees were visualized and edited in
507 the iTOL (<https://itol.embl.de>) online platform (Letunic and Bork 2019).

508

509 **Code availability**

510 All BASALT codes including homebrew scripts for quality checking against standard datasets are
511 available at (<https://github.com/EMBL-PKU/BASALT>).

512

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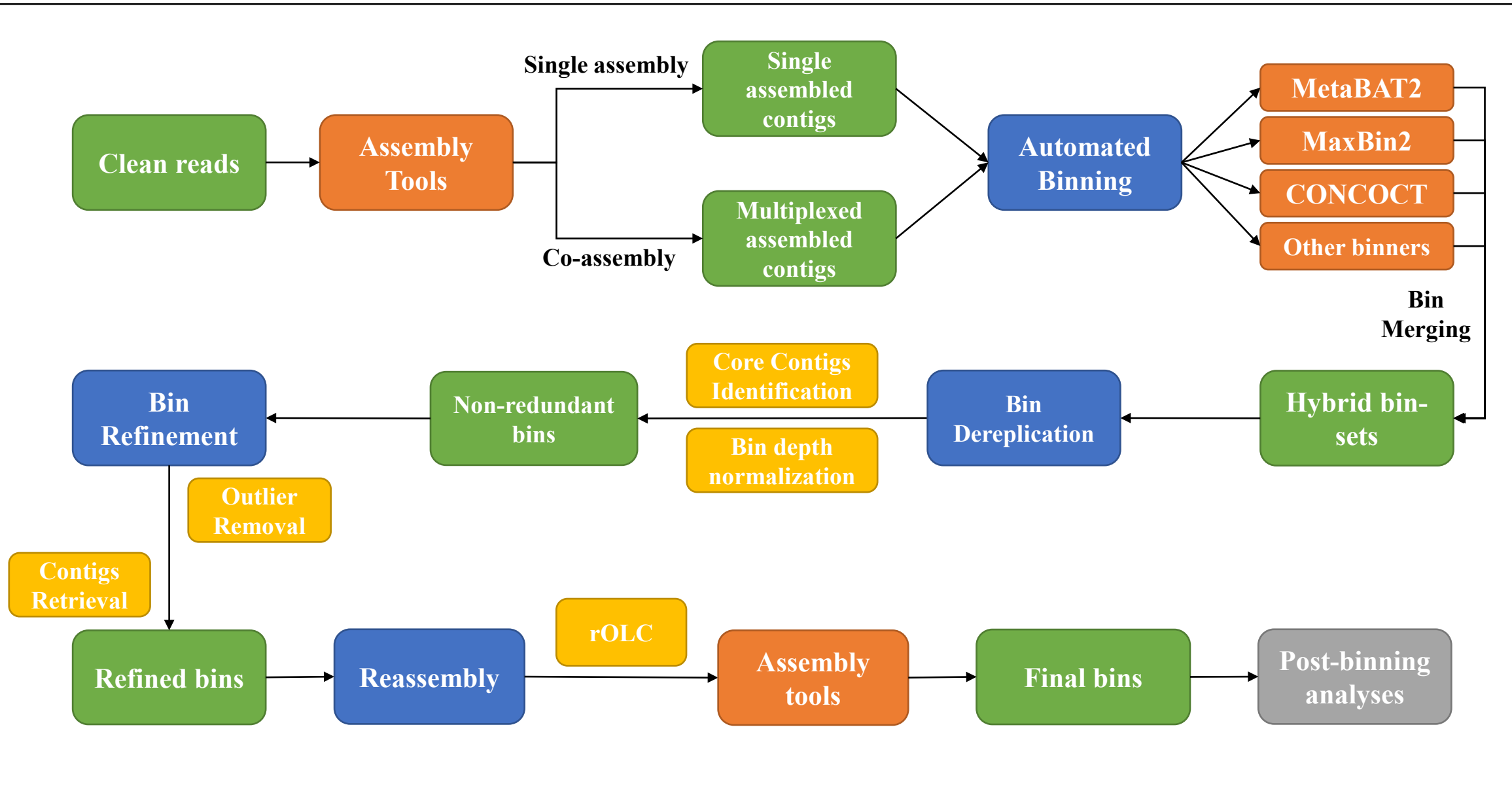
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Sequences



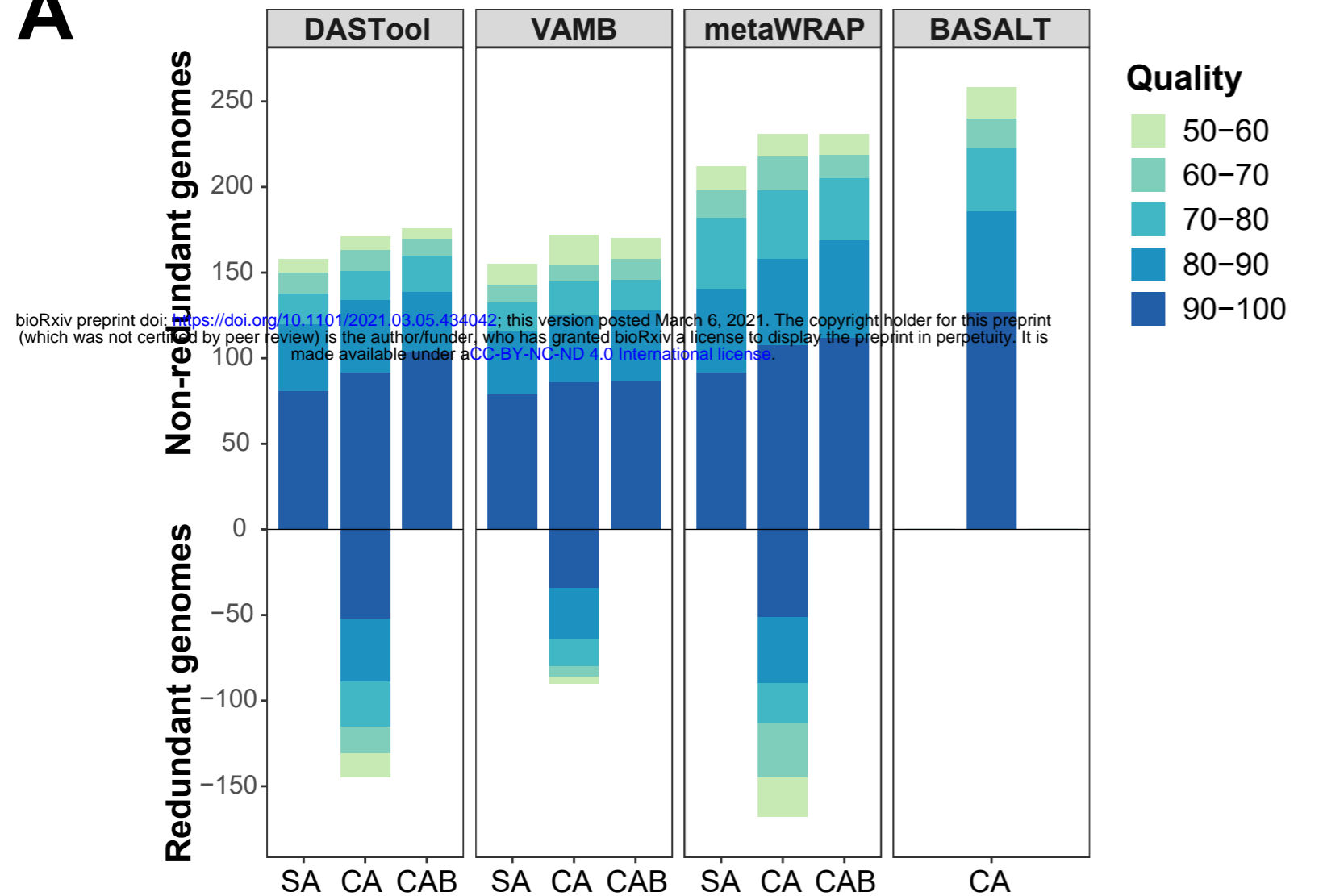
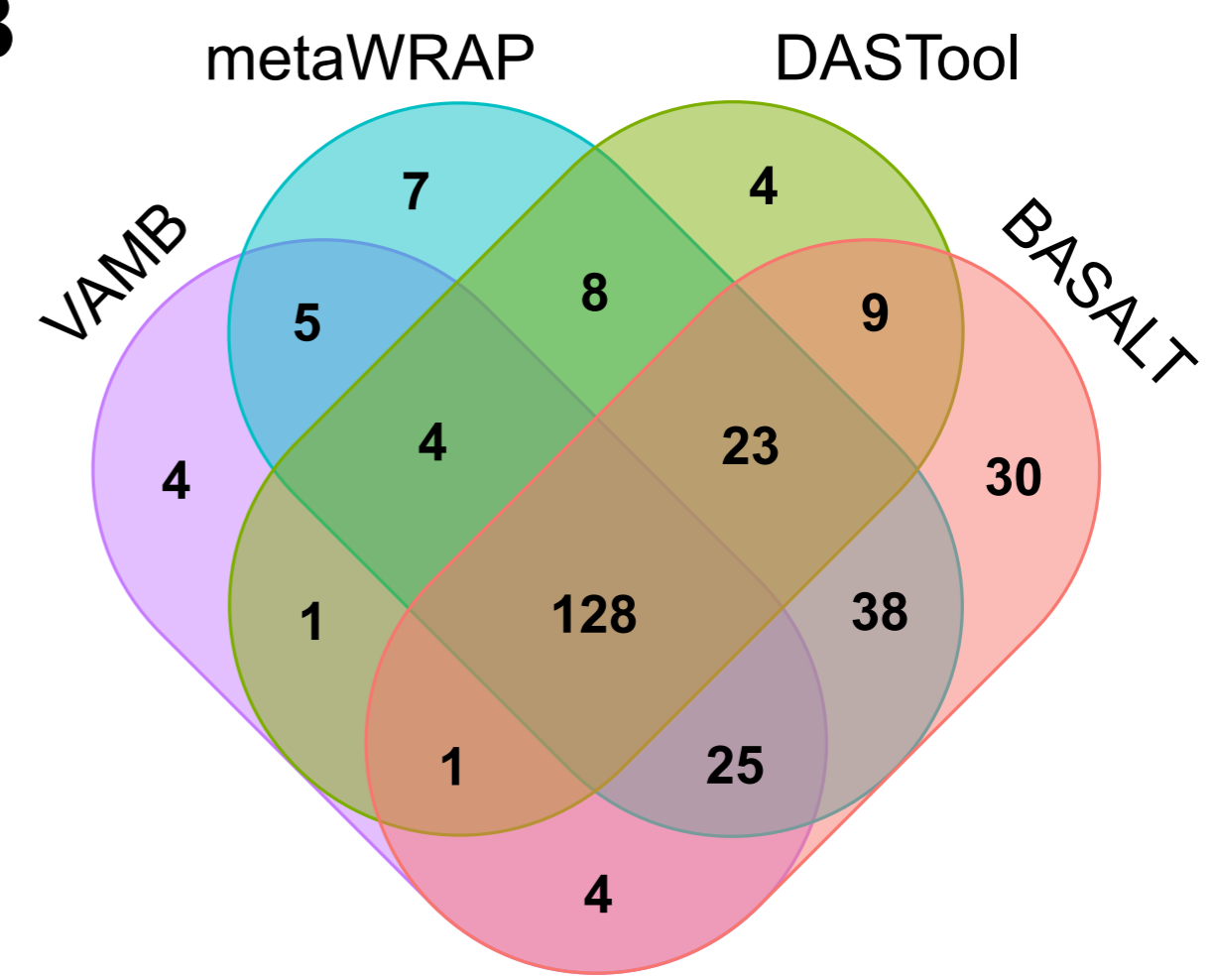
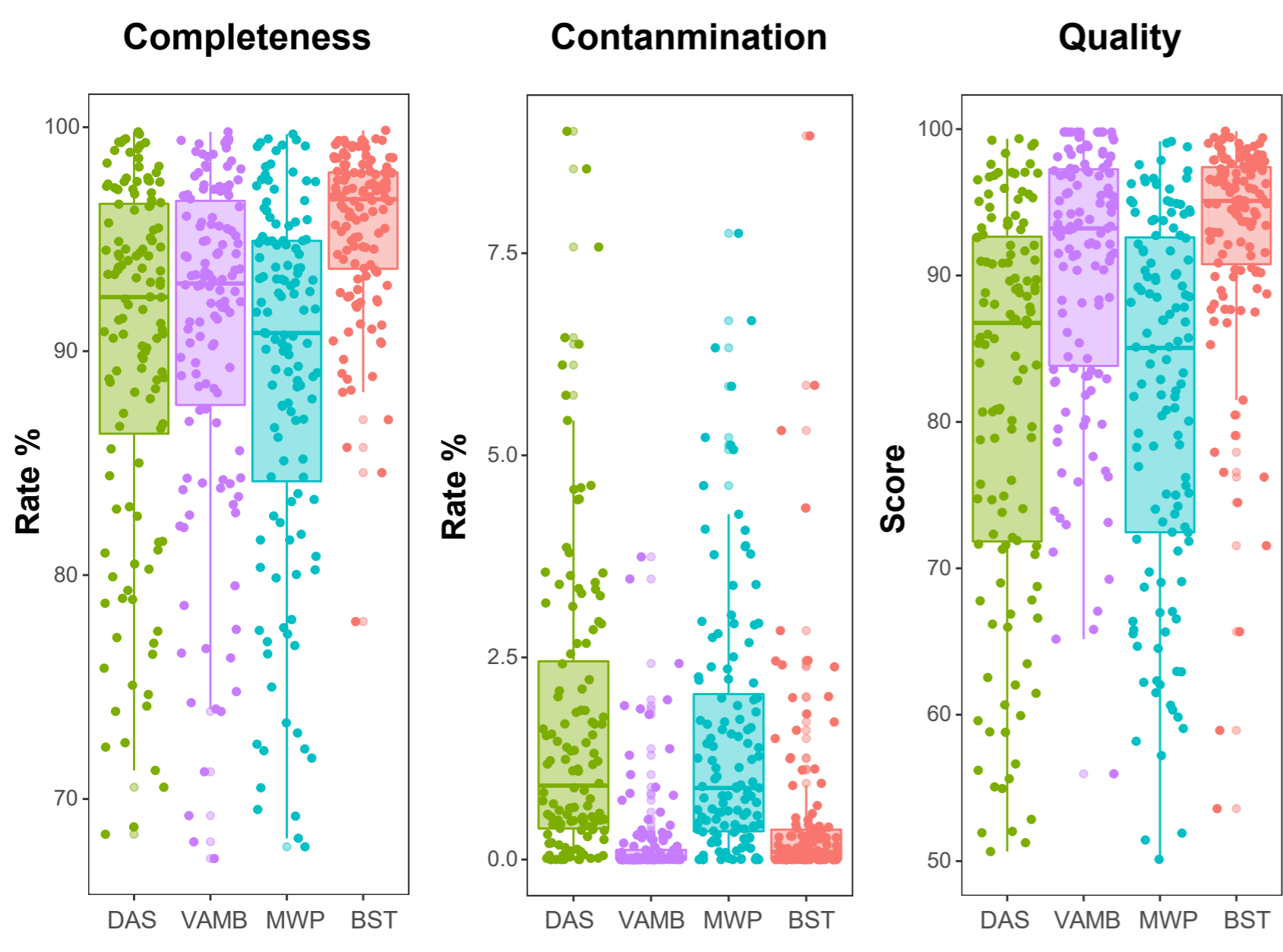
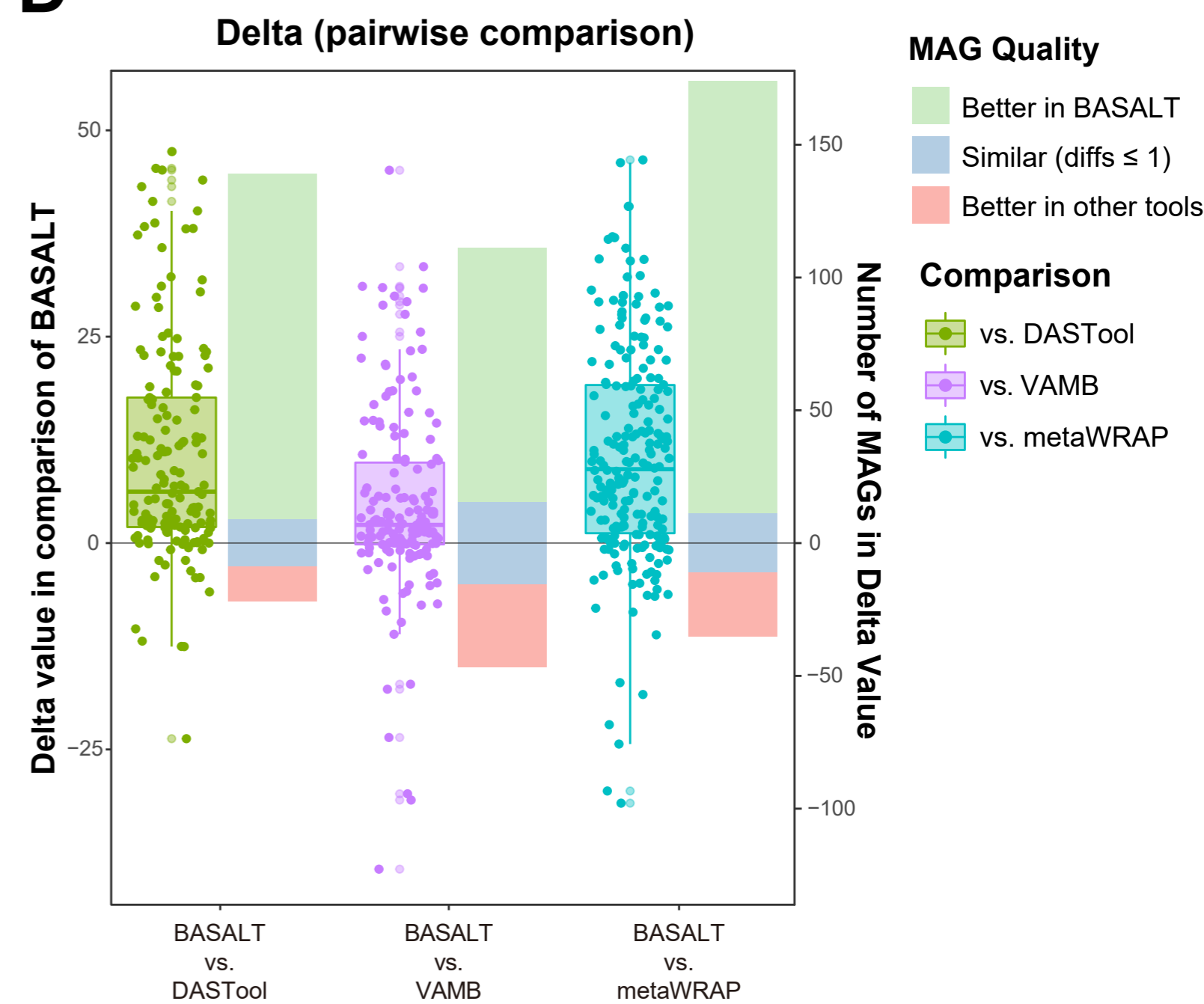
Modules



Assembly/Binning tools

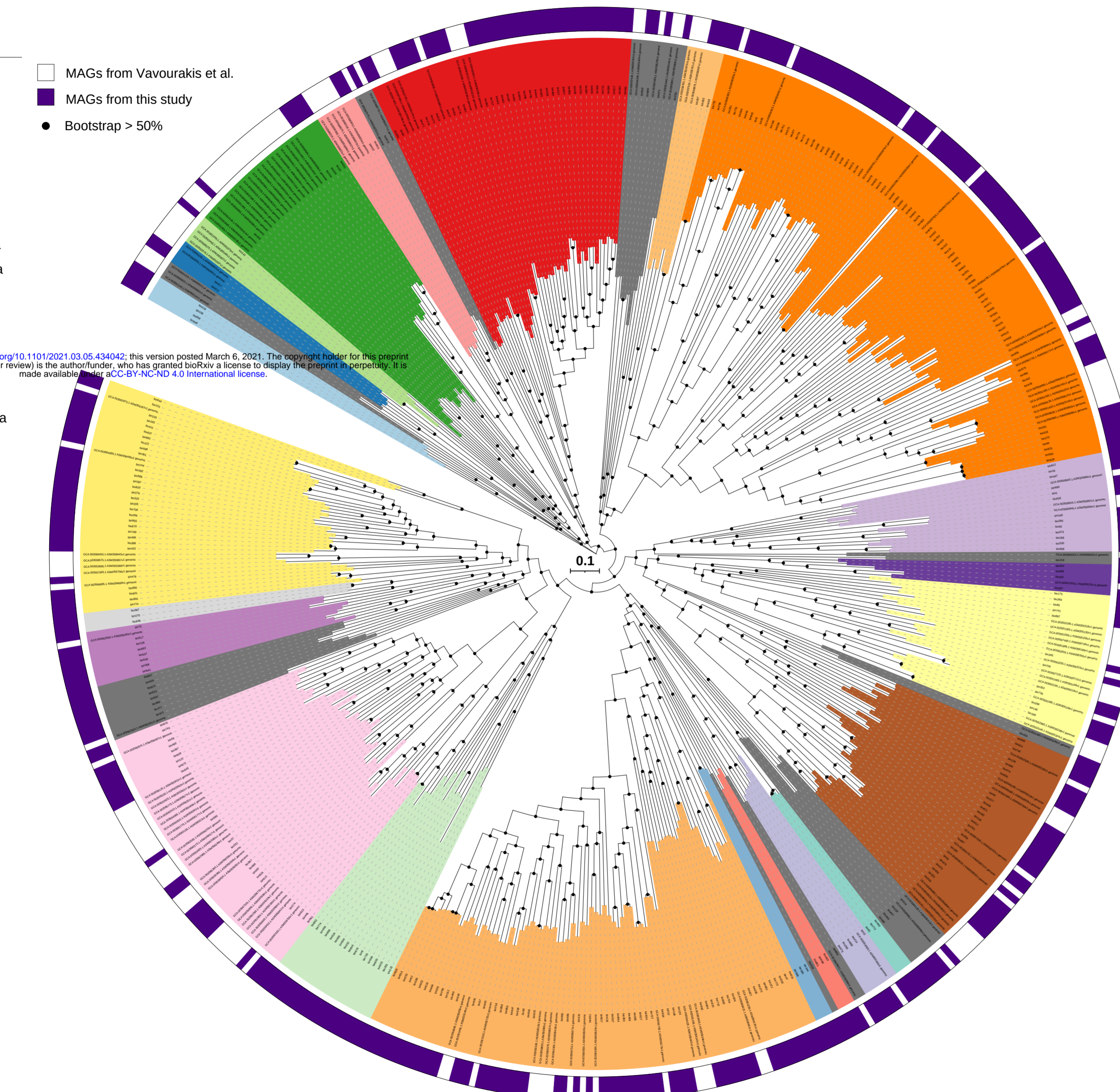


Homebrew algorithms

A**B****C****D**

A**Phylum**

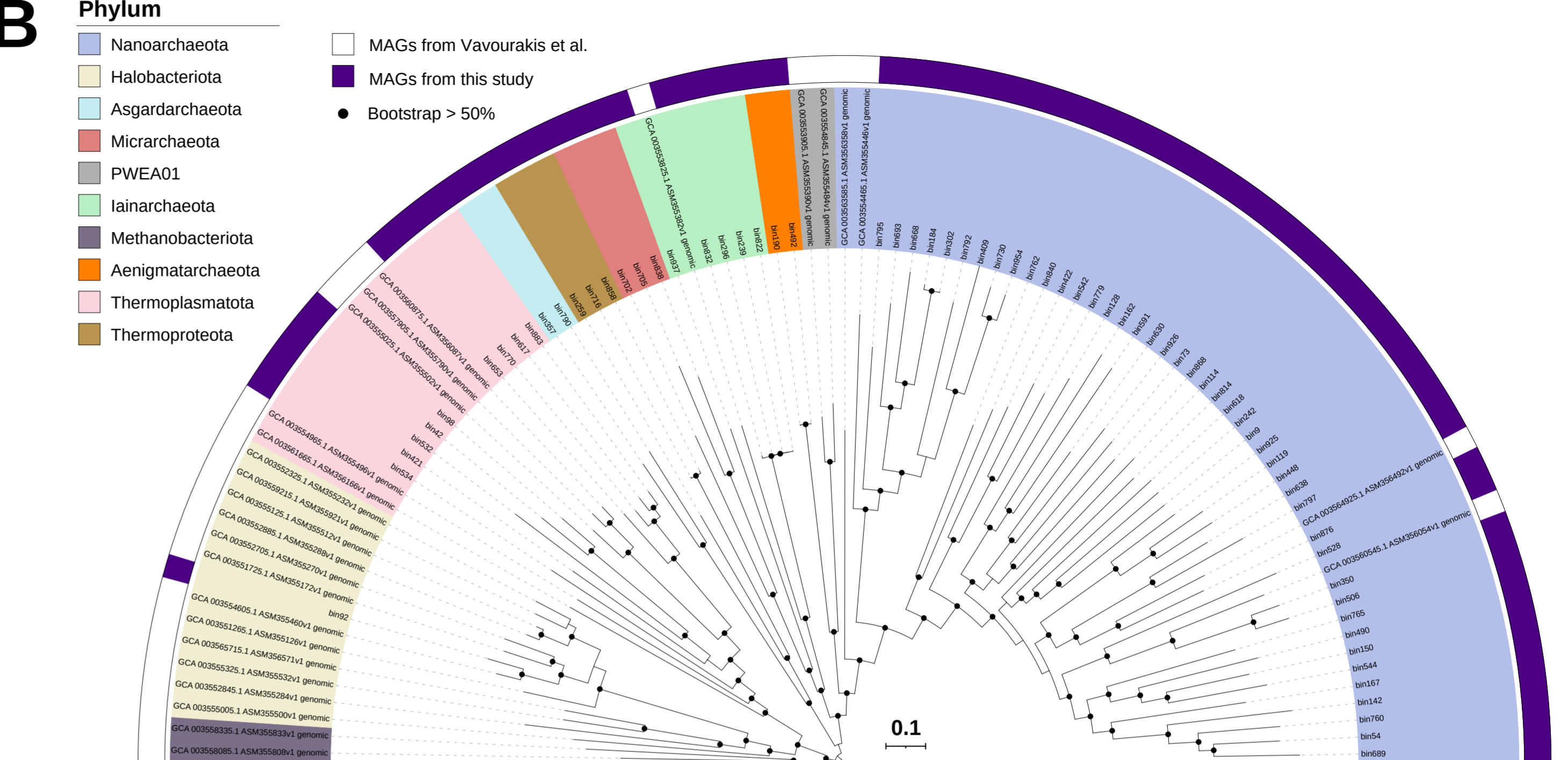
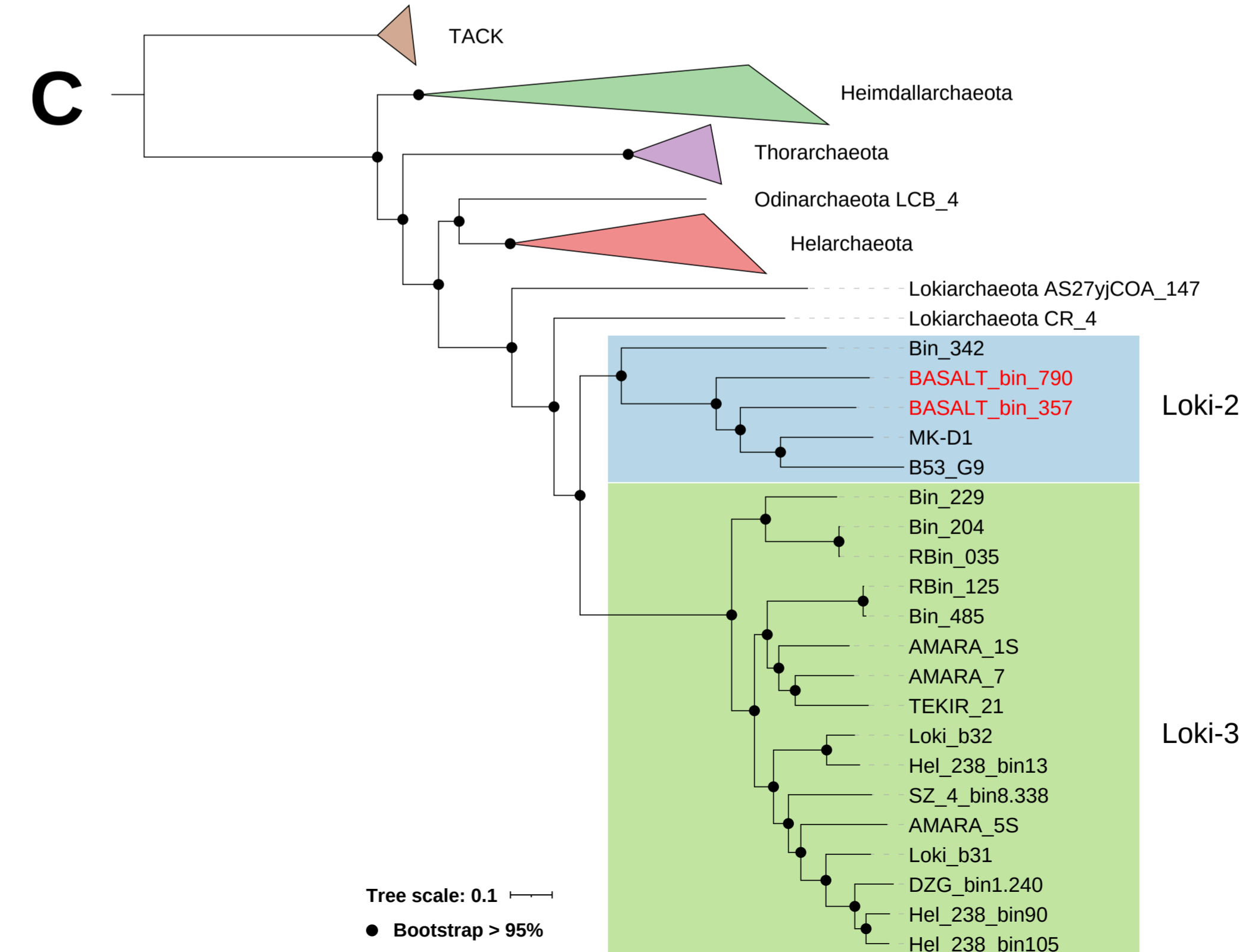
- Patescibacteria
- Zixibacteria
- Margulisbacteria
- Spirochaetota
- Myxococcota
- Planctomycetota
- Chloroflexota
- Verrucomicrobiota
- Hydrogenedentota
- Cyanobacteria
- WOR-3
- Omnitrophota
- Actinobacteriota
- Firmicutes
- Gemmimonadota
- Proteobacteria
- KSB1
- Others
- Desulfov bacterota
- Bipolaricau lota
- Bacteroidota
- Nitrospirota
- Acidobacteriota

**B****Phylum**

- Nanarchaeota
- Halobacteriota
- Asgardarchaeota
- Micrarchaeota
- PWEA01
- Iainarchaeota
- Methanobacteriota
- Aenigmarchaeota
- Thermoplasmata
- Thermoproteota

- MAGs from Vavourakis et al.
- MAGs from this study
- Bootstrap > 50%

Tree scale: 0.1
● Bootstrap > 95%

**C**

Loki-2

Loki-3