

1 Mapping-based genome size estimation

2 Boas Pucker^{1,2*}

3 1 Genetics and Genomics of Plants, Bielefeld University, Bielefeld, Germany

4 2 Center for Biotechnology (CeBiTec); Bielefeld University, Bielefeld, Germany

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6 Email: bpucker@cebitec.uni-bielefeld.de

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8 ORCID: 0000-0002-3321-7471

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10 * corresponding author: Boas Pucker, bpucker@cebitec.uni-bielefeld.de

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12 **Key words:** NGS, genome sequencing, k-mer profile, comparative genomics, *Arabidopsis*
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14 *mays*

15

16 Abstract

17 While the size of chromosomes can be measured under a microscope, the size of genomes
18 cannot be measured precisely. Biochemical methods and k-mer distribution-based approaches
19 allow only estimations. An alternative approach to predict the genome size based on high
20 contiguity assemblies and short read mappings is presented here and optimized on *Arabidopsis*
21 *thaliana* and *Beta vulgaris*. *Brachypodium distachyon*, *Solanum lycopersicum*, *Vitis vinifera*, and
22 *Zea mays* were also analyzed to demonstrate the broad applicability of this approach. Mapping-
23 based Genome Size Estimation (MGSE) and additional scripts are available on github:
24 <https://github.com/bpucker/MGSE>.

25

26 Introduction

27 Nearly all parts of the plant are now tractable to measure, but assessing the size of a plant
28 genome is still challenging. Although chromosome sizes can be measured under a microscope
29 [1], the combined length of all DNA molecules in a single cell is still unknown. Almost 20 years
30 after the release of the first *Arabidopsis thaliana* genome sequence, this holds even true for one
31 of the most important model species. Initially, biochemical methods like reassociation kinetics
32 [2], Feulgen photometry [3], quantitative gel blot hybridization [4], southern blotting [5], and flow
33 cytometry [6, 7] were applied. Unfortunately, these experimental methods rely on a reference
34 genome [8]. The rise of next generation sequencing technologies [9] enabled new approaches
35 based on k-mer profiles or the counting of unique k-mers [10, 11]. JellyFish [11], Kmergenie
36 [12], Tallymer [13], Kmerlight [14], and genomic character estimator (gce) [15] are dedicated
37 tools to analyze k-mers in reads. Next, genome sizes can be estimated based on unique k-mers
38 or a complete k-mer profile. Many assemblers like SOAPdenovo [16] and ALLPATHS-LG [17]
39 perform an internal estimation of the genome size to infer an expected assembly size. Recently,
40 dedicated tools for the genome size estimation like GenomeScope [18] and findGSE [19] were
41 developed. Although the authors considered and addressed a plethora of issues with real data
42 [18], results from different sequencing data sets for the same species can vary. While some
43 proportion of this variation can be attributed to accession-specific differences as described e.g.
44 for *A. thaliana* [19, 20], specific properties of a sequencing library might have an impact on the
45 estimated genome size. For example, high levels of bacterial or fungal contamination could bias
46 the result if not removed prior to the estimation process. Due to high accuracy requirements, k-
47 mer-based approaches are usually restricted to high quality short reads and cannot be applied
48 to long reads of third generation sequencing technologies. The rapid development of long read
49 sequencing technologies enables high contiguity assemblies for almost any species and is
50 therefore becoming the standard for genome sequencing projects [21, 22]. Nevertheless, some
51 highly repetitive regions of plant genomes like nucleolus organizing region (NOR) and
52 centromeres remain usually unassembled [20, 23, 24]. Therefore, the genome size cannot be
53 inferred directly from the assembly size, but the assembly size can be considered a lower
54 boundary when estimating genome sizes.

55 Extreme genome size estimates of *A. thaliana* for example 70 Mbp [2] or 211 Mbp [25] have
56 been proven to be inaccurate based on insights from recent assemblies [20, 24, 26–28].
57 However, various methods still predict genome sizes between 125 Mbp and 165 Mbp for diploid
58 *A. thaliana* accessions [26, 29–31]. Substantial technical variation is observed not only between

59 methods, but also between different labs or instruments [32]. As described above, extreme
60 examples for *A. thaliana* display 3 fold differences with respect to the estimated genome size.
61 Since no assembly is representing the complete genome, the true genome size remains
62 unknown. An empirical approach, i.e. running different tools and comparing the results, might be
63 a suitable strategy.

64 This work presents a method for the estimation of genome sizes based on the mapping of reads
65 to a high contiguity assembly. Mapping-based Genome Size Estimation (MGSE) is a Python
66 script which processes the coverage information of a read mapping and predicts the size of the
67 underlying genome. MGSE is an orthogonal approach to the existing tools for genome size
68 estimation with different challenges and advantages.

69

70

71 **Methods**

72 Data sets

73 Sequencing data sets of the *A. thaliana* accessions Columbia-0 (Col-0) [33–38] and
74 Niederzenz-1 (Nd-1) [31] as well as several *Beta vulgaris* accessions [39–41] were retrieved
75 from the Sequence Read Archive (AdditionalFile 1). Only the paired-end fraction of the two
76 included Nd-1 mate pair libraries was included in this analysis. Genome assembly versions
77 TAIR9 [42], AthNd-1_v1 [31], AthNd-1_v2 [24], and RefBeet v1.5 [39, 43] served as references
78 in the read mapping process. The *A. thaliana* assemblies, TAIR9 and Ath-Nd-1_v2, already
79 included plastome and chondrome sequences. These subgenome sequences of Ath-Nd-1_v2
80 were added to Ath-Nd-1_v1 as this assembly was previously cleaned of such sequences.
81 Plastome (KR230391.1, [44]) and chondrome (BA000009.3, [45]) sequences were added to
82 RefBeet v1.5 to allow proper placement of respective reads.

83 Genome sequences of *Brachypodium distachyon* strain Bd21 (GCF_000005505.3 [46]),
84 *Solanum lycopersicum* (GCA_002954035.1 [47]), *Vitis vinifera* cultivar Chardonnay
85 (QGNW01000001.1 [48]), and *Zea mays* cultivar DK105 (GCA_003709335.1 [49]) were
86 retrieved from the NCBI. Corresponding read data sets were retrieved from the Sequence Read
87 Archive (AdditionalFile1).

88

89 Genome size estimation

90 JellyFish2 v2.2.4 [11] was applied for the generation of k-mer profiles which were subjected to
91 GenomeScope [18]. Selected k-mer sizes ranged from 19 to 25. Results of different sequencing
92 data sets and different k-mer sizes per accession were compared. Genomic character estimator
93 (gce) [15] and findGSE [19] were applied to infer genome sizes from the k-mer histograms. If
94 tools failed to predict a value or if the prediction was extremely unlikely, values were masked to
95 allow meaningful comparison and accommodation in one figure. The number of displayed data
96 points is consequently a quality indicator.

97

98 Mapping-based genome size estimation

99 Despite some known biases [50–52], the underlying assumption of MGSE is a nearly random
100 fragmentation of the DNA and thus an equal distribution of sequencing reads over the complete
101 sequence. If the sequencing coverage per position (C) is known, the genome size (N) can be
102 calculated by dividing the total amount of sequenced bases (L) by the average coverage value:
103 $N = L / C$. Underrepresented repeats and other regions display a higher coverage, because
104 reads originating from different genomic positions are mapped to the same sequence. The
105 accurate identification of the average coverage is crucial for a precise genome size calculation.
106 Chloroplastic and mitochondrial sequences account for a substantial proportion of reads in
107 sequencing data sets, while contributing very little size compared to the nucleome. Therefore,
108 sequences with very high coverage values i.e. plastome and chondrome sequences are
109 included during the mapping phase to allow correct placement of reads, but are excluded from
110 MGSE. A user provided list of reference regions is used to calculate the median or mean
111 coverage based on all positions in these specified regions. Benchmarking Universal Single
112 Copy Orthologs (BUSCO) [53] can be deployed to identify such a set of *bona fide* single copy
113 genes which should serve as suitable regions for the average coverage calculation. Since
114 BUSCO is frequently applied to assess the completeness of a genome assembly, these files
115 might be already available to users. GFF files generated by BUSCO can be concatenated and
116 subjected to MGSE. As some BUSCOs might occur with more than one copy, MGSE provides
117 an option to reduce the predicted gene set to the actual single copy genes among all identified
118 BUSCOs.

119 BWA MEM v0.7 [54] was applied for the read mapping and MarkDuplicates (Picard tools v2.14)
120 [55] was used to filter out reads originating from PCR duplicates. Next, a previously described
121 Python script [56] was deployed to generate coverage files, which provide information about the
122 number of aligned sequencing reads covering each position of the reference sequence. Finally,
123 MGSE (<https://github.com/bpucker/MGSE>) was run on these coverage files to predict genome
124 sizes independently for each data set.

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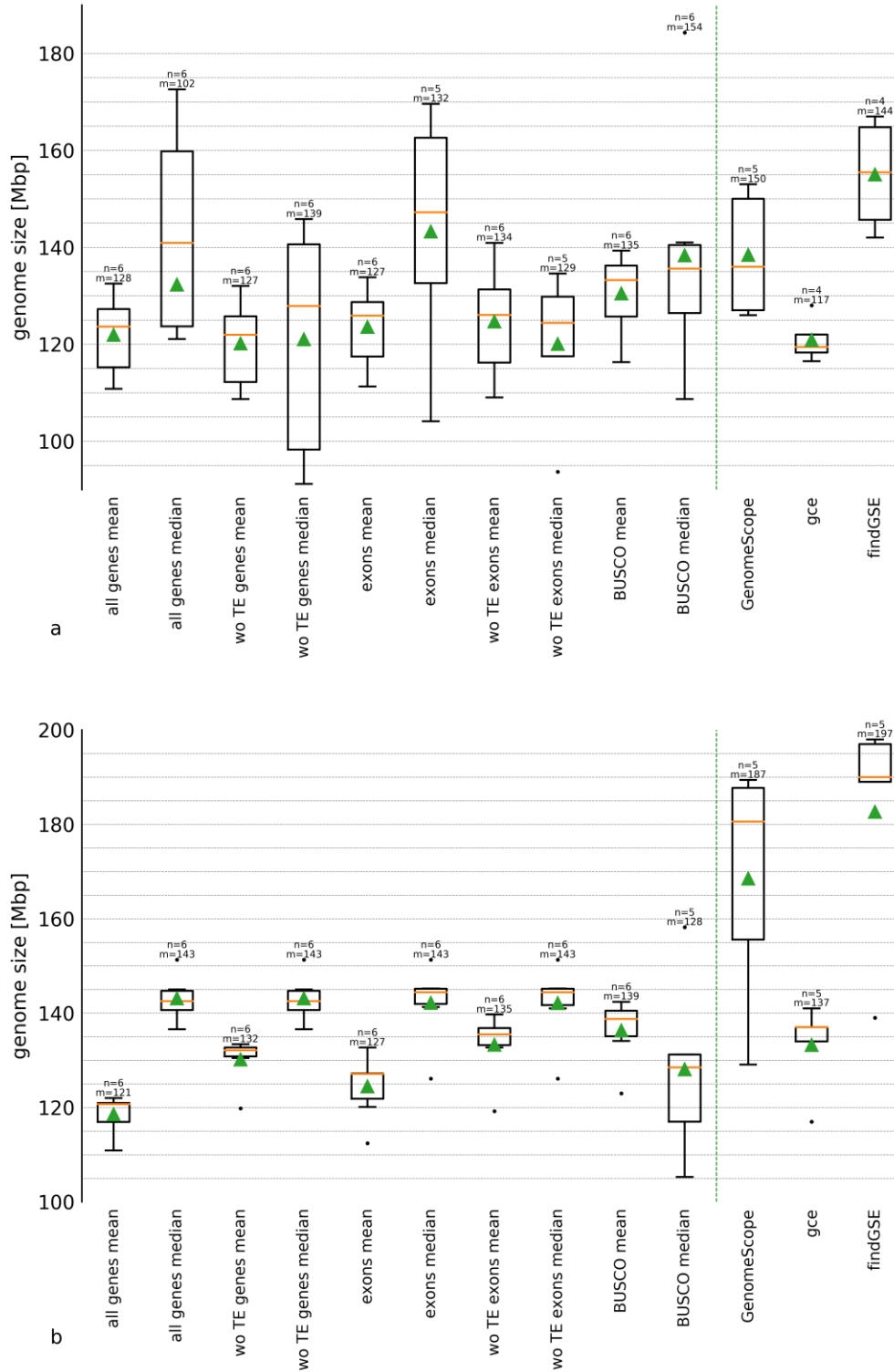
127 **Results & Discussion**

128 *Arabidopsis thaliana* genome size

129 MGSE was deployed to calculate the genome size of the two *A. thaliana* accessions Col-0 and
130 Nd-1 (Fig. 1). In order to identify the best reference region set for the average coverage
131 calculation, different reference region sets were tested. Manually selected single copy genes, all
132 protein encoding genes, all protein encoding genes without transposable element related genes,
133 only exons of these gene groups, and BUSCOs were evaluated (AdditionalFile2). The results
134 were compared against predictions from GenomeScope, gce, and findGSE for k-mer sizes 19,
135 21, 23, and 25.

136 Many estimations of the Col-0 genome size are below the assembly size of 120 Mbp [26] and
137 display substantial variation between samples (Fig. 1a). Due to low variation between different
138 samples and a likely average genome size the BUSCO-based approaches appeared promising.
139 GenomeScope predicted a similar genome size, while gce reported consistently much smaller
140 values. findGSE predicted on average a substantially larger genome size. Final sample sizes
141 below six indicated that prediction processes failed e.g. due to insufficient read numbers.

142 The variation among the estimated genome sizes of Nd-1 was smaller than the variation
143 between the Col-0 samples (Fig. 1). BUSCO-based estimations differed substantially between
144 mean and median with respect to the variation between samples (Fig. 1b). Therefore, the
145 average coverage is probably more reliably calculated via mean than via median. While gce
146 predicted as reasonable genome size for Nd-1, the average predictions by GenomeScope and
147 findGSE are very unlikely, as they contradict most estimations of *A. thaliana* genome sizes [6,
148 19, 24, 31].



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150

151 **Fig. 1: Comparison of *Arabidopsis thaliana* genome size estimations.**

152 Genome sizes of the *A. thaliana* accessions Col-0 (a) and Nd-1 (b) were predicted by MGSE,

153 GenomeScope, gce, and findGSE. Different MGSE approaches were evaluated differing by the set of

154 regions for the average coverage calculation (e.g. all genes) and the methods for the calculation of this
155 value (mean/median). Multiple read data sets (n) were analyzed by each tool/approach to infer an average
156 genome size given as median (m, yellow line) and mean (green triangles). transposable elements = TE,
157 without = wo.

158

159 The genome size estimation of about 139 Mbp inferred for Nd-1 through integration of all
160 analyses is slightly below previous estimations of about 146 Mbp [31]. Approximately 123.5 Mbp
161 are assembled into pseudochromosomes which do not contain complete NORs or centromeric
162 regions [24]. Based on the read coverage of the assembled 45S rDNA units, the NORs of Nd-1
163 are expected to account for approximately 2-4 Mbp [31]. Centromeric repeats which are only
164 partially represented in the genome assembly [24] account for up to 11 Mbp [31]. In summary,
165 the Nd-1 genome size is expected to be around 138-140 Mbp. The BUSCOs which occur
166 actually with a single copy in Ath-Nd1_v2 emerged as the best set of reference regions for
167 MGSE.

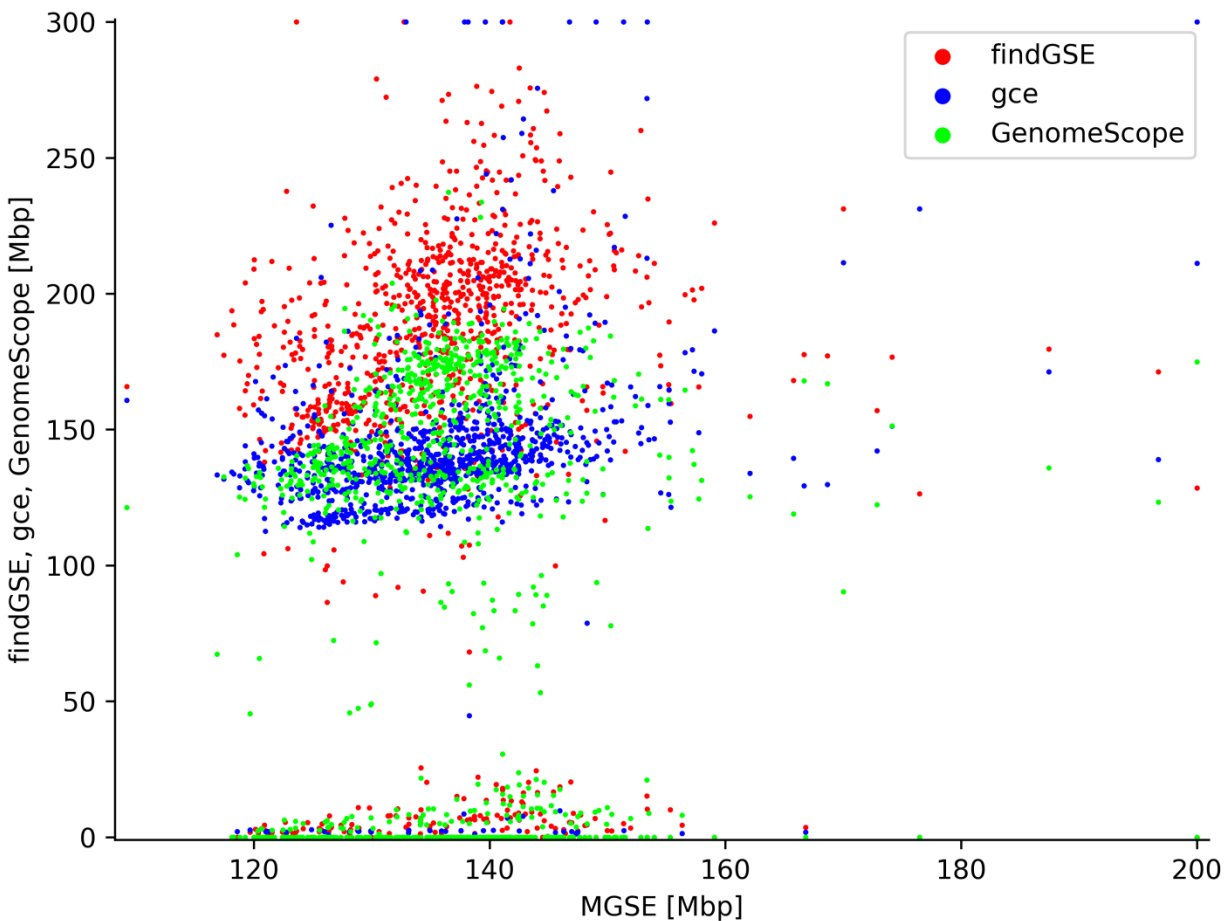
168 The relevance of very high assembly contiguity was assessed by comparing results of
169 AthNd-1_v1 (AdditionalFile3), which is based on short Illumina reads, to results of AthNd-1_v2
170 (AdditionalFile2), which is based on long Single Molecule Real Time sequencing (PacBio)
171 reads. The genome size predictions based on AthNd-1_v2 were substantially more accurate.
172 Reads are not mapped to the ends of contigs or scaffolds. This has only a minor influence on
173 large contigs, because a few small regions at the ends with lower coverage can be neglected.
174 However, the average coverage of smaller contigs might be biased as the relative contribution
175 of contig ends weights stronger. In addition, the representation of centromeric repeats and
176 transposable elements increases with higher assembly size and contiguity [24].

177 The feasibility of MGSE was further demonstrated by estimating the genome sizes of 1,028
178 *A. thaliana* accessions (Fig. 2, AdditionalFile4) which were analyzed by re-sequencing as part of
179 the 1001 genome project [57]. Most predictions by MGSE are between 120 Mbp and 160 Mbp,
180 while all other tools predict most genome sizes between 120 Mbp and 200 Mbp with some
181 outliers showing very small or very large genome sizes. MGSE differs from all three tools when
182 it comes to the number of failed or extremely low genome size predictions. All k-mer-based
183 approaches predicted genome sizes below 50 Mbp, which are most likely artifacts. This
184 comparison revealed systematic differences between findGSE, gce, and GenomeScope with
185 respect to the average predicted genome size. findGSE tends to predict larger genome sizes

186 than gce and GenomeScope. Very large genome sizes could have biological explanations like
187 polyploidization events.

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191 **Fig. 2: Genome size estimations of *Arabidopsis thaliana* accessions.**

192 MGSE, findGSE, gce, and GenomeScope were deployed to predict the genome sizes of 1,028 *A. thaliana*
193 accessions based on sequence read data sets (AdditionalFile4). Extreme outliers above 200 Mbp (MGSE)
194 or 300 Mbp (other tools) are displayed at the plot edge to allow accommodation of all data points with
195 sufficient resolution in the center.

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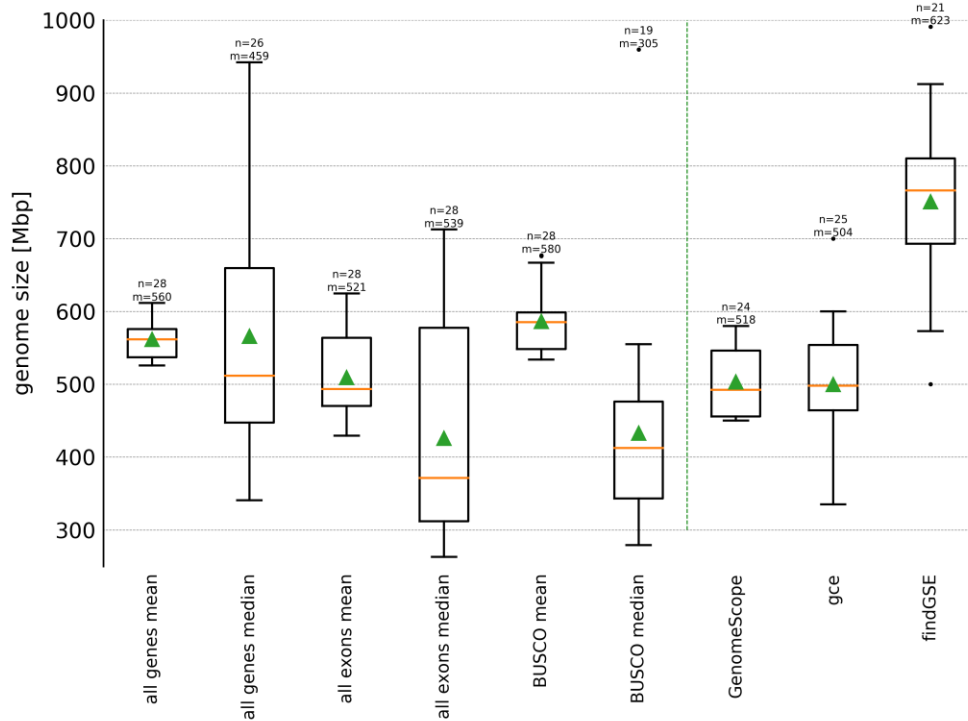
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199 *Beta vulgaris* genome size

200 Different sequencing data sets of *Beta vulgaris* were analyzed via MGSE, GenomeScope, gce,
201 and findGSE to assess the applicability to larger and more complex genomes (Fig. 3,
202 AdditionalFile5). Different cultivars served as material source for the generation of the analyzed
203 read data sets. Therefore, minor differences in the true genome size are expected. Moreover,
204 sequence differences like single nucleotide variants, small insertions and deletions, as well as
205 larger rearrangements could influence the outcome of this analysis. Since the current RefBeet
206 v1.5 assembly represents 567 Mbp [39, 43] of the genome, all estimations below this value can
207 be discarded as erroneous. Therefore, the mean-based approaches relying on all genes or just
208 the BUSCOs as reference region for the sequencing coverage estimation outperformed all other
209 approaches (Fig. 3). When comparing the *A. thaliana* and *B. vulgaris* analyses, the calculation
210 of an average coverage in all BUSCOs, which are actually present as a single copy in the
211 investigated genome, appears to be the most promising approach. While GenomeScope and
212 gce underestimate the genome size, the predictions by findGSE are extremely variable but
213 mostly around the previously estimated genome sizes [39, 43]. Based on results from the
214 *A. thaliana* investigation, the mean calculation among all single copy BUSCOs should be the
215 best approach. The prediction of slightly less than 600 Mbp is probably an underestimation, but
216 still the highest reliable estimate. When assuming centromere sizes of only 2-3 Mbp per
217 chromosome, this number could be in a plausible range. However, a previous investigation of
218 the repeat content indicates a larger genome size due to a high number of repeats which are
219 not represented in the assembly [58].

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222

223 **Fig. 3: Comparison of *Beta vulgaris* genome size estimations.**

224 The genome size of *B. vulgaris* was predicted by MGSE, GenomeScope, gce, and findGSE. Different
225 MGSE approaches were evaluated differing by the set of regions for the average coverage calculation
226 (e.g. all genes) and the methods for the calculation of this value (mean/median). Multiple read data sets
227 (n) were analyzed by each tool and approach to infer an average genome size given as median (m, yellow
228 line) and mean (green triangles).

229

230 Application to broad taxonomic range of species

231 After optimization of MGSE on *A. thaliana* (Rosids) and *B. vulgaris* (Caryophyllales), the tool
232 was deployed to analyze data sets of different taxonomic groups thus demonstrating broad
233 applicability. *Brachypodium distachyon* was selected as representative of grasses. *Solanum*
234 *lycopersicum* represents the Asterids, *Zea mays* was included as monocot species with high
235 transposable element content in the genome, and *Vitis vinifera* was selected due to a very high
236 heterozygosity. The predictions of MGSE are generally in the same range as the predictions
237 generated by GenomeScope, gce, and findGSE (AdditionalFile5, AdditionalFile6,
238 AdditionalFile7, AdditionalFile8, and AdditionalFile9). With an average prediction of 290 Mbp as

239 genome size of *B. distachyon*, the MGSE prediction is slightly exceeding the assembly size.
240 GenomeScope and gce predict genome sizes below the assembly size, while the prediction of
241 303 Mbp by findGSE is more reasonable. The *Z. mays* genome size is underestimated by all
242 four tools. However, MGSE outperforms GenomeScope and gce on the analyzed data set. The
243 *S. lycopersicum* genome size is underestimated by MGSE on most data sets. However, the
244 compared tools failed to predict a genome size for multiple read data sets. The highest MGSE
245 predictions are in the range of the expected genome size. MGSE failed for *V. vinifera* by
246 predicting only 50 Mbp. The high heterozygosity of this species could contribute to this by
247 causing lower mapping rates outside of important protein encoding genes i.e. BUSCO genes.

248

249 Considerations about performance and outlook

250 MGSE performs best on a high contiguity assembly and requires a (short) read mapping to this
251 assembly. Accurate coverage calculation for each position in the assembly is important and
252 contigs display artificially low coverage values towards the ends. This is caused by a reduction
253 in the number of possible ways reads can cover contig ends. The shorter a contig, the more is
254 the apparent coverage of this contig reduced. Since a read mapping is required as input, MGSE
255 might appear less convenient than classical k-mer-based approaches at first look. However,
256 these input files are already available for many plant species, because such mappings are part
257 of the assembly process [23, 24, 59, 60]. Future genome projects are likely to generate high
258 continuity assemblies and short read mappings in the polishing process.

259 One advantage of MGSE is the possibility to exclude reads originating from contaminating DNA
260 even if the proportion of such DNA is high. Unless reads from bacterial or fungal contaminations
261 were assembled and included in the reference sequence, the approach can handle such reads
262 without identifying them explicitly. This is achieved by discarding unmapped reads from the
263 genome size estimation. MGSE expects a high contiguity assembly and assumes all single copy
264 regions of the genome are resolved and all repeats are represented by at least one copy.
265 Although the amount of contamination reads is usually small, such reads are frequently
266 observed due to the high sensitivity of next generation sequencing [31, 61–64].

267 Reads originating from PCR duplicates could impact k-mer profiles and also predictions based
268 on these profiles if not filtered out. After reads are mapped to a reference sequence, read pairs
269 originating from PCR duplicates can be identified and removed based on identical start and end

270 positions as well as identical sequences. This results in the genome size prediction by GMSE
271 being independent of the library diversity. If the coverage is close to the read length or the
272 length of sequenced fragments, reads originating from PCR duplicates cannot be distinguished
273 from *bona fide* identical DNA fragments. Although MGSE results get more accurate with higher
274 coverage, after exceeding an optimal coverage the removal of apparent PCR duplicates could
275 become an issue. Thus, a substantially higher number of reads originating from PCR-free
276 libraries could be used if duplicate removal is omitted. Depending on the sequencing library
277 diversity completely skipping the PCR duplicate removal step might be an option for further
278 improvement. As long as these PCR duplicates are mapped equally across the genome, MGSE
279 can tolerate these artifacts.

280 All methods are affected by DNA of the plastome and chondrome integrated into the nuclear
281 chromosomes [65, 66]. K-mers originating from these sequences are probably ignored in many
282 k-mer-based approaches, because they appear to originate from the chondrome or plastome
283 i.e. k-mers occur with very high frequencies. The apparent coverage in the mapping-based
284 calculation is biased due to high numbers of reads which are erroneously mapped to these
285 sequences instead of the plastome or chondrome sequence.

286 Differences in the GC content of genomic regions were previously reported to have an impact
287 on the sequencing coverage [67, 68]. Both, extremely GC-rich and AT-rich fragments,
288 respectively, are underrepresented in the sequencing output mainly due to biases introduced by
289 PCR [69, 70]. Sophisticated methods were developed to correct coverage values based on the
290 GC content of the underlying sequence [70–72]. The GC content of genes selected as reference
291 regions for the coverage estimation is likely to be above the 36.3% average GC content of
292 plants [56]. This becomes worse when only exons are selected due to the even higher
293 proportion of coding sequence. Although a species specific codon usage can lead to some
294 variation, constraints of the genetic code determine a GC content of approximately 50% in
295 coding regions. The selection of a large set of reference regions with a GC content close to the
296 expected overall GC content of a genome would be ideal. However, the overall GC content is
297 unknown and cannot be inferred from the reads due to the above mentioned sequencing bias.
298 As a result, the average sequencing coverage could be overestimated leading to an
299 underestimation of the genome size. Future investigations are necessary to develop a
300 correction factor for this GC bias of reads.

301 Many plant genomes pose an additional challenge due to recent polyploidy or high
302 heterozygosity. Once high contiguity long read assemblies become available for these complex
303 genomes, a mapping based approach is feasible. As long as the different haplophases are
304 properly resolved, the assessment of coverage values should reveal a good estimation of the
305 genome size. Even the genomes of species which have recently undergone polyploidization
306 could be investigated with moderate adjustments to the workflow. Reference regions need to be
307 selected to reflect the degree of ploidy in their copy number.

308 The major issue when developing tools for the genome size prediction is the absence of a gold
309 standard. Since as of yet there is no completely sequenced plant genome, benchmarking with
310 real data cannot be perfect. As a result, how various estimation approaches will compare to the
311 first completely sequenced and assembled genome remains speculative. Although not
312 evaluated in this study, we envision that MGSE could be generally applied to all species and is
313 not restricted to plants.

314

315 **Data availability**

316 Scripts developed as part of this work are freely available on github:
317 <https://github.com/bpucker/MGSE> (<https://doi.org/10.5281/zenodo.2636733>). Underlying data
318 sets are publicly available at the NCBI and SRA, respectively.

319

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324

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- 489 **Supplements:**
- 490 AdditionalFile1: Sequencing data set overview.
- 491 AdditionalFile2: *A. thaliana* genome size prediction values for all different approaches.
- 492 AdditionalFile3: *A. thaliana* genome size prediction based on Ath-Nd1_v1.
- 493 AdditionalFile4: *A. thaliana* genome size predictions by MGSE, findGSE, gce, and
494 GenomeScope.

495 AdditionalFile5: *B. vulgaris*, *Zea mays*, *Brachypodium distachyon*, *Solanum lycopersicum*, and
496 *Vitis vinifera* genome size prediction values for all different approaches.

497 AdditionalFile6: Genome size estimation of *Brachypodium distachyon*.

498 AdditionalFile7: Genome size estimation of *Zea mays*.

499 AdditionalFile8: Genome size estimation of *Solanum lycopersicum*.

500 AdditionalFile9: Genome size estimation of *Vitis vinifera*.

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