1	Phylogenetic relationships in the genus Avena based on
2	the nuclear <i>Pgk1</i> gene
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22 Abstract

23	The phylogenetic relationships among 76 Avena taxa, representing 14 diploids,
24	eight tetraploids, and four hexaploids were investigated by using the nuclear plastid
25	3-phosphoglycerate kinase gene(<i>Pgk1</i>). A significant deletion (131 bp) was detected
26	in all the C genome homoeologues which reconfirmed a major structural divergence
27	between the A and C genomes. Phylogenetic analysis indicated the C_p genome is
28	more closely related to the polyploid species than is the C_{v} genome. Two haplotypes
29	of <i>Pgk1</i> gene were obtained from most of the AB genome tetraploids. Both types of
30	the <i>barbata</i> group showed a close relationship with the A_s genome diploid species,
31	supporting the hypothesis that both the A and B genomes are derived from an A_s
32	genome. Two haplotypes were also detected in A. agadiriana, which showed close
33	relationships with the A_{s} genome diploid and the A_{c} genome diploid, respectively,
34	emphasizing the important role of the A _c genome in the evolution of <i>A. agadiriana</i> .
35	Three homoeologues of the <i>Pgk1</i> gene were detected in five hexaploid accessions.
36	The homoeologues that might represent the D genome were tightly clustered with
37	the tetraploids A. marrocana and A. murphyi, but did not show a close relationship
38	with any extant diploid species.
20	Introduction

39 Introduction

The genus *Avena* L. belongs to the tribe Aveneae of the grass family
(Poaceae). It contains approximately 30 species [1-4] reflecting a wide range of
morphological and ecological diversity over the temperate and subtropical regions
[5]. The evolutionary history of *Avena* species has been discussed for decades, and

44	remains a matter of debate despite considerable research effort in this field.
45	Cytologically, three ploidy levels are recognized in the genus Avena: diploid,
46	tetraploid, and hexaploid, with a base number of seven chromosomes [6, 7]. The
47	diploids are divided clearly into two distinct lineages with the A and C genomes. All
48	hexaploid species share the same genomic constitution of ACD, corroborated by
49	fertile interspecific crosses among each other, as well as by their similar genome
50	sizes [8]. With less certainty, the tetraploids have been designated as AB or AA, AC or
51	DC, and CC genomes [9]. It is noteworthy that the B and D genomes within the
52	polyploid species have not been identified in any extant diploid species. There are
53	three C genome diploid species, which have been grouped into two genome types
54	$(C_{\rm p} \text{and} C_{\nu})$ according to their karyotypes [10]. Both types show a high degree of
55	chromosome affinity to the polyploid C genome [9-14], but none have been
56	undisputedly identified as the C genome progenitor of the polyploids.
57	The A genome origin of polyploid oats has also been under intense scrutiny.
58	However, there is no conclusive evidence regarding which the A genome diploid
59	contributed to the polyploid oats. There are up to 12 species designated as A
60	genome diploids. These species have been further subdivided into five sub-types of
61	A_c , A_d , A_l , A_p and A_s genomes, according to their karyotypes [6, 7]. Most research
62	based on karyotype comparisons [6, 15], in situ hybridization [11, 16-18], as well as
63	the alignments of nuclear genes [13, 14] suggest that one of the A_s genome species
64	may be the A genome donor of polyploid oats. Alternatively, some studies have

65 proposed the A_c genome diploid *A. canariensis* [19], or the A_l genome diploid *A.*

66 *longiglumis* [9, 12] as the most likely A genome donor.

67	The absence of diploids with the B and D genomes complicates the B and D
68	genome donor identification. It is generally accepted that both B and D genomes are
69	derived from A genomes, due to the high homology between the B and A genomes
70	[11, 20], as well as between the D and A genomes [16, 19, 21]. Our recent study
71	based on high-density genotyping-by-sequencing (GBS) markers [9] provided strong
72	evidence that the three tetraploid species formerly designated as AC genomes are
73	much closer to the C and D genomes of the hexaploids than they are to the
74	hexaploid A genome. These findings suggest that the hexaploid D genome exists in
75	the extant tetraploids. However, no extant diploid species, even the A_c genome
76	diploid A. canariensis, which was considered as the most likely D genome progenitor
77	based on direct evidence from morphological features [22] and indirect evidence
78	from fluorescent in situ hybridization (FISH) [18], showed enough similarity to the D
79	genome of tetraploid and hexaploid oats to warrant consideration as a direct D
80	genome progenitor.
81	In the case of the B genome, an initial study of chromosome pairing of
82	hybrids between the AB genome tetraploids and the A_{s} genome diploids suggested
83	that the B genome arose from the A_s genome through autoploidization [23].
84	Recently, another GBS study [19] showed that the AB genome tetraploid species fell

85 into a tight cluster with A_s genome diploids, also supporting the hypothesis that the

86 B genome arose through minor divergence following autoploidization. However,

87	other evidence from C-banding [24], FISH [17], RAPD markers [25], and DNA
88	sequence alignment [14] has indicated a clear distinction between A and B genomes,
89	suggesting an allotetraploid origin of the AB genome tetraploid species. The most
90	probable A genome progenitor of the AB genome tetraploids is assumed to be an A_s
91	genome diploid species, while the B genome of these species remains controversial.
92	Single or low copy nuclear genes are widely used in phylogenetic analyses
93	due to their bi-parental inheritance and to the informativeness of mutations. Such
94	studies have successfully revealed multiple polyploid origins, and clarified
95	hybridization events in a variety of plant families [26, 27]. In a previous study [14],
96	we investigated the relationships among Avena species by sequencing the
97	single-copy nuclear acetyl-coA carboxylase gene (Acc1). The results provided some
98	useful clues to the relationships of Avena species.
99	The <i>Pgk1</i> gene, which encodes the plastid 3-phosphoglyceratekinase, is
100	another nuclear gene that has been widely used to reveal the evolutionary history of
101	the Triticum/Aegilops complex due to its single copy status per diploid chromosome
102	in grass [26, 28, 29]. The <i>Pgk1</i> gene is now considered to be superior to the <i>Acc1</i>
103	gene in phylogenetic analysis, since it has more parsimony informative sites than the
104	Acc1 gene [26, 29]. In the present study, we sequenced cloned Pgk1gene copies
105	from 76 accessions representing the majority of Avena species, in an attempt to
106	further clarify evolutionary events in this important genus.
107	Materials and Methods

Materials and Methods

108 Plant materials

109 A total of 76 accessions from 26 Avena species were investigated to r	epresent
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- 110 the geographic range of six sections in *Avena*, together with one accession from
- 111 *Trisetopsis turgidula* as a functional outgroup (Table 1). All seeds were provided by
- 112 Plant Gene Resources of Canada (PGRC) or the National Small Grains Collection,
- 113 Agriculture Research Service, United States Department of Agriculture (USDA, ARS)
- 114 with the exception of the three accessions of *A. insularis*, which were kindly provided
- by Dr. Rick Jellen, Brigham Young University, Provo, UT, USA. The species A.
- 116 *atherantha*, *A. hybrida*, *A. matritensis* and *A. trichophylla* described in Baum's [1]
- 117 monograph and A. prostrata described by Ladizinsky [30] were not included due to a
- 118 lack of viable material.
- 119 Table 1. List of materials used in the present study including species, haplomes,
- 120 accession number, origin, abbreviation displayed in MJ network, and the sequence
- 121 number in Genbank (<u>https://www.ncbi.nlm.nih.gov</u>).

Таха	Haplomes	Accession	Origin [*]	Abbrev-	Genbank
IdXd		Number		iation	Accession
Section Ventricosa					
<i>A. clauda</i> Dur.	Cp	CN 19242	Turkey	CLA1_1	KU888786
		CN 21378	Greece	CLA2_1	KU888787
		CN 21388	Algeria	CLA3_1	KU888804
		CN 24695	Turkey	CLA4_1	KU888784
<i>A. eriantha</i> Dur. (syn <i>A. pilosa</i> Bieb.)	C _p	Clav 9050	United Kingdom	ERI1_1	KU888785
		PI 367381	Madrid, Spain	ERI2_1	KU888805
<i>A. ventricosa</i> Balansa ex Coss.	C _v	CN 21405	Algeria	VEN1_1	KU888806
		CN 39706	Azerbaijan	VEN2_1	KU888807
Section Agraria					
A. brevis Roth	A _s	Clav 1783	German	BRE1_1	KU888707

		Class 0112	F		KU 000740
		Clav 9113	Europe	BRE2_1	KU888718
A him mine And	•	PI 258545	Portugal	BRE3_1	KU888710
A. hispanica Ard.	A _s	CN 25676	Portugal	HIS1_1	KU888714
		CN 25727	Portugal	HIS2_1	KU888711
		CN 25766	Portugal	HIS3_1	KU888709
		CN 25778	Portugal	HIS4_1	KU888712
A. nuda L.	As	PI 401795	Netherlands	NUD1_1	KU888734
A. strigose Schreb.	As	PI 83722	Australia	STR1_1	KU888719
		PI 158246	Lugo, Spain	STR2_1	KU888713
		Clav 9066	Ontario, Canada	STR3_1	KU888708
Section Tenuicarpa					
<i>A. agadiriana</i> Baum & Fedak	AB	CN 25837	Africa: Morocco	AGA1_1	KU888753
				AGA1_2	KU888774
		CN 25854	Africa: Morocco	AGA2_1	KU888777
				AGA2_2	KU888754
		CN 25856	Africa: Morocco	AGA3_1	KU888776
				AGA3_2	KU888751
		CN 25863	Africa: Morocco	AGA4_1	KU888775
		CN 25869	Africa: Morocco	AGA5_1	KU888752
				AGA5_2	KU888778
<i>A. atlantica</i> Baum & Fedak	A _s	CN 25849	Africa: Morocco	ATL1_1	KU888757
		CN 25859	Africa: Morocco	ATL2_1	KU888756
		CN 25864	Africa: Morocco	ATL3_1	KU888739
		CN 25887	Africa: Morocco	ATL4_1	KU888737
		CN 25897	Africa: Morocco	ATL5 1	KU888736
<i>A. barbata</i> Pott ex Link	AB	PI 296229	Northern, Israel	BAR1_1	KU888723
		PI 337802	Izmir, Turkey	BAR2_1	KU888722
				BAR2_2	KU888732
		PI 337826	Greece	BAR3_1	KU888720
		PI 282723	Northern, Israel	_ BAR4_1	KU888729
			Macedonia,	_	
		PI 337731	Greece	BAR5_1	KU888731
		PI 367322	Beja, Portugal	BAR6_1	KU888730
A. canariensis Baum et al	A _c	CN 23017	Canary Islands	CAN1_1	KU888779
		CN 23029	Canary Islands	CAN2_1	KU888782
		CN 25442	Canary Islands	CAN3_1	KU888780
		CN 26172	Canary Islands	CAN4_1	KU888783
		CN 26195	Canary Islands	CAN5_1	KU888781
<i>A. damascena</i> Rajah & Baum	A _d	CN 19457	Syria	DAM1_1	KU888744
		CN 19458	Syria	DAM2_1	KU888745

		CN 19459	Syria	DAM3_1	KU888747
A. hirtula Lag.	As	CN 19530	Antalya, Turkey	HIR1_1	KU888738
		CN 19739	Algeria	HIR2_1	KU888762
		CN 21703	Morocco	HIR3_1	KU888717
A. longiglumis Dur.	Ai	Clav 9087	Oran, Algeria	LON1_1	KU888741
		Clav 9089	Libya	LON2_1	KU888749
		PI 367389	Setubal, Portugal	LON3_1	KU888750
<i>A. lusitanica</i> Baum	As	CN 25885	Morocco	LUS1_1	KU888746
		CN 25899	Morocco	LUS2_1	KU888748
		CN 26265	Portugal	LUS3_1	KU888742
		CN 26441	Spain	LUS4_1	KU888763
A. wiestii Steud.	As	PI 53626	Giza, Egypt	WIE1_1	KU888715
		Clav 9053	Ontario, Canada	WIE2_1	KU888716
Section Ethiopica					
A. abyssinica Hochst.	AB	PI 411163	Seraye, Eritrea	ABY1_1	KU888724
		PI 411173	Tigre, Ethiopia	ABY2_1	KU888740
				ABY2_2	KU888725
A. vaviloviana Mordv.	AB	PI 412761	Eritrea	VAV1_1	KU888743
				VAV1_2	KU888728
		PI 412766	Shewa, Ethiopia	VAV2_1	KU888726
				VAV2_2	KU888735
Section Pachycarpa					
A. insularis Ladiz.	AC(DC)	sn	Sicily, Italy	INS1_1	KU888794
				INS1_2	KU888705
		6-B-22	Sicily, Gela, Italy	INS2_1	KU888706
				INS2_2	KU888796
		INS-4	Sicily, Gela, Italy	INS3_1	KU888790
				INS3_2	KU888704
A. maroccana Grand.					
(syn. <i>A magna</i> Murphy	AC(DC)	Clav 8330	Morocco	MAR1_1	KU888773
et Terrell)					
				MAR1_2	KU888799
		Clav 8331	Khemisset,	MAR2_1	KU888721
			Morocco	1017 (112_1	10000721
				MAR2_2	KU888800
A. murphyi Ladiz.	AC(DC)	CN 21989	Spain	MUR1_1	KU888767
				MUR1_2	KU888802
		CN 25974	Morocco	MUR2_1	KU888769
				MUR2_2	KU888788
Section Avena					
A.fatua L.	ACD	PI 447299	Gansu, China	FAT1_1	KU888768
				FAT1_2	KU888795
		PI 544659	United States	FAT2_1	KU888764
		0			

				FAT2_2 FAT2_3	KU888760 KU888798
A.occidentalis Dur.	ACD	CN 4547	Canary Islands, Spain	OCC1_1	KU888791
		CN 23036	Canary Islands, Spain	OCC2_1	KU888755
				OCC2_2	KU888803
				OCC2_3	KU888771
		CN 25942	Morocco	OCC3_1	KU888733
				OCC3_2	KU888789
				OCC3_3	KU888758
		CN 25956	Morocco	OCC4_1	KU888801
				OCC4_2	KU888772
A. sativa L.	ACD	PI 194896	Gonder, Ethiopia	SAT1_1	KU888727
				SAT1_2	KU888759
				SAT1_3	KU888793
		PI 258655	Russian Federation	SAT2_1	KU888797
				SAT2_2	KU888766
				SAT2_3	KU888761
A. sterilis L.	ACD	PI 411503	Alger, Algeria	STE1_1	KU888765
		PI 411656	Tigre, Ethiopia	STE2_1	KU888792
				STE2_2	KU888770
Outgroup					
<i>Trisetopsis turgidula</i> Röser & A. Wölk		PI 364343	Maseru, Lesotho		KU888808

^{*} Origin represents the collection site of wild material where this information is

123 available, otherwise it represents the earliest source for which information is

124 available.

125 DNA isolation, cloning and sequencing

126 Genomic DNA was isolated from fresh leaves of single plants following a

127 standard CTAB protocol [31]. *Pgk1* gene sequences were amplified by using a pair of

- 128 *Pgk1*-specific primers, PGKF1 (5'-TCGTCCTAAGGGTGTTACTCCTAA-3') and PGKR1
- 129 (5'-ACCACCAGTTGAGATGTGGCTCAT-3') described by Huang et al. [28]. Polymerase

130	chain reactions (PCR) were carried out under cycling conditions reported previously
131	[26]. After estimating the size by 1.0% agarose gel, PCR products were purified using
132	the QIAquick gel extraction kit (QIAGEN Inc., USA). The purified products were
133	cloned into the pMD19-T vector (Takara) following the manufacturer's instructions.
134	Initially, 6-8 positive clones from each of four accessions from 4 diploid
135	species, including A. canariensis (A_c), A. longiglumis (A_l), A. strigosa (A_s), and A. clauda
136	(C _p), were sequenced to confirm that the <i>Pgk1</i> gene was present in <i>Avena</i> diploid
137	species as a single copy. After confirming its single copy status in diploid species, 2-3
138	positive clones were selected and sequenced from each accession of the remaining
139	diploid species. In order to isolate all possible homoeologous sequences in polyploid
140	species, 4-6 positive clones from each accession of the tetraploid species and 5-10
141	positive clones from each accession of the hexaploid species were selected and
142	sequenced. All the cloned PCR products were sequenced on both strands by a
143	commercial company (Sangon Biotech Co., Ltd., Shanghai, China) based on Sanger
144	sequencing technology.

145 Sequence alignment and phylogenetic analysis

146The homology of sequences was verified using the BLAST program in NCBI. In147order to reduce the matrix size of the dataset, redundant sequences were removed,148keeping one representative sequence if several identical sequences were derived149from the same accession. Sequences were aligned using ClustalW software with150default parameters [32] followed by manual correction. Substitution saturation of151*Pgk1* sequences was examined using DAMBE version 5 [33] by calculating and plotting10

152	pairwise rates of transitions and transversions against sequence divergence under
153	the TN93 model. Phylogenetic trees were created by using Maximum parsimony
154	(MP), and Bayesian inference (BI). MP analysis was performed on PAUP* 4.0b10 [34]
155	using the heuristic search with 100 random addition sequence replicates and Tree
156	Bisection-Reconnection (TBR) branch swapping algorithms. Bootstrapping with 1000
157	replicates was estimated to determine the robustness of formed branches [35]. Gaps
158	in the sequence alignment were disregarded using the option 'gapmode=missing',
159	which is consistent with an assumption that insertion/deletion events are an
160	independent stochastic process from SNP substitutions. BI analysis was carried out
161	by using MrBayes v3.2 [36]. The best-fit substitution model for BI analysis was
162	GTR+F+I, which was determined by using MrModelTest v2.3 under Akaike
163	information criteria (AIC) (http://www.ebc.uu.se/systzoo/staff/nylander.html). Four
164	Markov chain Monte Carlo (MCMC) chains with default priors settings were run
165	simultaneously. To ensure the two runs converged onto the stationary distribution,
166	6,000,000 generations were run to make the standard deviation of split frequencies
167	fall below 0.01. Samples were taken every 100 generations. The first 25% samples
168	from each run were discarded as the "burn-in". The 50% majority-rule consensus
169	tree was constructed from the remaining trees. Posterior probability (PP) values
170	were used to evaluate the statistical confidence of each node.
474	Notwork analysis

Network analysis

172 The median-joining (MJ) network [37] method has been demonstrated to be 173 an effective method for assessing the relationship in closely related lineages [38],

174	and thus was app	lied in this stud	y. As MJ alg	gorithms are d	lesigned for
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- non-recombining molecules [37], DNA recombination was test by using a pragmatic
- 176 approach-Genetic Algorithm Recombination Detection (GARD), described by Pond et
- al. [39]. The test was carried out on a web-based interface for GARD at
- 178 http://www.datamonkey.org/GARD/. Building upon this test, the intron data was
- used for MJ reconstruction due to the absence of recombination signal, while
- 180 potential recombination signals were detected in the exon regions. The MJ network
- analyses was performed using the Network 4.6.1.4 program (Fluxus Technology Ltd,
- 182 Clare, Suffolk, UK).
- **183 Results**

184 Sequence analysis

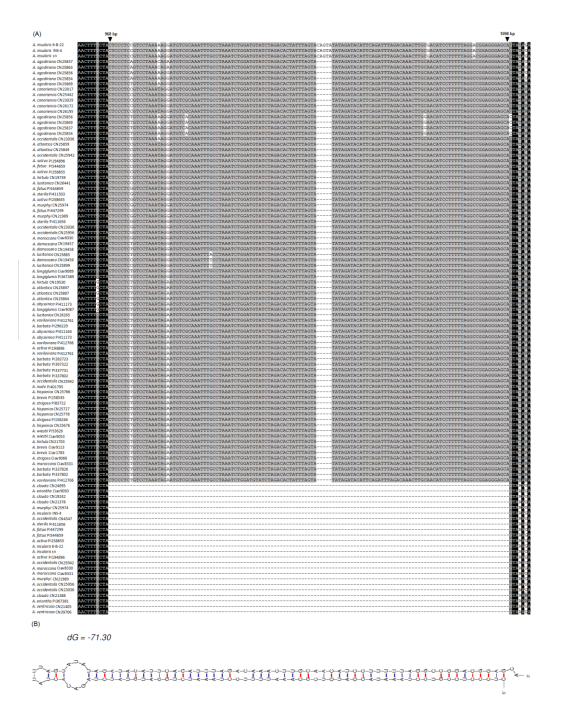
185 A total of 237 clones were sequenced from 76 accessions of 26 Avena species. Following removal of the redundant sequences within each accession, 104 186 sequences were identified, including one from each of the 44 diploid accessions, 37 187 188 unique sequences from 22 tetraploids, and 23 from 10 hexaploids. Theoretically, 44 189 homoeologues should be isolated from 22 tetraploid accessions, and 30 single-copy 190 homoeologues were expected from 10 hexaploid accessions. However, the full number of expected homoeologues were not isolated from every polyploid species 191 192 for various potential reasons. In particular, within the AB genome tetraploid species A. barbata, only one copy was detected in five of its six accessions, whereas two very 193 194 similar (only one site varied in exon 2) copies were detected in the sixth accession. This also happened in the hexaploid species A. sterilis, for which two accessions 195

provided only two homoeologues each. For these taxa, the missing genome type
might be detected by screening a larger number of positive clones, but it is also
possible that these accessions contain genomes of high similarity or autopolyploid
origin. Another possibility that cannot be ruled out within the polyploids is the loss
of one gene copy through homoeologous recombination or deletion.

201

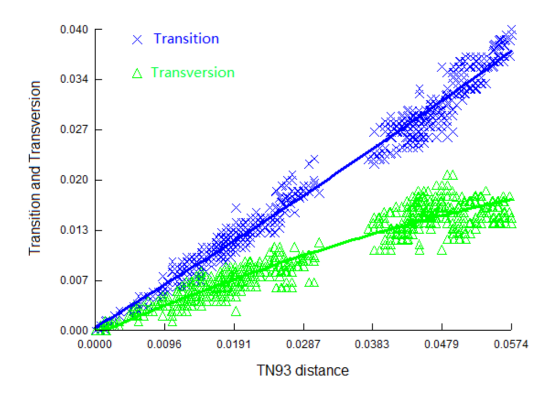
All of the Pgk1 gene sequences isolated in this study contain 5 exons and 4

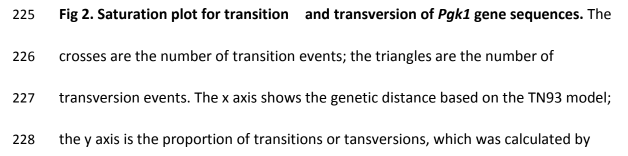
introns, covering a total length from 1391 bp to 1527 bp, which is consistent with 202 203 previous studies of this gene in wheat [28] and Kengyilia [26]. The alignment of Pak1 204 sequences including both exons and introns resulted in a matrix of 1539 nucleotide positions, of which 11.6% (179/1539) were variable, and 10.1% (155/1539) were 205 206 parsimony informative. The nucleotide frequencies were 0.264 (A), 0.304 (T), 0.199 207 (C), and 0.232 (G). A significant (131-bp) insertion/deletion feature (Fig 1A) occurred at position 968, whereby all non-C genome type sequences contained the inserted 208 209 (or non-deleted) region. Further analysis indicated that this region is likely an 210 inserted inverted repeat, which belongs to the MITE stowaway element. Its 211 secondary structure is shown in Fig 1B. This insertion/deletion event could be used 212 as a genetic marker for rapid diagnosis of Avena species containing the C genome.



- Fig 1. Pgk1 gene sequence analysis. (A) Partial alignment of the amplified Pgk1 gene
- of Avena species (B) Secondary structure of the deletion sequence between the A
- and C genomes.
- 216 **Phylogenetic analyses**

217	The substitution plot for <i>Pgk1</i> (Fig 2) indicated that the <i>Pgk1</i> gene was not
218	saturated and that it could be used for phylogenetic analysis. Phylogentic trees of 76
219	Avena accessions with the oat-like species Trisetopsis turgidula as outgroup were
220	generated through maximum parsimony and Bayesian inference approaches on the
221	non-redundant dataset. The parsimony analysis resulted in 80 equally parsimonious
222	trees (consistency index (CI) =0.632, retention index (RI) =0.954). BI analysis inferred
223	an almost identical tree topology as the MP analysis, so the MP results were selected
224	to describe this study (Fig 3).





- using the number of transitions or transversions divided by the sequence length. The
- 230 curves show the trends of the variance of transitions and transversions with the

231 genetic distance increasing.

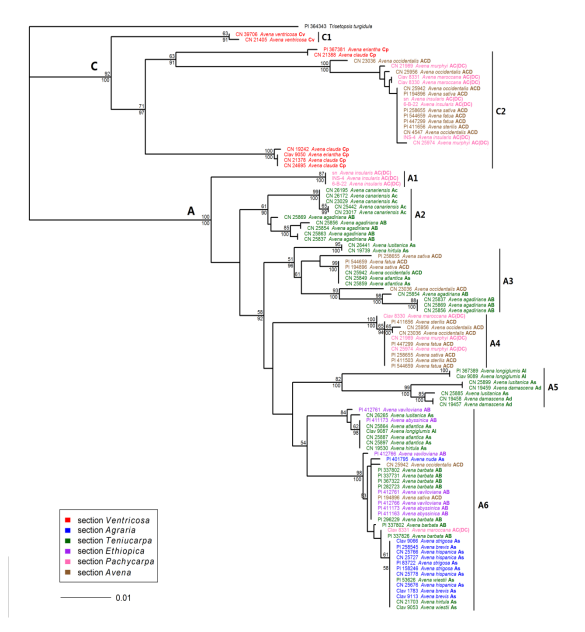


Fig 3. Maximum parsimony tree derived from *Pgk1* sequence data. The tree was
 constructed using a heuristic search with TBR branch swapping. Numbers above and
 below the branches are bootstrap support (BS) values ≥50% and Bayesian posterior

probability (PP) values ≥90%. Accession number, species name and haplome are
indicated for each taxon.

237	Fig 3 shows that the <i>Pgk1</i> gene sequences from 76 Avena accessions were
238	split into two distinct clades with high BS (100% and 92%) and PP (100% and 100%)
239	support. One clade contained all C-genome type sequences, hence referred to as the
240	C genome clade. The other clade contained all sequences from the species carrying
241	the A genome, henceforth, referred to as the A genome clade. The C genome clade
242	was composed of two major subclades. All C_{ν} genome diploids and two C_{p} genome
243	diploid accessions formed the subclade C1 with 63% BS and 91% PP support, while
244	subclade C2 included four C_p diploids accessions, seven AC(DC) genome tetraploid
245	accessions and nine hexaploid accessions with 71% BS and 97% PP support. The Pgk1
246	gene sequences in the A genome clade were further split into six major subclades.
247	The AC(DC) genome tetraploid species A. insularis was distinct from the other
248	species, consequently forming a monophyletic clade (A1) with high BS (87%) and PP
249	(100%) support. All five accessions of the A_c genome diploid species A. canariensis
250	and one genome homoeologue of the AB genome tetraploid species A. agadiriana
251	clustered together into subclade A2. Subclade A3 was composed of four accessions
252	of the AB genome tetraploids A. agadiriana, five hexaploid accessions (A.
253	occidentalis CN 23036 and CN 25942, A. sativa PI 194896 and PI 258655, A. fatua PI
254	544659) and four A_s genome diploid accessions (<i>A. atlantica</i> CN25849 and CN 25859,
255	A. lusitanica CN 26441, and A. hirtula CN 19739). One genome sequence of the
256	AC(DC) genome tetraploids (without <i>A. insularis</i>) and the hexaploids formed a 17

257	homogeneous clade (A4) that was separated from other species with high BS (100%)
258	and PP (100%) support. The subclade A5 consisted of the A_d genome diploid A.
259	damascena, the A_1 genome diploid A. longiglumis, and the A_s genome diploid A.
260	lusitanica. The remaining sequences from the A genome diploids and the AB genome
261	tetraploids (without A. agadiriana) formed a relatively broader cluster A6, together
262	with two hexaploid accessions (A. sativa PI 194896 and A. occidentalis CN 25942)
263	and one AC (DC) genome tetraploid accession (A. maroccana Clav 8831).
264	Three groups of haplotypes of <i>Pgk1</i> sequences were identified in five
265	hexaploid accessions (A. fatua PI 544659, A. occidentalis CN 25942, CN 23036, and A.
266	sativa PI 194896, PI 258655). These sequences fell into four subclades. One group
267	clustered with the C genome diploids in subclade C2, and one group clustered with
268	AC(DC) genome tetraploids in subclade A4. We hypothesize that these two types
269	represent homoeologues from the C and D genomes, respectively. A third and fourth
270	group fell into subclades A3 and A6. Since these two groups are highly separated, it
271	is possible that they represent different A-genome events leading to different
272	hexaploid lineages.

273 Network analysis

To gain better insight into relationships within closely related lineages, MJ network reconstruction based on the haplotypes of *Pgk1* sequences was employed. Due to the potential presence of recombination in the exon regions, the intron data was used for MJ network reconstruction. A total of 40 haplotypes were derived from 104 *Pgk1* gene sequences (Fig 4). This low level of haplotype diversity demonstrates

279	the high conservation of this gene within genus Avena. The MJ network recovered a
280	nearly identical phylogenetic reconstruction to that based on the MP and BI trees,
281	therefore we identified the clades from the MP results (Fig 3) within the MJ network
282	(Fig 4). Based on the topology and frequency of haplotypes, the MJ network was split
283	into two main groups. The two major groups representing two distinct types of
284	haplotypes (A and C genomes) were distinguished due to the 131 bp
285	insertion/deletion. Ten C genome haplotypes were observed, which were much less
286	diverse than the 30 A genome haplotypes. The two main groups were further
287	subdivided into clusters corresponding to the eight MP-based subclades discussed
288	earlier. The only divergence was that the AC(DC) genome tetraploids A. insularis,
289	which formed a separate clade (A1) in MP and BI trees, fell into together with the AB
290	genome tetraploid A. agadiriana and the A_c genome diploid A. canariensis to form a
291	relatively broad cluster in the MJ network (A1&A2).

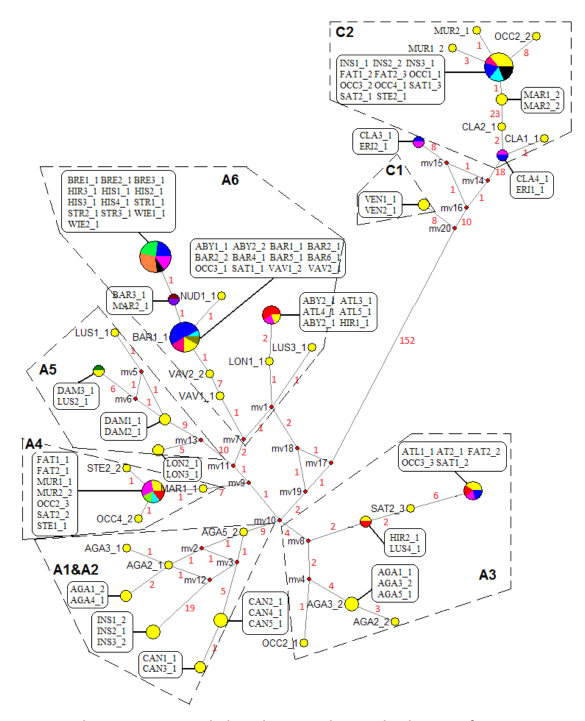


Fig 4. Median-joining networks based on 40 *Pgk1* gene haplotypes of intron regions

293 derived from 26 Avena species. Each circular node represents a single haplotype,

- with relative size being proportional to the frequency of that haplotype. Distinct
- 295 colors in the same haplotype node represent different species sharing the same
- 296 haplotype (colors are arbitrary). Median vectors (mv) represent the putative missing

297	intermediates. Numbers along network branches indicate the number of bases
298	involved in mutations between two nodes. Clusters (surrounded by dashed lines) are
299	named based on clade names shown in the MP tree (Fig 3). Three-letter
300	abbreviations of species names are listed in Table 1. The numbers immediately after
301	each species abbreviation represent different accessions of the same species, and
302	the number following the underscore identifies different haplotypes from the same
303	accession.

304 **Discussion**

Two distinct diploid lineages exist in genus *Avena.*

306 A significant 131 bp insert/deletion separated all Avena diploid species into two distinct groups representing the A and C genomes, respectively (Figs 1 and 4). 307 308 These groups were also separated based on the MP or BI analysis that ignored gaps (Fig 3), indicating that the separation of A and C genomes is the most ancient major 309 articulation in the genus Avena, a result that is consistent with most other literature 310 311 [13, 14, 40]. MJ network analysis revealed that the C genome diploids have much 312 lower levels of haplotype diversity than the A genome diploids. Within the C genome 313 diploids, the C_p genome haplotypes were relatively more diverse than those of the C_v genome. These results might be explained by the geographic distribution of these 314 315 species. The A genome diploids are distributed in a large region between latitude 20 and 40° N, while the C genome diploid species are restricted to a narrow territory 316 along the Mediterranean shoreline [1]. The geographic distributions of the C genome 317

diploid species are overlapping, but the range of the C_p genome diploid species is much broader than that of the C_v genome diploid species [41].

320 The A genome diploid species are the most diverse set of species in genus Avena, and chromosome rearrangements have occurred during the divergence of 321 A-genomes from a common progenitor [41], resulting in the subdivision of the A 322 genome into five types, of which we have investigated four. Our results showed that 323 324 species with genome types A_c , A_l , and A_d formed groups that correspond well with 325 previously reported structural differences. However, the A_s genome diploids appear 326 to be much more diverse than previously reported, and are scattered into different 327 subclades (Fig 3). Baum [1] divided all A_s genome diploids into two sections, section Agraria and section Tenuicarpa. All species of section Agraria have florets with a 328 329 domesticated (non-shattering) base, whereas the other A_s species share relatively 330 narrow spikelets. However, classification based on simple morphological traits is increasingly controversial. In this study, the A_s genome diploid species of section 331 332 Agraria showed high degree of genetic homogeneity, consistently forming their own 333 subclade A6, but other As genome species in section *Tenuicarpa* did not have their own subclade. A. wiestii showed a close relationship with the species of section 334 Agraria, suggesting that it may be better-classified within that section. This result is 335 336 in agreement with previous studies based on RAPD (Perchuk et al. 2002) and karyotypic comparisons (Badaeva et al. 2005). Accessions of the other two A_s 337 338 genome species of section Tenuicarpa (A. atlantica and A. hirtula) were scattered into different subclades. These results were also observed in other studies (Peng et 339

340	al. 2010, Yan et al. 2014). A. lusitanica, another A _s species of section Tenuicarpa, was
341	diverged from other A_{s} species, but showed a close relationship to those with the A_{d}
342	genome species A. damascena. This divergence has also been observed in many
343	other studies [8, 9, 14, 40]. These, and other incongruences between morphological
344	characters and genetic differences raise questions about appropriate taxonomical
345	classifications among A _s genome species.

346 The A_s and A_c genomes played roles in the AB tetraploid

347 formation.

348 Four recognized species have been proposed to have an AB genome composition. Of these, A. barbata, A. abyssinica and A. vaviloviana are grouped into 349 a biological species known as the barbata group, while A. aqadiriana is distinct [25, 350 351 42]. Our results confirmed the reported structural differences between these two groups (Fig 3). Two different Pak1 gene sequences were detected from most of the 352 AB genome tetraploids, supporting their allotetraploid origins. However, the 353 354 genomes of A.barbata showed the least divergence, with only one of six A. barbata 355 accessions providing multiple sequences, both of which were very similar. It seems 356 that little divergence has occurred within the genome of A. barbata compared with that of A. abyssinica and A.vaviloviana, suggesting that A. barbata is the ancestral 357 358 version of the species within the *barbata* group. This is supported by two lines of evidence. First, both A. abyssinica and A.vaviloviana are semi-domesticated forms 359 360 that occur almost exclusively in Ethiopia, whereas the wild A. barbata are more geographically distributed, but can still be found close to the abyssinica and 361

362	vaviloviana forms [43]. The second line of evidence was provided by FISH and
363	Southern hybridization [17], which found some B chromosomes of A. vaviloviana are
364	involved in inter-genomic translocations, while these rearrangements were not
365	detected in <i>A. barbata</i> . There is little doubt that the A genome diploids have been
366	involved in the formation of the <i>barbata</i> species. Some studies have suggested that
367	both the A and B genomes of <i>barbata</i> species are diverged A _s genomes [16, 23, 44],
368	while some others proposed that the B genome might have originated from other A
369	genome diploid species [24, 25, 45]. In this study, both types of <i>Pgk1</i> sequences
370	detected from the barbata group showed high degree of genetic homogeneity with
371	the A_s genome diploids (Fig 3), thus it was impossible to determine which type
372	represents the A or B genome.
373	The recently discovered tetraploid species A. agadiriana was also proposed
374	to have an AB genome composition because of its high affinity with A. barbata [23].
375	However, this designation has been questioned due to chromosomal divergences
376	between A. agadiriana and the barbata species, as revealed by cytological studies
377	
	[45, 46] and by molecular data [9, 13, 14]. In the current study, two distinct types of
378	[45, 46] and by molecular data [9, 13, 14]. In the current study, two distinct types of <i>Pgk1</i> sequences were obtained in <i>A. agadiriana</i> . One copy clustered with the A_c
378 379	
	<i>Pgk1</i> sequences were obtained in <i>A. agadiriana</i> . One copy clustered with the A _c
379	<i>Pgk1</i> sequences were obtained in <i>A. agadiriana</i> . One copy clustered with the A _c genome species <i>A. canariensis</i> , whereas the other copy fell into cluster A3 with the
379 380	<i>Pgk1</i> sequences were obtained in <i>A. agadiriana</i> . One copy clustered with the A _c genome species <i>A. canariensis</i> , whereas the other copy fell into cluster A3 with the A _s species <i>A. atlantica</i> , <i>A. hirtula</i> , <i>A. lusitanica</i> , and the hexaploids <i>A. occidentalis</i> , <i>A</i> .

from the *barbata* group, and that one of its two genomes originates from the A_c
genome species *A. canariensis*, whereas the other one is closely related to the A_s
species.

387 The tetraploid species A. maroccana and A. murphyi are

closely related to the hexaploids, while *A. insularis* is

389 diverged.

The other tetraploid group (Avena sect. Pachycarpa) contains three species, A. 390 391 maroccana, A. murphyi, and the recently discovered A. insularis. Initial studies based 392 on genomic in situ hybridization [47] supported an AC genome designation for these species. However, this designation has been challenged by FISH analysis, which has 393 revealed that this set of tetraploid species, like the D chromosomes of the hexaploid 394 395 oats, lacks a repetitive element that is diagnostic of the A genome [18]. This, together with other molecular evidence [14, 48] and our recent whole-genome 396 analysis based on GBS markers [9], suggests that these tetraploid species contain the 397 398 genome designated as D in hexaploid oats, and that they are more properly designated as DC genome species. 399 400 In the present study, two distinct *Pqk1* homoeologues were detected in each of the three AC(DC) species, with each pair falling consistently into two clusters 401 402 within the C and the A genome clades, respectively (Fig 3). The C-copy sequences of these tetraploids clustered consistently with the C-type homoeologues of the 403 hexaploids, while the A/D genome homoeologues, with the exception of these from 404 A. insularis and one sequence from A. maroccana (Clav 8331) fell into subclade A4 405 25

406	along with a set of sequences from the hexaploid oats (Fig 3). Considering that the
407	other <i>Pgk1</i> gene sequences from the hexaploid oats clustered with the C or A
408	genome diploids, we deduced that the sequences falling in subclade A4 must
409	represent the D genome homoeologues of the hexaploids and of the AC(DC) species
410	A. maroccana and A. murphyi. This result is not fully consistent with our previous
411	GBS study: although A. maroccana and A. murphyi were very similar to hexaploid oat
412	and were designated as DC genomes, our GBS work suggested that A. insularis was
413	also a DC genome that was even more similar to the hexaploids [9]. Examining the
414	existing literature, all three of these tetraploid species have variously been
415	considered as the tetraploid ancestor of the hexaploids [4, 9, 49]. In view of the
416	genome structure of these tetraploids [24, 50] and the meiotic chromosome paring
417	of their interspecific hybrids [51], all of these tetraploids are proposed to have
418	diverged from a common ancestral tetraploid after the occurrence of some large
419	chromosome rearrangements [24, 50]. However, it cannot be ruled out that these
420	tetraploids might have originated independently from different diploid ancestors,
421	since they have shown close relationships with different diploid species [40].
422	Interestingly, in network analysis (Fig 4), the A/D-type homoeologues of A. insularis
423	fell into a group with the A_c genome species A. canariensis and the AB genome
424	species A. agadiriana. In fact, previous studies have revealed that A. canariensis is
425	closely related to the DC genome tetraploids [15]. These results suggest a possibility
426	that A. canariensis was involved in contributing an early version of a D genome in all
427	three AC(DC) genome tetraploids. Nevertheless, we do not have an explanation for ²⁶

428 why the D genome copy of *Pqk1* in *A. insularis* could have diverged so far from the

- 429 version found in the hexaploids, especially since the C genome copies remain
- 430 identical.

432

The genome origins of the hexaploid species. 431

It is now generally accepted that two distinct steps were involved in the evolution of hexaploid oats. The first step would have been the formation of a DC 433 genome hybrid from ancestral D and C genome diploids, followed by doubling of the 434 chromosomes to form an allotetraploid. The second step would have involved 435 436 hybridization of a DC tetraploid with a more recent A genome diploid, followed by

doubling of the triploid hybrid [9, 13]. 437

The diploid progenitor of the hexaploid C genome was probably restricted to 438

439 the narrow geographic range where the three extant C genome diploids are

distributed. However, numerous inter-genomic translocations among hexaploid 440

chromosomes [9, 11, 52, 53] have deceased the homology between the C genome 441

442 diploids and the hexaploid C genome, making the identification of the C genome

443 donor of the hexaploids challenging. In this study, the C_{p} genome species shared the

highest degree of genetic similarity with both the DC genome tetraploids, as well as 444

with the hexaploids, leading us to conclude that a C_p genome species was the C 445

446 genome donor of the polyploids. This conclusion is supported by other evidence

from nuclear genes [13, 54]. This is important, since it was recently demonstrated 447

448 that the maternal tetraploid and hexaploid genomes originated from an A genome

species, not from a C genome species [55], rendering comparisons to the C_v vs C_p 449

450 maternal genomes irrelevant in determining the origin of the nuclear C genome in451 the hexaploids.

452	The A genome origin of the hexaploids remains a matter of debate, and many
453	A genome diploids have been suggested as putative diploid progenitors, as
454	summarized by Peng et al [13]. FISH analysis showed that an A_s -specific DNA repeat
455	was restricted to the A_s and A_l genomes, as well as the hexaploid A genome [18]. In
456	this study, a close relationship between the A_s genome diploid A. atlantica was
457	observed for some hexaploid haplotypes in the phylogenetic tree (Fig 3) and the MJ
458	network (Fig 4). An A. atlantica genome origin is consistent with previous studies
459	based on IGS-RFLP analysis [12] and the <i>ppcB1</i> gene [40]. However, there is evidence
460	in our work that some hexaploids may have an alternate A genome origin, closer to
461	the Agraria section of A_s diploids. The presence of multiple A genome origins could
462	explain variable results that have been reported in studies of hexaploid phylogeny.
463	In this study, strong evidence is presented for a D genome origin in the
464	tetraploids A. maroccana and A. murphyi (Figs 3-4). However, these D genome
465	sequences did not show a close relationship with any diploid species investigated in
466	this study. Other than the discrepancy with A. insularis, this result is consistent with
467	our recent GBS study [9]. One factor that may hinder the discovery of a D genome
468	progenitor is the presence of inter-genomic translations among all three genomes in
469	the hexaploid [9, 53]. Two hexaploid accessions (A.occidentalis CN 25942 and A.
470	sativa PI 194896) did not contribute haplotypes that clustered with the putative D
471	genome sequences (Subclade A4 in Fig 3). Although this may be a result of 28

472 incomplete sampling, it may also result from inter-genomic translations that have473 duplicated or eliminated copies of *Pgk1*.

474	In conclusion, this is the most comprehensive study to date that investigates
475	a phylogeny in genus Avena using a single informative nuclear gene. It confirms or
476	clarifies most previous work, and presents strong evidence in support of a working
477	hypothesis for the origin of hexaploid oat. However, many questions still remain, and
478	these will be best addressed through further studies involving multiple nuclear genes
479	or whole genomes. We are collaborating on work that will provide exome-based
480	gene diversity studies, but this work will require complete hexaploid reference
481	sequences before it can be properly analyzed. Such reference sequences are
482	currently in progress, so the next few years may see a revolution in our
483	understanding of Avena phylogeny. Nevertheless, as many researcher in this field are
484	aware, the polyploid species in this genus have experienced extensive chromosome
485	rearrangement, which will continue to complicate phylogenetic studies. It may even
486	be necessary to generate a pan-genome hexaploid reference sequence before
487	definitive statements can be made.

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