# 1 Title

# 2 Genome improvement and genetic map construction for Aethionema arabicum,

# 3 the first divergent branch in the Brassicaceae family

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### 25 Abstract

# 26 Background

The genus *Aethionema* is a sister-group to the core-group of the Brassicaceae family that includes *Arabidopsis thaliana* and the Brassica crops. Thus, *Aethionema* is phylogenetically well-placed for the investigation and understanding of genome and trait evolution across the family. We aimed to improve the quality of the reference genome draft version of the annual species *Aethionema arabicum*. Secondly, we constructed the first *Ae. arabicum* genetic map. The improved reference genome and genetic map enabled the development of each other.

### 34 **Results**

We started with the initially published genome (version 2.5). PacBio and MinION 35 sequencing together with genetic map v2.5 were incorporated to produce the new 36 reference genome v3.0. The improved genome contains 203 MB of sequence, with 37 approximately 94% of the assembly made up of called bases, assembled into 2,883 38 scaffolds. The N<sub>50</sub> (10.3 MB) represents an 80-fold over the initial genome release. 39 We generated a Recombinant Inbred Line (RIL) population that was derived from two 40 ecotypes: Cyprus and Turkey (the reference genotype. Using a Genotyping by 41 Sequencing (GBS) approach, we generated a high-density genetic map with 749 (v2.5) 42 and then 632 SNPs (v3.0) was generated. The genetic map and reference genome 43 were integrated, thus greatly improving the scaffolding of the reference genome into 44 11 linkage groups. 45

#### 46 **Conclusions**

We show that long-read sequencing data and genetics are complementary, resulting in an improved genome assembly in *Ae. arabicum*. They will facilitate comparative

- 49 genetic mapping work for the Brassicaceae family and are also valuable resources to
- 50 investigate wide range of life history traits in Aethionema.
- 51 Keywords
- 52 Aethionema arabicum, Brassicaceae, genome improvement, genetic map, PacBio,
- 53 MinION, Genotyping by Sequencing.

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### 73 Background

The genus Aethionema belongs to the important plant family Brassicaceae. The 74 crucifers contain many species of interest, such as the Brassica crop plants (e.g. B. 75 rapa, B. oleracea and B. napus), ornamental plants (such as the genera Aubrieta, 76 Iberis, Lunaria and Draba) and research model plant species (including Arabidopsis 77 thaliana, A. lyrata, Capsella rubella and Arabis alpina). Phylogenetic studies have 78 established Aethionema as the sister-group of the core-group in the family [1]. Thus, 79 Aethionema holds an essential phylogenetic position for studies on genome and trait 80 evolution across the Brassicaceae family. 81

The monogeneric tribe *Aethionemeae* consists of 57 species and is distributed mainly 82 in the Irano-Turanian region, a hot spot for species radiation and speciation [2-4]. This 83 tribe displays various interesting morphological and ecological characteristics, 84 especially fruit and seed heteromorphism. Heteromorphism is defined as the 85 production of two or more distinct fruit or seed morphs on the same individual [5], which 86 includes morphological size, shape and color; physiological dormancy and germination 87 of fruits and seeds. Aethionema arabicum is one of the seven reported heteromorphic 88 species of Aethionema [6, 7]. Aethionema arabicum is a small diploid annual, with a 89 short life cycle starting from seed germination to the end of the vegetative development 90 in spring, followed by reproduction and the end of life cycle in summer [8]. Both annual 91 life history and heteromorphism probably evolved as adaptive responses to 92 unpredictable environments, especially dry arid habitats, indicating a wide range of 93 natural variation for ecologically adaptive traits in Ae. arabicum. 94

Owing to its unique phylogenetic position and interesting characteristics, *Ae. arabicum* is an ideal sister-group model for research. Therefore, *Aethionema* genome and
 genetic resources are desirable. The initially published *Ae. arabicum* draft genome

(v1.0) contains 59,101 scaffolds with an N50 of 115,195 bp while the genome was 98 predicted to be 320 Mbp in size with n=11 [9]. Here we first aimed to (i) improve the 99 quality of the reference genome and (ii) to construct the first Ae. arabicum genetic map. 100 A higher quality version of the genome assembled by the VEGI consortium was later 101 released as version 2.5, which is used as the starting point of our analyses. 102 High throughput sequencing using Pacific Biosciences (PacBio) and Oxford Nanopore 103 MinION (MinION) technology followed and resolved many uncalled gaps in the v2.5 104 genome and supported further super-scaffolding, which resulted in genome v3.0. 105 The genetic map was constructed using Genotyping by Sequencing (GBS) on a 106 Recombinant Inbred Line (RIL) population. The 216 RILs were derived from a cross 107 between Turkey (reference ecotype) and Cyprus ecotypes. The first version of genetic 108 map v2.5 was obtained based on genome v2.5 with 746 Single Nucleotide 109 Polymorphism (SNP) markers. The later genetic map v3.0 was built with 626 SNPs 110 generated based on genome v3.0. 111

Here we show that the long-read genome assembly and the genetic map of *Ae. arabicum* supported the development of each other. They will serve as a substantial resource for further research on *Aethionema* as well as the Brassicaceae family.

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### 120 Data description and Methods

## 121 **Overview of the workflow**

An overview of the improvements of the genome of Aethionema arabicum and the 122 generation and improvement of its genetic map are depicted in Figure 1. The genome 123 draft version 1.0 was first improved by Ray [10] and AllPathsLGs [11] and led to the 124 release of genome v2.5 (available on genomevolution.org). Genome v2.5 was used as 125 a basis for SNP calling after GBS of the RILs. This generated SNP markers used to 126 construct the genetic map v2.5. Scaffolds were ordered with AllMaps [12] based on the 127 maximum co-linearity to genome v2.5 and genetic map v2.5. This resulted in genome 128 vAM. Gap filling and super-scaffolding improvement for genome vAM was obtained by 129 PacBio sequencing leading to genome v2.6. PBjelly2 [13] run using the MinION reads 130 further improved genome v2.6 to v3.0. We revisited the genetic map v2.5 by recalling 131 SNPs according to genome v3.0 and constructed a genetic map v3.0 with the newly 132 called SNP markers. Below we describe the workflow in detail in the three following 133 sections: (i) the initial genome assembly, (ii) genetic map construction and (iii) genome 134 improvement. 135

- (i) The intial genome (v2.5): The starting point
- 137 Genome re-assembly using AllPathsLG

The version 1.0 assembly generated by the Ray assembler [10] was fragmented *in silico* into a set of artificial overlapping reads, combined with paired end and mate pair data (described in [9]), and re-assembled using the AllPathsLG assembler [11]. Gap closing was then performed using GapCloser, part of the SOAPdenovo2 package [14]. Gene annotations were lifted over from assembly version 1.0 to version 2.5 using the LiftOver tool from the UCSC Genome Browser tools package [15].

Genome version 2.5 contains 3,166 scaffolds, has an N50 of 564,741 bp and was published as version 2.5 on <u>https://genomevolution.org/coge/</u>.

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### 147 (ii) Genetic map construction

#### 148 Plant material

Two Aethionema arabicum ecotypes were used, Turkey (TUR) and Cyprus (CYP). The
TUR accession comes from the living plant collections at the Botanical Garden in Jena,
Germany (Botanischer Garten Jena). The seeds for this genotype were derived from
a plant in the Botanical Garden in Nancy, France. The CYP ecotype was collected in
2010 near Kato Moni (coordinates UTM WGS 84: 508374 - 3879403) at an altitude
410 m on pillow lava by Charalambos S Christodoulou [16].

These two ecotypes were used as parents for the development of the recombinant inbred line population, where TUR was the father and CYP the mother. Seeds from initial  $F_1$  plants were used to generate an initial  $F_2$  population. For each of the 216 segregating  $F_2$  plants, a single seed was randomly chosen to further grow and reproduce the next generation. The procedure was repeated until  $F_8$ , when the experiment was performed with 216 RILs.

To grow the plants for the GBS, F<sub>8</sub> seeds of 216 RILs were placed on filter paper, wetted with distilled water, in petri dishes. Imbibed seeds were incubated at 4°C in dark for 3 days, followed by germination in the light at 20°C for 2 days. Seedlings were transferred to soil pots (10.5 cm diameter 10 cm height) in November 2014. Plants were grown in greenhouse (Wageningen University and Research, the Netherlands) in partially controlled conditions, long day (16 h light and 8 h dark) and at 20°C.

#### 167 Genotyping By Sequencing (GBS)

168 DNA isolation

Young tissues from leaves and flower buds were collected from each F9 plant for DNA 169 isolation. The DNA isolation was done according to a modified CTAB protocol [17]. In 170 brief, plant material was frozen with liquid nitrogen and ground into powder. Each 171 sample was incubated with 500 µl of CTAB buffer at 65°C in the water bath for 30 min. 172 cooling at room temperature, equal volume (500 µl) After 30 min 173 of chloroform: isoamylalcohol (24:1 v/v) was added, and vigorously hand-mixed for a min. 174 400 µl of supernatant was recovered after centrifuging at maximum speed for 5 min. 175 The supernatant was cleaned again with a chloroform: isoamylalcohol step. DNA 176 precipitation was performed by adding an equal volume of cold isopropanol with 30 177 min incubation on ice and centrifugation at maximum speed for 15 min. The DNA pellet 178 was cleaned twice with 1 ml of 70% ethanol and centrifugation at maximum speed for 179 5 min. Dry DNA pellet was dissolved in Milli-Q water. 180

181 Constructing GBS libraries

DNA was treated with RNAse overnight at 37°C with RNAse one by Promega. Quality 182 was checked on a 1% agarose gel and DNA quantity was checked with Pico Green. 183 Based on this, DNA was diluted down to 20 ng/µl with MQ water and used in further 184 analysis. GBS was performed in general by following the procedure described in [18]. 185 Oligonucleotides for creation of common as well as 96 barcoded ApeKI adapters were 186 obtained from Integrated DNA Technologies and diluted to 200 µM. For each barcoded 187 and common adapter, top and bottom strand oligos were combined to a 50 µM 188 annealing molarity in TE to 100 µl total volume. Adapter annealing was carried out in 189 a thermocycler (Applied Biosystems) at 95°C for 2 min, ramp to 25°C by 0.1 degree 190 per second, hold at 25°C for 30 min and 4°C forever. Annealed adapters were further 191 diluted to a 0.6 ng/µl concentrated working stock of combined barcoded and common 192 adapter in 96 well microtiter plate and dried using a vacuum oven. For each genomic 193

<sup>194</sup> DNA sample 100 ng (10 ng/µl) was used and added to lyophilized adapter mix and <sup>195</sup> dried down again using a vacuum oven.

Adapter DNA mixtures were digested using 2.5 Units ApeKI (New England Biolabs) for 196 2 hours at 75°C in a 20 µl volume. Digested DNA and Adapters were used in 197 subsequent ligation by 1.6 µl (400 Units/µl) T4DNA Ligase in a 50 µl reaction volume 198 at 22°C for one hour followed by heat inactivation at 65°C for 30 min. Sets of 96 199 digested DNA samples, each with a different barcode adapter, were combined (10µl 200 each) and purified using a Qiaquick PCR Purification columns (Qiagen). Purified 201 pooled DNA samples were eluted in a final volume of 10µl. DNA Fragments were 202 amplified in 50 µl volume reactions containing 2 µl pooled DNA, 25 µl KAPA HiFi 203 HotStart Master Mix (Kapa Biosystems), and 2 µl of both PCR primers (12.5 µM). PCR 204 cycling consisted of 98°C for 30 seconds, followed by 18 cycles of 98°C for 30 seconds, 205 65°C for 30 seconds, 72°C for 30 seconds with a final extension for 5 minutes and kept 206 at 4°C. Amplified libraries were purified as above but eluted in 30 µl. Of the amplified 207 libraries 1 µl was loaded onto a Bioanalyzer High Sensitivity DNA Chip (Agilent 208 technologies) for evaluation of fragment sizes and 1 µl was used for quantification 209 using Qubit (Life Technologies). Amplified library products were used for extra size 210 selection using 2% agarose gel cassette on a blue pippin system (Sage Science) to 211 remove fragments smaller than 300 bp. Eluted size selected libraries were purified by 212 AmpureXP beads (Agencourt). Final libraries were used for clustering on five lanes of 213 an illumina Paired End flowcell using a cBot. Sequencing used an illumina HiSeg2000 214 instrument using 2\*100 nt Paired End reads. 215

216 Sequencing and processing raw GBS data

Raw sequencing data was processed using the TASSEL software package [19]
 version 5.2.37 using the GBSv2 pipeline. For quality filtering and barcode trimming,

the GBSSeqToTagDBPlugin was run with the following parameters: kmerLength: 64, 219 minKmerL: 20, mnQs: 20, mxKmerNum 100000000. Tags were dumped from the 220 produced database using TagExportToFastgPlugin and mapped to the reference 221 genome using the bwa software package [20] in single-ended mode (samse). 222 Positional information from aligned SAM files was stored in the TASSEL database 223 using the SAMToGBSdbPlugin. The DiscoverySNPCallerPlugin was run using the 224 following parameters: mnLCov: 0.1, mnMAF: 0.01. Found SNPs were scored for 225 quality using SNPQualityProfilerPlugin and the Average taxon read depth at SNP was 226 used as a quality score for filtering in the next step (minPosQS parameter), these 227 scores were written to the TASSEL database using UpdateSNPPositionQualityPlugin. 228 Finally, the ProductionSNPCallerPluginV2 was run with the following parameters: Avg 229 Seq Error Rate: 0.002, minPosQS: 10, mnQS: 20. 230

231 Genetic map calculation

We used JoinMap v4.1 for the genetic map construction [21, 22]. The genetic map v2.5 was built with 749 SNPs generated by GBS based on genome v2.5 (unprocessed and processed data available as S1 and S2). A set of 632 SNPs called according to genome 3.0 was used for the genetic map v3.0 (unprocessed and processed data available as S3 and S4). Regression and Maximum likelihood mapping were used to calculate these maps (the linkage group information for both 2.5 and 3.0 genetic maps are available as S5).

#### 239 (iii) Genome improvement

#### 240 Genome version vAM: AllMaps

We ran AllMaps [12] with default setting to combine genetic map v2.5 and physical map genome v2.5. This step resulted in genome vAM, in which scaffolds were ordered and oriented to reconstruct chromosomes.

### 244 Contamination removal

The Ae. arabicum scaffolds v2.5 were checked for contaminations. The genome 245 scaffolds were split into 197,702 1 kbp fragments and blasted against the NCBI nt 246 database [23] using Tera-BLAST (TimeLogic® Tera-BLAST™ algorithm, Active Motif 247 Inc., Carlsbad, CA). The output was then analyzed by MEGAN 6 [24]. All scaffolds for 248 which more than 50% of their entire length was found in bacteria and with no hit in 249 Viridiplantae were marked as putatively contaminated. A hit was counted with a 250 minimum bit score of 50. Additionally, Bisulfite sequencing (BS-seq) CpG and 251 Chromatin ImmunoPrecipitation DNA-Sequencing (ChIP-seq) H3 data were checked 252 to identify contaminants (Aethionema contamination.xlsx). We ensured that these 253 scaffolds were not combined with another scaffold by PBjelly2. After screening, three 254 v2.5 scaffolds were removed: Scaffold 2406, Scaffold 2454 and Scaffold 2594. They 255 had a total length of 1,758 bp without any annotated genes. A summary of the 256 contamination screen is available as S6. 257

## 258 Long read generation for genome improvement

259 PacBio reads

Genomic DNA (gDNA) for *Ae. arabicum* was obtained from leaves of the Cyprus and Turkey ecotypes. DNA was extracted using a modified protocol [25] based on [26]. For the Turkey ecotype 35.70 µg and for the Cyprus ecotype 21.45 µg high molecular weight DNA were sent to the Max Planck-Genome-Centre, Cologne, and sequenced using the PacBio RS II machine (library insert size was 10-15 kbp gDNA). Four flow cells for Cyprus and six for Turkey were sequenced. Table 1 summarizes the statistics of the reads. The CG content of the pooled reads was 38%.

### **Table 1: Overview of the Aethionema arabicum PacBio reads**

Total reads	Cyprus	Turkey	
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Number of reads	381,069	152,415	228,654
Length variation	11 - 57,910	11 - 55,919	11 - 57,910
Average length	5,845.12	5,794.88	5,878.8
Average quality	10.48	10.13	10.72

<sup>268</sup> The lengths are given in nucleotides and the quality as phred score.

269 MinION reads

To obtain MinION reads, gDNA was extracted from the Turkey ecotype (four leaf 270 samples, 100 mg each) as outlined above. After pooling the samples, the gDNA 271 concentration was measured using Hoechst 33258 DNA dye and resulted in 73.85 272 ng/µl. The library preparation was done using the Oxford Nanopore SQK-NSK007 273 protocol and R9.4 chemistry to design a 8 kbp 2D library. The sequencing run was 274 carried out using Oxford Nanopores MinION technology. The flow cell sequenced 275 30,935 reads (122,362,072 nt) at -205 mV and 24 hours of runtime. After base calling 276 with the MinKNOW 1.6 software (Oxford Nanopore Technologies Ltd.) the read length 277 ranged from 5 to 63,441 nt with an average length of 3,955 nt. The average phred 278 quality score was 11 and the GC content 41%, reads were not filtered or trimmed. The 279 initial sequence format FAST5 was converted to FASTQ format by using the R package 280 poRe version 0.21 [27]. Because the MinION flow cell had previously been used for 281 Physcomitrella patens DNA in the same run, the 30,935 reads were filtered for putative 282 P. patens contamination. The reads were mapped with the long read mapper GMAP 283 version 2017-08-15 [28] against the P. patens genome V3 [29]. All settings were kept 284 at default. 1,447 reads were characterized as putative *P. patens* reads and therefore 285 removed. 286

### 287 Genome improvement using long reads

To perform super-scaffolding and gap filling, the program PBjelly 2 version 15.8.24 was used [13]. It internally uses BLASR v5.1 [30] for mapping reads to a reference. The BLASR parameters internally used for mapping were: "-minMatch 12 -bestn 1 noSplitSubreads".

### 292 Genome version 2.6: PacBio sequencing incorporation

We ran PBjelly2 with 381,022 (152,398 CYP, 228,624 TUR) PacBio reads which where head-cropped with 20 (due to suspicious per base sequence content suggestion presence of adapters) using Trimmomatic version 0.36 [31].

PBjelly2 was used to improve genome v2.5 and vAM. Comparing the results, we found 296 some scaffold connections which were made by PBjelly2 (v2.5) were no longer 297 possible for vAM (these scaffolds were already connected). Five connections formed 298 for v2.5 scaffolds were already introduced by the genetic map approach (see above). 299 Twelve connections which could be established in v2.5 were not formed in the PBjelly2 300 output for improving vAM, because the scaffolds were already connected with other 301 scaffolds. Since PBjelly2 only fills gaps with reads and is not able to place whole 302 scaffolds in gaps, it was necessary to split the vAM genome at certain points to be able 303 to obtain the twelve connections which were not present in the PBjelly2 output for vAM 304 (visualized in Figure 2). Split scaffolds were reconnected again after running PBjelly2, 305 using N-stretches of length 100 to keep all improvements introduced in vAM if they 306 were not formed by PBjelly2 (scaffolds in the vAM genome were combined using 307 stretches of 100 Ns to denote a gap of unknown length). Since it is possible that 308 PBjelly2 only fills a gap partially, we had to identify the positions of the gaps introduced 309 by AllMaps in the new genome version and checked if they were filled completely or 310

not. If the gap length was reduced, it was extended to have a length of 100 again. This
 approach produced genome v2.6.

#### 313 Genome version 3.0: MinION sequencing incorporation

After improving the genome to v2.6 using the PacBio reads, the same approach was

applied for 30,935 MinION TUR reads to obtain the Ae. arabicum genome v3.0. The

316 MinION reads were also checked for contamination. The genome version 3.0 is

available at https://genomevolution.org/coge/GenomeView.pl?gid=36061.

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### 319 Name convention of Ae. arabicum v3.0 genome scaffolds

Scaffolds of genome v3.0 were named and ordered according to their length from long 320 to short. The longest eleven scaffolds were named linkage group (LG) based on the 321 genetic map. Scaffolds which were combined are named csc for concatenated scaffold 322 and the other ones are named sc (scaffold). The v3.0 scaffold names therefore follow 323 the scheme type-number v2.5 scaffold[.v2.5 scaffold...]. I.e., the scaffold type (LG, 324 CSC, SC), followed by a minus and the number of the scaffold, separated by a blank, 325 followed a list of scaffolds denoting the v2.5 scaffolds or the v3.0 scaffolds. This naming 326 system resulted in a shift in LG order between v2.5 and v3.0 (Supplementary file 327 linkage group map.xlsx). 328

## 329 Migration of proteins to new genome version

To perform the lift over of the gene models from v2.5 to v3.0, a combination of Gene Model Mapper (GeMoMa) v1.4 [32] and flo (flo - same species annotations lift over pipeline, https://github.com/wurmlab/flo) were used. The results of both programs were concatenated. flo results were preferred over GeMoMa results if the results of the two programs differed, because flo works with alignments on nucleotide level while GeMoMa works with blasting proteins on amino acid level. If a protein could not be lifted completely, it is marked as partial in the resulting GFF (v3.0). A total of 34 genes

had to be lifted manually, because they were either not lifted at all or only partially. If 337 an intron could not be lifted, it was added by hand. If an exon or CDS could not be 338 lifted, the new location was deriving from neighboring features which could be migrated 339 to the new genome version. The location was then used to extract the nucleotide 340 sequence from the genome using samtools v1.4 [33]. Only if the sequence was 341 identical to the original sequence extracted from v2.5, the feature was migrated. This 342 check was performed with ClustalW v2.1 [34]. After the migration step, the GFF file 343 was checked and corrected. Genes which did not contain a start or a stop, contained 344 internal stops or whose CDS sequence had a length not dividable by 3 were marked 345 as potential pseudogenes with "pseudo=true". To check if a gene contains internal 346 stops each of its CDS features was checked individually for having at least one frame 347 which results in no stop codons. Genes which were identical to other genes (start and 348 stop position are equal) or were contained in other genes were removed. If the 3' CDS 349 of a gene did not contain a stop codon but could be added by extending the CDS by 350 three nucleotides, the CDS was corrected. The lifted genes were classified as shown 351 inTable 2. 352

353 Table 2: Overview of gene liftover: GFF migration statistics

Lifted only by flo	4,346
Lifted only by GeMoMa	36
Lifted with both programs	18,177
Manually lifted	34
Partially lifted	14
Number of corrected CDS	10,259
Marked as pseudo	3,230

Most genes could be lifted by flo and GeMoMa. The reason why flo was able to lift more proteins is that GeMoMa works with protein sequences and the program was not able to generate proteins for 20,056 CDS features, either because a gene did not possess a CDS or because of faulty CDS sequences.

### 358 Name convention of v3.0 genes

Old gene IDs were kept in the note attribute of the genes in the GFF and the linkage 359 group numbers of the genetic map are also noted. The names of the genes were 360 changed into Aa3typeNumberGenenumber: Aa for Aethionema arabicum, indicator 361 genome version 3, followed by the type of scaffold, its number and the number of the 362 gene (starting with 1 at the 5' end), e.g. Aa3LG1G2 or Aa3SC2601G1). For transcript 363 isoforms (splice variants) this locus nomenclature can be extended by the number of 364 the isoform (.X). Version 3.0 of the genome and all gene models are available at 365 https://genomevolution.org/coge/GenomeView.pl?gid=36061. 366

## 367 Data Availability Statement

- The genome version 2.5 is available at:
- https://genomevolution.org/coge/GenomeInfo.pl?gid=23428.
- The genome version 3.0 is available at:
- https://genomevolution.org/coge/GenomeView.pl?gid=36061.
- 372 For Review: the data (as mapped BAM files) have been made available in CoGe.
- 373 Account: agr\_reviewer Password: GoCoGe'!
- 374 <u>https://auth.iplantcollaborative.org/cas4/login?service=https://genomevolution.org/coge/index.pl</u>
- 375 https://genomevolution.org/coge/GenomeInfo.pl?gid=36061
- The GBS unprocessed and processed reads for genome mapping 2.5 and 3.0 are
- available as supplmental files S1-S4. The linkage group information for both 2.5 and
- 378 3.0 genetic maps are available as S5. A summary of the contamination screen is
- available as S6.

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## 383 Results and Discussion

### **Reference genome improvement**

The published draft version of the Ae. arabicum genome utilized the Ray assembler 385 and contained 59,101 scaffolds with an N50 of 115,195bp [9]. Reassembly using the 386 the AllPathsLG assembler and gap-closing using the SOAPdenovo GapCloser tool 387 were used as a starting point for super-scaffolding. This resulted in a reassembly with 388 3,166 scaffolds, and a scaffold  $N_{50}$  of 564,741bp labeled and published as version 2.5 389 on https://genomevolution.org/coge/. The subsequent genome versions (vAM, v2.6 390 and v3.0) were obtained using linkage map and long read correction. The quality 391 improvement of the genome is presented as the increase in total number of bases, 392 reduced number of scaffolds and number of gaps, as well as bigger N<sub>50</sub> and smaller 393 L<sub>50</sub> parameters (Table 3). In comparison with the starting genome v2.5, the final 394 genome v3.0 has 9% less scaffolds (from 3,166 to 2,883). The overall length of 395 genome v3.0 was extended from 196,005,095 to 203,449,326 bases (17% more) and 396 the number of uncalled based was reduced from 25,768,296 to 13,790,434 (from 397 13.2% to 6.8%) (Table 3). 398

**Table 3**: Statistic overview of *Aethionema arabicum* genome versions

Genome version	Draft	v2.5	vAM	v2.6	v3.0
# Bases		196,005,095	196,022,695	203,150,143	203,449,326
# Scaffolds	59,101	3,166	2,990	2,895	2,883
# Scaffolds containing Ns		1,910	1,734	1,542	1,539
# Ns		25,768,296	25,785,896	13,946,922	13,790,434

N50	115,195	564,741	10,141,718	10,328,388	10,328,388
L50		56	9	9	9

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#### 401 Genome improvement using long reads

### 402 Read mapping efficiency

The results of the mapping of the reads to the genome using PBjelly2 are summarized 403 in Table 4. Almost the same number of PBjelly2 reads were mapped to genome v2.5 404 and vAM. However, it was important to apply PBjelly2 on both genomes in order to find 405 scaffold connections which were not possible due to a combination of certain scaffolds 406 in vAM (see supplementary file 407 combination comparison pbjelly for v2.5 vs pbjelly for vGM.ods for details). The 408 genome v2.6 resulted by improving the split vAM genome using PBjelly2, reconnecting 409 scaffolds and resizing gaps if needed. We also compared the results for improving v2.5 410 with vAM, but there were no new scaffold connections which were missed by improving 411 the v2.6 version, so we did not perform a split step for improving the genome using the 412 MinION reads. The mapping efficiency for the MinION reads is lower than for the 413 PacBio reads due to a contamination of the reads (see Methods for details). There are 414 5.9% more MinION reads which were mapped to v2.6 than v2.5, demonstrating that 415 the changes done to the genome are supported by the very long reads. 416

## 417 Table 4: Mapping efficiency of PBjelly2's mapping step.

The percentages in brackets give the percentage of the total number of reads (CYP, TUR or CYP + TUR). The line "# covered scaffolds" gives the number of scaffolds in which at least one read was mapped. Here, the number in brackets gives the percentage of the total number of scaffolds.

Setup	PacBio vs. v2.5	PacBio vs.	MinION vs.	MinION vs.
		vAM	v2.5	v2.6
# mapped TUR	198,675	198,629	14,098 (45.6%)	15,886 (51.4%)
reads	(86.9%)	(86.9%)		
# mapped CYP	131,976	131,942	-	-
reads	(86.6%)	(86.6%)		
# total reads	330,651	330,571	14,098 (45.6%)	15,886 (51.4%)
mapped	(86.8%)	(86.8%)		
# unmapped	50,371 (13.2%)	50,451 (13.2%)	16,837 (54.4%)	15,049 (48.6%)
reads				
# scaffolds	3,166	2,990	3,166	2,895
input genome				
# covered	2,971 (93.8%)	2.804 (93.8%)	1.689 (53.3%)	1.429 (49.4%)
scaffolds				

423 The effect of PBjelly2 runs applied to the different assembly versions

Most scaffolds were combined in the vAM which resulted in the L<sub>50</sub> valuelowered from 424 56 to 9 and the N<sub>50</sub> value almost doubled. PBjelly2 was not as good as using the 425 genetic map in combining scaffolds. The increase of the N<sub>50</sub> value in case of the 426 PBjelly2 result (using the PacBio reads for improving scaffolds) results from more 427 improvements of the shorter scaffolds. Comparing the PBjelly2 result for applying the 428 PacBio reads to v2.5 and vAM shows that the reduction of scaffold number and 429 increase of number of bases in the genome is similar (Table 5). MinION reads could 430 also be used for v2.5 assembly improvement, however the results were not as good 431 as for using PacBio reads, due to a much smaller number of reads. Improving the 432 genome v2.6 with the MinION reads is also possible, but the improvement is not as 433

- 434 good as for v2.5. This demonstrates that connections done using the PacBio reads are
- 435 also supported by MinION reads.

Setup	v2.5	PacBio vs. v2.5	MinION vs. v2.5
# scaffolds	3,166	3,066	3,123
# bases	196,005,095	203,024,676	196,600,700
N50	564,741	542,490	564,741
L50	56	58	56
Setup	vAM	PacBio vs. vAM	
# scaffolds	2,990	2,905	
# bases	196,022,695	203,137,854	
N50	10,141,718	10,314,234	
L50	9	9	
Setup	v2.6		MinION vs. v2.6
# scaffolds	2,895		2,886
# bases	203,150,143		203,450,934
N50	10,328,388		10,323,234
L50	9		9

### Table 5: Overview of the PBjelly2 result statistics for the different setups.

Comparison of values for the different genome versions with the values for the PBjelly2
output is shown. The PBjelly2 outputs are denoted in the form "read type" vs. "genome
version" to show which reads were used to improve which version of the genome. The
result for PacBio vs. vAM was the basis for v2.6 and MinION vs. v2.6 was the basis for
v3.0.

While PBjelly2 does not do as good a job as the genetic map approach in connecting scaffolds, its power is revealed by the gap filling. In genome v2.5 a total of 1,910 scaffolds contained uncalled bases. This number was reduced to 1,711 scaffolds (by

7.3%; Table 6) using PBjelly2 with PacBio reads. The exact number of uncalled bases 445 in the v2.5 A. arabicum genome was 25,768,296 (Table 6). In the PBjelly2 result only 446 13,940,203 Ns (Table 6) are left, a reduction by 45.9%. Comparing this with the 447 PBjelly2 result for the improvement of the vAM genome using PacBio reads (Table 6), 448 more gaps were removed from the connected genome. The number of scaffolds 449 containing Ns was reduced by 10.8% and the overall number of Ns was reduced by 450 50.0%, while the overall percentage of Ns in the genome remained the same in the 451 two results. Due to the small number of MinION reads, the improvement of the 452 assembly is less pronounced. 453

Setup	v2.5	PacBio vs. v2.5	MinION vs. v2.5
# scaffolds	1,910 (60.3%)	1,711 (56.0%)	1,901 (60.0%)
containing Ns			
# Ns	25,768,296 (13.2%)	13,940,203 (7.1%)	25,142,571 (12.8%)
Setup	VAM	PacBio vs. vAM	
# scaffolds	1,734 (58.0%)	1,546 (51.7%)	
containing Ns			
# Ns	25,785,896 (13.2%)	13,942,094 (7.1%)	
Setup	v2.6		MinION vs. v2.6
# scaffolds	1,542 (53.3%)		1,539 (53.2%)
containing Ns			
# Ns	13,946,922 (6.9%)		13,790,284 (6.8%)

454 **Table6: Gap/N analysis of different genome versions.** 

This table gives an overview of the number of Ns in the different genome versions and the PBjelly2 results.. For the number of scaffolds containing Ns, the percentage is given relative to the total number of scaffolds is given in brackets. For the number of Ns, the percentage is relative to the total number of bases in the respective assembly.

459

### 460 Migration of proteins to new genome version

The genome v2.5 harbors 23,594 annotated protein coding genes. Eight of them could 461 not be lifted because they were located next to a gap in the genome. Since it is possible 462 that PBjelly2 changes the sequences around gaps, the sequences of the genes were 463 not identical anymore and the programs were therefore not able to migrate some genes 464 from one assembly version to another. We checked the expression of the genes which 465 could not be lifted using Illumina RNA-seq data representing several developmental 466 stages (data not shown) and found that all of them had almost no expression, as a 467 result they were not lifted manually. In addition to some unlifted genes, there were 17 468 genes which could be lifted only partially due to the same reason. All the other 23,569 469 470 genes could be lifted. A set of 579 genes were removed due to being identical with other genes, and 140 genes were removed because they were located completely in 471 another gene. A total of 1,202 genes have no starting methionine, 2,055 have no stop, 472 132 genes contain internal stops and for 1,019 genes the length of the CDSs is not 473 dividable by three. In the end 19,363 genes were lifted which were not marked as 474 potential pseudogene or partial. 475

We find that that start point (assembly version) for improvement is not relevant. 476 PBjelly2 is able to make more improvements using the PacBio data than with the 477 MinION data due to the higher number of PacBio reads. The number of added bases 478 per read is higher for PacBio than MinION reads (18.71 vs. 9.67) and also the number 479 of removed Ns is higher (31.07 vs. 5.06) while the MinION reads lead to more scaffold 480 connections per read (2.49 x 10<sup>-4</sup> vs. 3.88 x 10<sup>-4</sup> connected scaffolds). Using the 481 MinION reads for improvement makes only a few changes, but they show that they 482 support the changes which were made to the genome using the PacBio data. Since 483 the Ae. arabicum genome was almost not contaminated at all, only three small 484

scaffolds had to be removed. Gene models were filtered for multiple genes and genes
contained in other genes. If a problem with a gene was found, it was marked in the
resulting GFF file. We note that there are gene models which are probably not correct
and need to be fixed in future studies.

The combination of genetic mapping and long reads significantly improved the structure of the assembly, reducing the total scaffold number and decreasing the number of gaps.

- 492 Genetic map of Aethionema arabicum
- 493 Genetic map v2.5:

494 SNP calling

A GBS approach [35] was used to generate genetic variation data for genetic mapping. 495 Illumina sequencing of the parental lines and the RILs resulted in 442,101,405 raw 496 reads after quality filtering. Using the TASSEL package [19] to match sequence tags 497 to markers, 160,379 SNPs could be called based on genome v2.5. SNPs identified 498 through the GBS method often take the form of many SNP 'islands', where a multitude 499 of SNPs is present over only a few kbp of sequence with the same states across 500 individuals. This makes genetic mapping difficult as it results in a very large number of 501 markers that are mostly redundant. We reduced this SNP amount using a sliding 502 window approach collapsing a group of SNPs that all have the same states across 503 individuals into one single marker over windows of 10 kbp, thus the bigger the scaffold 504 the more selected SNPs. This, together with filtering non-informative SNPs (missing 505 data in more than 30% many individuals) resulted in a core group of markers of 5,428 506 SNPs. 507

508 Genetic map calculation

We used JoinMap 4.1 to calculate the genetic linkage map for the Ae. arabicum RIL 509 population. For map v2.5, we first checked the marker similarity among the initial set 510 of 5,428 SNPs that were obtained from GBS based on genome v2.5 by a pairwise 511 comparison. SNPs that were highly similar (higher than 90%) were represented by one 512 marker, which refined the number of markers to 1,818. Grouping was selected at a 513 LOD threshold of 9.0, which led to the grouping of the expected 11 Linkage Groups 514 (LGs) (Figure 3). We further optimized each LG to avoid inflation of the map distance 515 due to saturating SNPs using a Maximum likelihood model. Markers that were not more 516 than 0.1 cM away were also eliminated. As a result, a final set of 746 SNP markers 517 was used on 11LGs. Out of 11 LGs, there are three LGs (4, 7 and 11) containing cluster 518 of segregation distorted SNPs (more than 50% of SNPs, supplemental file 519 linkage group map.xlsx). 520

The Ae. arabicum genetic map v2.5 consists of 11 LGs with average distance size of 521 162.5 cM, covered by 746 SNP markers with average of 67 markers per LG. The 522 average marker spacing was 2.4 cM, equivalent to approximately 169 kbp. The 523 centromere is suggested by the high density of SNP markers within a small genetic 524 distance (e.g. a low recombination frequency). These markers typically belong to 525 relatively small scaffolds, consistent with a high-repetitive DNA content, where only a 526 few SNPs were called. LG4 centromere is located at the end of the linkage, suggesting 527 an assembly problem or that LG4 is a telocentric chromosome. 528

529 Overall the markers are distributed relatively dense and even in v2.5, with the biggest 530 gap smaller than 18 cM. SNPs that reside in the same scaffold were in agreement 531 among each other on the direction of their scaffold.

532 Genetic map v3.0

The procedure to build genetic map v3.0 was similar to v2.5. SNP calling was 533 performed based on genome v3.0 instead of v2.5 resulting in a raw set of 141,914 534 SNPs. After similar quality control strategy as for v2.5, we construct v3.0 with a core 535 set of 632 SNPs (Figure 4). The 11 LGs were maintained with the total size of~ 1945 536 cM, average marker distance of 3.1 cM. This inflation of genetic distance in genetic 537 map v3.0 compared with v2.5 can be explain by the newly retrieved SNPs due to 538 resolved Ns in the genome. These new SNPs are mainly located in the centromeric 539 regions. In general, SNP order and orientation in LGs are in agreement with map v2.5 540 (Figure 5). We have made adjusted linkage group order in map v3.0 compared with 541 map v2.5 (linkage group map.xlsx). Based on the size of the group, the biggest one 542 is LG 1, and the smallest is LG 11 (Figure 4). 543

However, there is a significant difference between v3.0 and v2.5 at three LGs that harbor clusters of segregation distorted SNPs, LG 5, 6, and 9 (equivalent LG 4, 11 and 7 in v2.5, respectively): the reduced number of markers as well as the increased marker distance (Figure 3 and 4). In order to maintain 11 LGs and certain degree of newly called SNP incorporation, we had to reduced number of distorted markers in those LGs in v3.0, as a result the dearth of markers was observed (see supplemental file. linkage group map.xlsx)

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- 552

## 553 Conclusions and potential implications

Aethionema is becoming an outgroup model for the Brassicaceae core. Studies on its genome, relevant life-history traits and their evolution rely on genome and genetic map resources. Thus, this work helps pave the way for future research of the Brassicaceae family. We have provided an advanced version of the *Aethionema arabicum* genome

- and its first genetic map, which allows for pseudochromosome construction needed for
- analysis of genome evolution. Finally, quantitative trait loci (QTL) mapping for the wide
- range of traits in *Aethionema* (e.g. flowering time, plant fitness, chemical defense and
- <sup>561</sup> heteromorphism) will be greatly enabled by this genetic map.
- 562

#### 563 **Declarations**

None of the authors have any competing interests in the manuscript.

#### 565

566 List of abbreviations

BAC	Bacterial Artificial Chromosome
BS-seq	Bisulfite sequencing
ChIP-seq	Chromatin ImmunoPrecipitation DNA-Sequencing
CYP	Cyprus ecotype
FPC	Finger Printed Contigs
GFF	Generic/General Feature Format
LG	Linkage group
MinION	Oxford Nanopore MinION
PacBio	Pacific Biosciences
QTL	Quantitative trait loci
RIL	Recombinant Inbred Line
SNP	Single Nucleotide Polymorphism
TUR	Turkey ecotype
WGP	Whole Genome Profiling

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568

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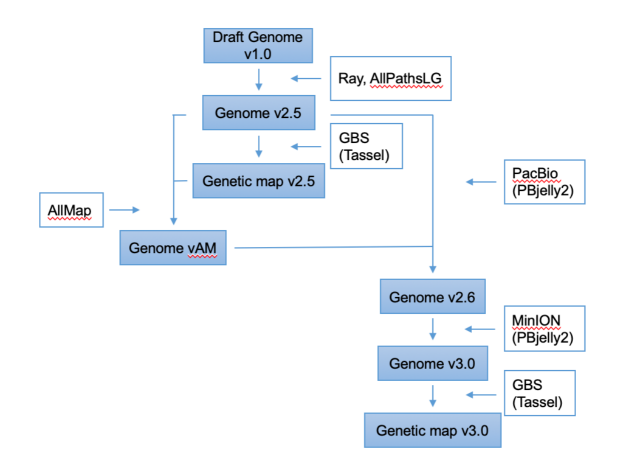
#### 574 Authors' contributions

TPN and ES designed the GBS experiment and performed the genetic map analyses. SM contributed to the development of the RIL population. AP did the initial genome assemblies. CM analysed the PacBio data, used the long-read data for superscaffolding and carried out the liftover of gene models to v3.0. FBH analysed the MinION data and carried out the contamination check. SAR supervised the work done by CM and FBH and conceived of this part of the work. EvdB performed GBS SNP calling. TPN, CM, SAR and ES wrote the manuscript with contributions by all authors.

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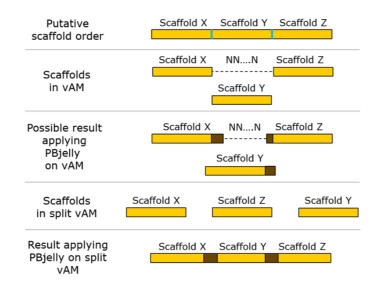
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## 589 Figure legends



590

- 591 **Figure 1: Overview of the analyses performed in this study**. In filled boxes are data
- sets, approaches and companying tools are in open boxes.



594

Figure 2: Problem arising from applying PBjelly2 on vAM. Scaffold borders are 595 visualized in blue and extensions of scaffolds introduced by PBjelly2 are shown in 596 brown. Assuming the true order of the scaffolds is shown on top of the figure, but 597 scaffold X and scaffold Z were already combined in the vAM assembly (second bar 598 from top) this could lead to a partial filling of the N-stretch and maybe an extension of 599 scaffold Y. However, PBjelly2 would not be able to place scaffold Y between the two 600 other scaffolds (middle bar). If the scaffolds were thus split again (second bar from 601 bottom), it is possible that the connections are made correctly applying PBjelly2 on the 602 split version (bottom bar). This only visualizes a theoretical case, in this work it 603 appeared every time that scaffold X and Y were connected by PBjelly2 and scaffold Z 604 had to be reconnected afterwards. 605

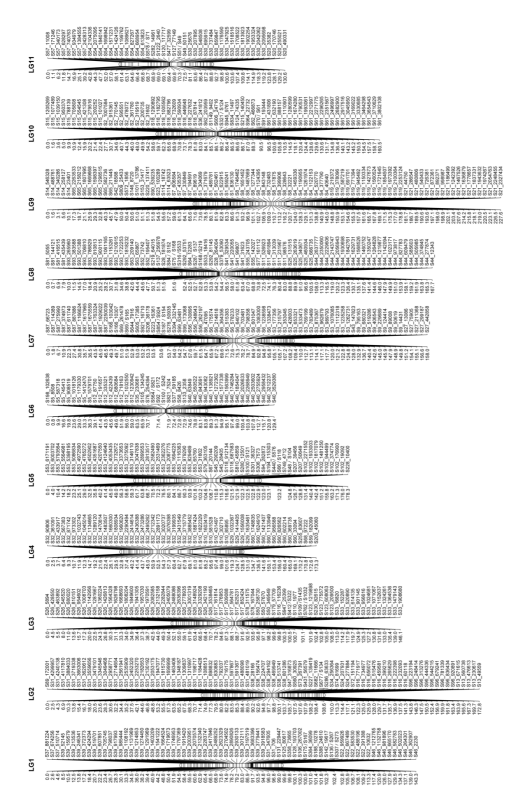


Figure 3: Aethionema arabicum genetic map v2.5. Genetic map version 2.5
 consists of eleven linkage groups. On each linkage group, genetic distance in cM is
 present on the left and SNP markers on the right.

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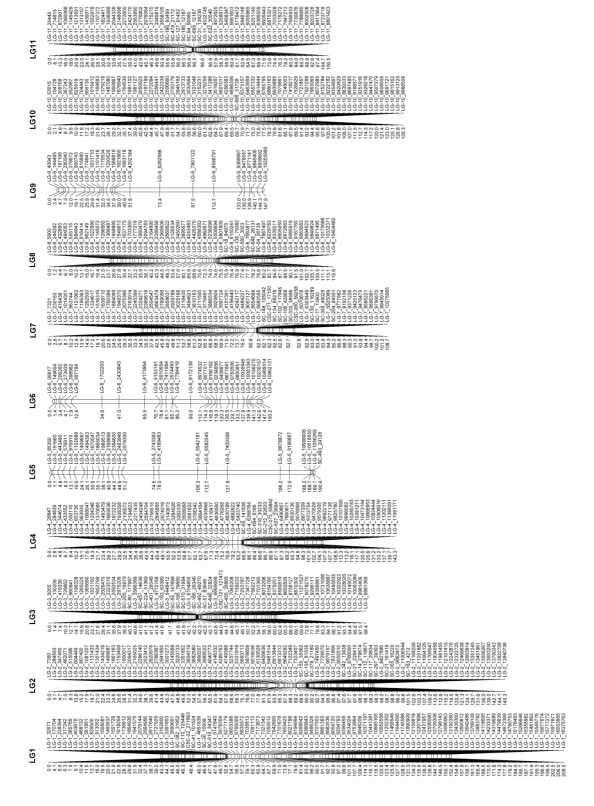


Figure 4: Aethionema arabicum genetic map v3.0. Genetic map version 3.0 consists of eleven linkage groups. On each linkage group, genetic distance in cM is present on the left and SNP markers on the right.

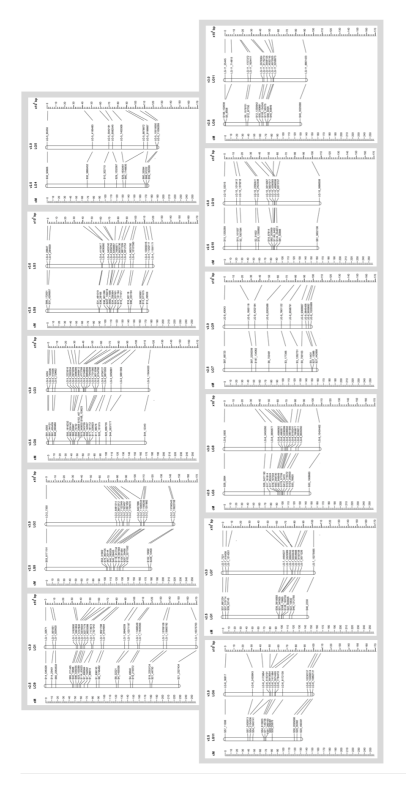


Figure 5: The alignment of genetic map v2.5, v3.0 and physical map. The alignment of the genetic map v2.5 and v3.0 were based on relative SNPs. The left ruler indicates genetic distance in cM and the right indicates physical distance in bp according to genome v3.0

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