The cuticular wax inhibitor locus $Iw2$ in wild diploid wheat *Aegilops tauschii*: phenotypic survey, genetic analysis, and implications for the evolution of common wheat

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Abstract

**Background:** Cuticular wax production on plant surfaces confers a glaucous appearance and plays important roles in plant stress tolerance. Most common wheat cultivars, which are hexaploid, and most tetraploid wheat cultivars are glaucous; in contrast, a wild wheat progenitor, *Aegilops tauschii*, can be glaucous or non-glaucous. A dominant non-glaucous allele, $Iw2$, resides on the short arm of chromosome 2D, which was inherited from *Ae. tauschii* through polyploidization. $Iw2$ is one of the major causal genes related to variation in glaucousness among hexaploid wheat. Detailed genetic and phylogeographic knowledge of the $Iw2$ locus in *Ae. tauschii* may provide important information and lead to a better understanding of the evolution of common wheat.

**Results:** Glaucous *Ae. tauschii* accessions were collected from a broad area ranging from Armenia to the southwestern coastal part of the Caspian Sea. Linkage analyses with five mapping populations showed that the glaucous versus non-glaucous difference was mainly controlled by the $Iw2$ locus in *Ae. tauschii*. Comparative genomic analysis of barley and *Ae. tauschii* was then used to develop molecular markers tightly linked with *Ae. tauschii* $Iw2$. Chromosomal synteny around the orthologous $Iw2$ regions indicated that some chromosomal rearrangement had occurred during the genetic divergence leading to *Ae. tauschii*, barley, and *Brachypodium*. Genetic associations between specific $Iw2$-linked markers and respective glaucous phenotypes in *Ae. tauschii* indicated that at least two non-glaucous accessions might carry other glaucousness-determining loci outside of the $Iw2$ locus.

**Conclusion:** Allelic differences at the $Iw2$ locus were the main contributors to the phenotypic difference between the glaucous and non-glaucous accessions of *Ae. tauschii*. Our results supported the previous assumption that the D-genome donor of common wheat could have been any *Ae. tauschii* variant that carried the recessive $iw2$ allele.

**Keywords:** Allopolyploid speciation, Cuticular wax inhibitor, Synthetic wheat, Wheat evolution

Background

Cuticular wax production on aerial surfaces of plants has important roles in various physiological functions and developmental events; the wax prevents non-stomatal water loss, inhibits organ fusion during development, protects from UV radiation damage, and imposes a physical barrier against pathogenic infection [1-4]. The trait, the coating of leaf and stem surfaces with a waxy whitish substance, is called glaucousness. In common wheat (*Triticum aestivum* L., 2n = 6x = 42, genome constitution BBAADD), dominant alleles $W1$ and $W2$, control the wax production and have been assigned to chromosomes 2B and 2D, respectively [5,6]. Additionally, dominant homoeoalleles for non-glaucousness, $lw1$ and $lw2$, have also been mapped to the short arms of chromosomes 2B and 2D, respectively [6-9]. Wheat plants with either the $w1$, $w2$, $lw1$ or $lw2$ allele show the non-glaucous phenotype, indicating that $W1$ and $W2$ are functionally redundant for the glaucous phenotype and that a single $lw$ dominant allele is sufficient to inhibit the glaucous
phenotype even in the presence of a W1 or W2 allele [3,6]. Wax composition in wheat plants with one Iw dominant allele is biochemically different from that in glaucous plants of any genotype; ß-diketones are completely absent from extracts of cuticular wax from Iw plants, while aldehydes and primary alcohols are very abundant in these extracts [3,10]. A fine map around the Iw1 region on 2BS was constructed using an F2 population of tetraploid wheat (Triticum turgidum L., 2n = 4x =28, BBAA), and three markers tightly linked to Iw1 were developed [10,11]. A high-resolution map of Iw2 on 2DS has been developed in hexaploid wheat, and two markers tightly linked to Iw2 were also developed [11]. Comparative mapping of Iw1 and Iw2 shows that the two loci are homoeologous to each other and orthologous to the same chromosomal region of Brachyypodium distachyon (L.) P. Beauv. [11]. Recently, a third wax-inhibitor locus Iw3 was identified on chromosome 1BS from wild emmer wheat [12], and a fine map of the Iw3 locus is available [13]. Iw2 is located on 2DS in Aegilops tauschii Coss. (2n = 2x = 14, DD), which is diploid and the progenitor of the D-genome of common wheat [14], but to our knowledge, a high-resolution genetic map of the Iw2 region in Ae. tauschii has not been constructed.

Common wheat is an allohexaploid species derived from interspecific hybridization between tetraploid wheat with a BBAA genome and Ae. tauschii. Most cultivated varieties of tetraploid wheat are glaucous, even though non-glaucous types are frequently found among wild tetraploid accessions [6,15]; this variation indicates that the glaucous phenotype might have been a target of artificial selection during the domestication of tetraploid wheat. Glaucous accessions of Ae. tauschii are found in the area ranging from Transcaucasia to the southern coastal region of the Caspian Sea [5,16]. Almost all varieties of common wheat carry W1 and W2 and lack Iw1 and Iw2; therefore, the D-genome donor of common wheat is assumed to have had the recessive iw2 allele [5]. Glaucous Ae. tauschii accessions have the W2 and iw2 alleles. Non-glaucous accessions of Ae. tauschii that have the W2 and iw2 alleles have been recovered from a wide distribution range in central Eurasia [5]. Moreover, discovery of a non-glaucous Ae. tauschii accession with the w2 recessive allele has not yet been reported.

Therefore, analysis of the Iw2 locus may provide important information that improves our understanding of the evolution of common wheat. Population structure analyses of Ae. tauschii indicate that the whole species Ae. tauschii can be divided into three major genealogical lineages, tauschii lineage 1 (Taul1), Taul2, and Taul3, and that genetically genomes of Taul2 accessions are most closely related to the D genome of common wheat [17-19]. Recently, a whole-genome shotgun strategy was used to generate a draft genome sequence of Ae. tauschii that has been published; this draft anchors 1.72 Gb of the 4.36 Gb genome to chromosomes [20]. A physical map of the Ae. tauschii genome that covers 4 Gb is also available [21]. The objectives of this study were (1) to examine the natural variation in glaucousness among a species-wide set of Ae. tauschii accessions, (2) to use F2 populations of Ae. tauschii accessions and synthetic hexaploid wheat lines to fine-map Iw2 locus on 2DS, (3) to develop molecular markers that are closely linked to Iw2 based on chromosomal synteny between barley and wheat chromosomes, and (4) to provide novel insights into the evolutionary relationship between the Ae. tauschii genome and the D genome of common wheat on the basis of the detailed genetic and phylogeographic knowledge of the Iw2 chromosomal region.

Methods

Plant materials and phenotype evaluation

In all, 210 Ae. tauschii accessions were used in this study [22]. Their passport data, including geographical coordinates, have been provided in previous reports [23,24]. Previously, 206 of the Ae. tauschii accessions were grouped into the three lineages, TauL1, TauL2, and TauL3, based on DArT marker genotyping analysis [19]. Of the 210 accessions, 12 were previously identified as subspecies strangulata based on the sensu-strico criteria [25,26]. Seeds from two Ae. tauschii hybrid F2 populations (n = 116 from each population) were sown in November 2011; one F2 population resulted from a cross between KU-2154 (non-glaucous) and KU-2126 (glaucous), the other from a KU-2003 (non-glaucous) by KU-2124 (glaucous) cross. In the 2012–2013 season, 169 additional F2 individuals of the KU-2154/KU-2126 population were grown to increase the size of the mapping population.

Previously, 82 synthetic hexaploid wheat lines were produced from crosses between a tetraploid wheat (T. turgidum subspecies durum (Desf.) Husn.) cultivar Langdon (Ldn) and 69 Ae. tauschii accessions [26,27]. These synthetic hexaploid wheat lines were used for crossing and phenotypic studies conducted in a glasshouse at Kobe University. Ldn shows the glaucous phenotype and is homozygous for the iw1 allele [10]. Each synthetic hexaploid thus contained the A and B genomes from Ldn and one of many diverse D genomes originating from the Ae. tauschii pollen parents. In the present study, four F3 plants derived from one F2 plant of each synthetic hexaploid were grown individually during the 2007–2008 season in pots that were arranged randomly in the glasshouse; these 276 F3 plants were used for crossing and phenotypic observation. The following three pairs of synthetic hexaploids were used to generate three F3 mapping populations: Ldn/P1476874 (non-glaucous) and Ldn/KU-2069 (glaucous), Ldn/JG126387 (non-glaucous) and Ldn/KU-2159 (glaucous), and Ldn/KU-2124 (glaucous)
and Ldn/IG47259 (non-glaucous). The first population (Ldn/P1476874//Ldn/KU-2069) comprised 106 F2 individuals grown in the glasshouse during the 2009–2010 season. Seeds from the other two populations were sown in November 2011, with the numbers of individuals in each being 100 (Ldn/KU-2159//Ldn/IG126387) and 82 (Ldn/KU-2124//Ldn/IG47259).

For analysis of the D genome of common wheat, 17 landraces collected in Iran were supplied from the National BioResource Project (NBRP) KOMUGI (http://www.shigen.nig.ac.jp/wheat/komugi). These Iranian landraces—KU-3097, KU-3098, KU-3121, KU-3126, KU-3136, KU-3162, KU-3184, KU-3189, KU-3202, KU-3232, KU-3236, KU-3274, KU-3289, KU-10393, KU-10439, KU-10480, and KU-10510—each showed the glaucous phenotype.

Glaucousness was evaluated based on the presence or absence of wax production on the surface of peduncles and spikes in both Ae. tauschii and synthetics. Wax production was clearly visible and whitish.

**Genotyping and construction of linkage maps**

To amplify PCR fragments containing molecular markers, some of which were simple sequence repeats (SSRs), total DNA was extracted from leaves of the parental strains and F2 individuals. For SSR genotyping, 40 cycles of PCR were performed using 2x Quick Taq HS DyeMix (TOYOBO, Osaka, Japan) and the following conditions: 10 s at 94°C, 30 s at the respective annealing temperature (72, 73, or 75°C), and 30 s at 68°C. The last step was a 1-min incubation at 68°C. Information on SSR markers and the respective annealing temperatures was obtained from the NBRP KOMUGI web site (http://www.shigen.nig.ac.jp/wheat/komugi). These Iranian landraces—KU-3097, KU-3098, KU-3121, KU-3126, KU-3136, KU-3162, KU-3184, KU-3189, KU-3202, KU-3232, KU-3236, KU-3274, KU-3289, KU-10393, KU-10439, KU-10480, and KU-10510—each showed the glaucous phenotype.

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For genotyping; two STS markers for each locus, and these markers were previously developed along with the Iw2-linked markers [7]. The amplified PCR products were separated via electrophoresis through a 2% agarose or 13% non-denaturing polyacrylamide gel and then stained with ethidium bromide.

**Development of additional markers linked to Iw2**

In our previous studies, we conducted deep-sequencing analyses of the leaf and spike transcriptomes of two Ae. tauschii accesses that represented two major lineages, and discovered more than 16,000 high-confidence single nucleotide polymorphisms (SNPs) in 5,808 contigs [31,32]. Contigs with the SNPs were searched with blastn against Ae. tauschii genome sequences [20] and barley genome sequences [33]; these genome sequences included high-confidence genes with an E-value threshold of $10^{-5}$ and hit length ≥ 50 bp, fingerprinted contigs, and whole genome shotgun assemblies.

To choose scaffolds for Ae. tauschii sequences throughout the Iw2 chromosomal region, all the genes contained in each scaffold were searched with blastn against the barley genomic sequence using parameters described above. Scaffolds containing at least one gene aligned on the distal region of chromosome 2HS (between 3.66 Mb and 5.51 Mb) were considered possible candidates for marker development. Scaffolds without genes were anchored based on respective results from the blastn searches against the barley genome. First, high-confidence SNPs [31,32] plotted in this 2HS chromosomal segment were used for marker development to refine the target region. Next, SciRoKo version 3.4 [34] was used with search mode setting “mismatched; fixed penalty” to identify additional SSR markers in sequence data of candidate scaffolds. Additional SNPs were also identified on candidate scaffolds by sequencing approximately 700 bp of amplified DNA of two Ae. tauschii accesses, KU-2154 and KU-2126. The nucleotide sequences were determined using an Applied Biosystems 3730xl DNA Analyzer (Applied Biosystems, Foster City, CA, USA), and SNPs were found via sequence alignments constructed and searched with GENETYX-MAC version 12.00 software (Whitehead Institute for Biomedical Research, Cambridge, MA, USA).

For genotyping, total DNA was extracted from leaves taken from each of the 210 Ae. tauschii accesses and the 17 Iranian wheat landraces. SSR amplification and detection of polymorphisms at these loci were conducted as described above. The identified SNPs were then further developed into cleaved amplified polymorphic sequence (CAPS) or high resolution melting (HRM) markers. The primer sequences for each SNP marker and any relevant restriction enzymes are summarized in Additional file 1. PCR and subsequent analyses were performed as described previously [31,32,35].
Blast analysis of the *Ae. tauschii* genes relative to the *Brachypodium* genome

Nucleotide sequences and annotation information of the selected *Ae. tauschii* scaffolds were analyzed with reference to the *Ae. tauschii* draft genome data, which was published by Jia et al. [20]. Reference sequences from *Brachypodium* [36] were searched against the National Center for Biotechnology Information (NCBI) NR protein database using the blastx algorithm with an E-value cut-off of $10^{-3}$.

Association analysis of the linked markers with glaucousness

The Q + K method was conducted using a mixed linear model (MLM) function in TASSEL ver 4.0 software [37] for an association analysis by incorporating phenotypic and genotypic data and information on population structure. In a previous report, the Bayesian clustering approach implemented in the software program STRUCTURE 2.3 [38] was used with the setting $k = 2$ to predict the population structure of the *Ae. tauschii* accessions [19]. The Q-matrix of population membership probabilities was served as covariates in MLM. Kinship (K) was calculated in TASSEL based on the genotyping information of the 169 DArT markers for the 206 *Ae. tauschii* accessions [19]. We performed the $F$-statistics and calculated the $P$-values for the $F$-test, and the threshold value was set as 1E-3 for the significant association. We omitted the target markers from the association analysis when their minor allele frequencies were less than 0.05.

Results

Wax production variation among *Ae. tauschii* accessions and among synthetic wheat lines

Of the 210 *Ae. tauschii* accessions examined, only 20 (9.5%) exhibited the glaucous phenotype and produced whitish wax on the surfaces of peduncles and spikes (Figure 1A-D, Additional file 2). Wax production for each accession was completely consistent between the Fukui and Kobe environments. Each glaucous accession belonged to *Ae. tauschii* subspecies *tauschii*; in other words, none belonged to *Ae. tauschii* subspecies *strangulata*; the geographic distribution of glaucous accessions was limited to the area that spans from Transcaucasia to the southern coastal region of the Caspian Sea (Figure 1H). In the eastern habitats (central Asia, Afghanistan, Pakistan, India, and China) of the species range, no glaucous accession was found. Of the 20 glaucous accessions, 19 belonged to the TauL2 lineage, and only one (IG127015 collected in Armenia) belonged to the TauL1 lineage (Additional file 2).

Of the 82 synthetic wheat lines that we examined, 15 exhibited whitish wax production on the peduncle and spike surface (Figure 1E-G), whereas no wax production was evident in any of the 67 other lines (Additional file 2). Of the 15 lines that showed the glaucous phenotype, 13 were produced by crossing Ldn with glaucous *Ae. tauschii* accessions, and each of the 67 non-glaucous lines was produced by crossing Ldn with a non-glaucous *Ae. tauschii* accession. Notably, two synthetic lines, Ldn/KU-2104 and Ldn/KU-2105, exhibited the glaucous phenotype even though their parental *Ae. tauschii* accessions were non-glaucous.

Mapping of the *Iw2* locus in *Ae. tauschii* and synthetic wheat

Two *F2* populations of *Ae. tauschii* and three *F2* populations from the synthetic wheat lines were analyzed to map the loci that control inhibition of wax production. Each *F1* plant used for the five cross combinations exhibited the non-glaucous phenotype. In each *F2* population, the ratio of non-glaucous to glaucous individuals was 3:1; these findings were statistically significant and consistent with Mendelian segregation of alleles of a single gene (Table 1). These results indicated that a single genetic locus was associated with the phenotypic difference between non-glaucous and glaucous surfaces on peduncles and spikes, and that allele conferring the non-glaucous phenotype was dominant and the allele conferring the glaucous phenotype was recessive.

A single locus that controlled inhibition of wax production in *Ae. tauschii* was mapped to the same region of the short arm of chromosome 2D in each *F2* mapping population (Figure 2). In the KU-2003/KU-2124 population, the locus that controlled inhibition of wax production, together with the loci for 25 SSR markers and *Ppd-D1*, was assigned to chromosome 2D, and the map length was 230.0 cM with an average inter-loci interval of 8.85 cM. In the KU-2154/KU-2126 population, the locus that controlled inhibition of wax production, together with 14 SSR and 2 STS markers and *Ppd-D1*, was assigned to chromosome 2D, and the map length was 175.4 cM with average inter-loci spacing of 10.32 cM. In the three synthetic wheat populations, Ldn/KU-2159/Ldn/IG126387, Ldn/KU-2124/Ldn/IG47259, and Ldn/Pl476874/Ldn/KU-2069, the locus that controlled inhibition of wax production was mapped to a similar position on the short arm of chromosome 2D (Figure 2). In these three synthetic wheat populations, the locus that controlled inhibition of wax production was mapped together with 11 to 13 SSR markers, 0 to 2 STS markers, and *Ppd-D1*; additionally, the map lengths ranged from 79.4 to 93.8 cM with an average inter-loci spacing of 4.96 to 8.53 cM.

*WE6* and *TE6* are EST-derived STS markers that are linked to *Iw2* in two mapping populations [7,9]. In three of our mapping populations, linkage of the non-glaucousness loci to *WE6* and *TE6* were confirmed. Thus, the position of one locus that controlled inhibition of wax production in *Ae. tauschii* corresponded to the well-known wax inhibitor.
Table 1 Segregation analysis of the non-glaucous phenotype in the five F$_2$ mapping populations

| F$_2$ population     | N  | Non-glaucous type | Glaucous type | $\chi^2$ value* | P value |
|----------------------|----|-------------------|---------------|-----------------|---------|
| KU-2003/KU-2124      | 116| 89                | 27            | 0.184           | 0.668   |
| KU-2154/KU-2126      | 116| 78                | 38            | 3.724           | 0.054   |
| Ldn/KU-2159/Ldn/IG126387 | 100| 71                | 29            | 0.853           | 0.356   |
| Ldn/KU-2124/Ldn/IG47259 | 82 | 65                | 17            | 0.797           | 0.372   |
| Ldn/PI476874/Ldn/KU-2069 | 106| 77                | 29            | 0.314           | 0.575   |

*Expected segregation ratio was 3:1.
gene, \textit{Iw2}, on chromosome 2D \cite{6, 7}. Therefore, hereafter, all glaucousness-related loci mapped in this study were considered to be identical to \textit{Iw2}.

**Fine mapping of the \textit{Iw2} locus**

The high-confidence SNPs derived from \textit{Ae. tauschii} RNA-seq data have been plotted onto barley chromosomes \cite{32}, and physical map information for the barley genome is available \cite{33}. Additionally, physical map information for \textit{Ae. tauschii} and 16,876 scaffolds that constitute 1.49 Gb from the draft \textit{Ae. tauschii} genome sequence are anchored to the \textit{Ae. tauschii} linkage map \cite{20, 21}. The RNA-seq-derived SNP information \cite{31, 32} was used to map seven high-confidence SNPs, represented as \textit{Xctg} loci in Figure 3, throughout the \textit{Iw2} chromosomal region in the KU-2154/KU-2126 F\textsubscript{2} population. Of the seven \textit{Xctg} loci, four were located within the 8.8 cM chromosomal region immediately surrounding \textit{Iw2}. Nucleotide sequences of the four cDNAs corresponding to these \textit{Xctg} loci were used as queries to select the carrier scaffolds from \textit{Ae. tauschii} sequences.

We selected the \textit{Ae. tauschii} scaffolds that mapped near the \textit{Xctg}-carrying \textit{Ae. tauschii} scaffolds based on synteny between the wheat and barley genomes and the barley physical map \cite{39}. In all, 18 \textit{Ae. tauschii} scaffolds were assigned \textit{in silico} to an area of the \textit{Ae. tauschii} genome that corresponded to the \textit{Iw2} region in the physical map of barley chromosome 2H (Figure 3). Using a previously developed physical map of the \textit{Ae. tauschii} 2DS chromosome \cite{21}, we mapped six \textit{Ae. tauschii} scaffolds \textit{in silico} to the corresponding region in the 2DS physical map. Nucleotide sequences of the selected scaffolds were used to design CAPS or SSR markers for each scaffold, and the markers that were polymorphic between KU-2154 and KU-2126 were then mapped in the F\textsubscript{2} population (Figure 3).

Of the selected scaffolds, 23 were mapped to the \textit{Iw2} chromosomal region on 2DS, and the remaining three scaffolds were assigned to other chromosomes. In the KU-2154/KU-2126 population with 115 F\textsubscript{2} individuals, the \textit{Iw2} locus was mapped within the 1.1 cM interval between the most closely linked markers (Figure 3). A dominant
marker (SS1038-8), derived from the Ae. tauschii scaffold 51038 sequence, was located 0.2 cM distal to Iw2, and the WE6 SSR marker was located 0.9 cM proximal to Iw2. Five co-dominant markers, derived from two Ae. tauschii scaffolds 10812 and 82981, co-localized with Iw2. The marker order in the KU-2154/KU-2126 linkage map was generally conserved with that in the barley 2H physical map. However, barley scaffold 9655 was more closely linked to the barley Iw2 ortholog than were two corresponding Ae. tauschii scaffolds, 13577 and 33766, to the tauschii Iw2 ortholog; this positioning indicated that a local inversion had occurred in the region proximal to Iw2 during the divergence between barley and tauschii.

Next, F₂ individuals of the KU-2154/KU-2126 population and 12 markers from five Ae. tauschii scaffolds were used to construct a fine map of Iw2 (Figure 4A). Based on this linkage map, Iw2 was located within the 0.7 cM between Xctg216249/SS1038-8 and WE6 and co-localized with five markers derived from two scaffolds, 10812 and 82981. Each of the five scaffolds was 63 to 334 kb in length and included one to 16 putative protein-coding genes [20,21]; marker positions of each scaffold are indicated in Figure 4B. Of the 12 markers, eight were derived from intergenic regions, the other four from open reading frames.

In all, 36 genes were evident on the five scaffolds, and gene annotation could be confirmed for 27 of the 36 genes (Table 2). Of these 27 Ae. tauschii genes, 10 putatively encoded cytochrome P450 monooxygenase proteins, and eight encoded disease-related proteins. Additionally, genes encoding laccase, agmatine coumaroyltransferase, receptor...
kinase, and cell number regulator 2-like were found on the
two scaffolds that co-localized with *Iw2*.

The *Ae. tauschii* scaffolds that included protein coding
genes were used as queries to search the *Brachypodium*
genomic information via a blastn search. Of the *Ae. tauschii*
genes on the five scaffolds, 18 had obvious
orthologs in the *Brachypodium* genome (Figure 4C).
Putative orthologs of the *Ae. tauschii* genes from the four
scaffolds were assigned to the 987 to 1068 kb region
of *Brachypodium* chromosome 5. In addition, three
*Brachypodium* paralogs (Bradi5g01220.1, Bradi5g01220.2,
and Bradi5g01230.1) positioned in the 1133 to 1143 kb
region were orthologous to an *Ae. tauschii* gene, AEGT
A20985; additionally, Bradi5g01280.1 at 1186 kb was ortho-
logous to AEGTA28084 in scaffold 6859. The locations of
two *Ae. tauschii* genes, AEGTA20985 and AEGTA28084,
were 3 and 3.9 cm, respectively, distal to *Iw2* (Figure 3);
therefore, the distal part of *Iw2* showed chromosomal
synteny to *Brachypodium* chromosome 5. Thus, the
*Iw2* chromosomal region on 2DS was generally syntenic
to *Brachypodium* chromosome 5. However, putative
orthologs of the *Ae. tauschii* genes from scaffold 43829
were assigned to *Brachypodium* chromosomes 1 and 2. Two paralogous *Ae. tauschii* genes, AEGTA19771 and AEGTA19772, on scaffold 10812 were orthologous to three paralogous *Brachypodium* genes (Bradi3g02290.1, Bradi3g02300.1, and Bradi3g02370.1) on *Brachypodium* chromosome 3. Therefore, the chromosomal synteny between *Ae. tauschii* and *Brachypodium* around the *Iw2* orthologs was complex with regard to chromosome structure.

**Iw2-linked marker genotypes in *Ae. tauschii***

To determine the genetic associations among the developed markers and glaucousness, 13 *Iw2*-linked PCR markers—including five CAPSs, five SSRs, one HRM, one insertion/deletion (indel), and one dominant (presence or absence) marker—were used to genotype the 210 *Ae. tauschii* accessions (Table 3). For eight of the 13 markers, the 210 accessions exhibited just two apparent alleles; additionally, the set of accessions exhibited just three distinct electrophoresis patterns—including the KU-2154-type, the KU-2126-type, and one other type—at one SSR marker for *WE6*. The other four SSR markers were highly polymorphic among the accessions; specifically, each marker gave rise to more than three distinct electrophoresis patterns.

### Table 2 Colinearity between *Ae. tauschii* and *Brachypodium* in the syntenic genomic regions around *Iw2* (Continued)

| *Ae. tauschii* gene | *Brachypodium* gene | Annotation |
|---------------------|---------------------|------------|
| AEGTA17439          | Bradi5g01135.1      | probable pectate lyase 15-like |
| AEGTA17438          | Bradi5g01110.1      | disease resistance rpp13-like protein 1-like |
| AEGTA17437          | Bradi5g01070.1      | disease resistance rpp13-like protein 1-like |
| AEGTA17436          | Bradi5g01080.1      | disease resistance rpp13-like protein 1-like |
| AEGTA17435          | Bradi5g01110.1      | disease resistance rpp13-like protein 1-like |
| AEGTA17434          | Bradi5g01080.1      | disease resistance rpp13-like protein 1-like |
| AEGTA23449          | Bradi5g01110.1      | disease resistance rpp13-like protein 1-like |
| AEGTA03244          | Bradi5g01110.1      | disease resistance rpp13-like protein 1-like |

**Table 2 Colinearity between *Ae. tauschii* and *Brachypodium* in the syntenic genomic regions around *Iw2***

| *Ae. tauschii* gene | *Brachypodium* gene | Annotation |
|---------------------|---------------------|------------|
| AEGTA20795          | Bradi1g15030.1      | cytochrome p450 85a1 |
| AEGTA20794          | Bradi1g15030.1      | cytochrome p450 85a1 |
| AEGTA25164          | Bradi1g15030.1      | cytochrome p450 85a1 |
| AEGTA22963          | Bradi1g15030.1      | cytochrome p450 85a1 |
| AEGTA20793          | Bradi1g15030.1      | cytochrome p450 85a1 |
| AEGTA20792          | f-box domain containing protein |
| AEGTA04539          | hypothetical protein F775_04539 |
| AEGTA09742          | Bradi1g15010.1      | probable fructokinase-1-like |
| AEGTA20791          | Bradi2g39120.1      | hypothetical protein F775_20791 |
| AEGTA20790          | Bradi2g39100.1      | hypothetical protein F775_20791 |
| AEGTA32301          | Bradi3g18920.1      | cytochrome p450 |
| AEGTA20789          | cytochrome p450 monoxygenase cyp71d70 |
| AEGTA20788          | cytochrome p450 |
| AEGTA09741          | cytochrome p450 71c4 |
| AEGTA09740          | sulfotransferase 16-like |
| AEGTA32300          | deleted in split hand split foot protein 1 |
| AEGTA24906          | brach planthopper-induced resistance protein 1 |
| AEGTA19771          | cytochrome p450 84a1 |
| AEGTA19777          | laccase-15-like |
| AEGTA17435          | disease resistance rpp13-like protein 1-like |
| AEGTA17434          | disease resistance rpp13-like protein 1-like |
| AEGTA23449          | hypothetical protein F775_23449 |
| AEGTA03244          | hypothetical protein F775_03244 |
The association analysis showed that four SSR markers (S43829-13, S43829-12, S10812-1, and S82981-2), an HRM marker (Xctg216249), the dominant marker (S51038-8), and an indel marker (S10812-14), and two CAPS markers (S10812-12, and S10812-13) co-localized with Iw2 in the Ae. tauschii linkage map, were significantly (P < 1E-3) associated with variation in glaucousness; in contrast, the other three genotyped markers were not significantly associated with variation in glaucousness (Table 3). The CAPS marker S43829-3 was removed from this association analysis because of the low-frequency (<0.05) allele. In particular, the KU-2126-type allele of the SSR locus S10812-1 was found only in 15 of the 20 glaucous accessions; moreover, none of the 190 non-glaucous accessions carried this KU-2126-type allele. The other five glaucous accessions carried a third allele of the S10812-1 locus. In 55 of the 190 non-glaucous accessions, only four carried the third allele of the S10812-1 locus, and the other 135 accessions carried different S10812-1 alleles. Of the four exceptional non-glaucous accessions that carried the third S10812-1 allele, two were KU-2104 and KU-2105, and these had each been used to generate a synthetic hexaploid wheat line Ldn/KU-2104 and Ldn/KU-2105, respectively; both synthetic lines showed the glaucous phenotype (Additional file 2). However, the phenotype of each synthetic hexaploid line (Ldn/KU-2074 and Ldn/KU-2079) derived from the remaining two of the exceptional accessions (KU-2074 and KU-2079) was non-glaucous.

Therefore, phenotypic differentiation in glaucousness was almost completely explained by the allelic configuration at the S10812-1 locus in these natural populations of Ae. tauschii.

The 17 Iranian wheat landraces showed the KU-2154-type alleles at S43829-3, Xctg216249, and S51038-8, whereas they exhibited the KU-2126-type alleles at C141566873, S10812-12, S10812-14, S10812-13, and Xctg202354. In addition, these landraces exhibited various genotypes that differed from the allelic combinations found in Ae. tauschii accessions at WE6 and four SSR marker loci, S43829-13, S10812-1, S82981-2 (Table 3). At S43829-12, 15 landraces showed the KU-2126-type genotype, and two exhibited other genotypes.

**Discussion**

**Natural variation for wax production in Ae. tauschii**

Glaucousness is presumably among the components of the domestication syndrome in tetraploid wheat [5,6]. Therefore, glaucousness was apparently a target of artificial selection during tetraploid domestication and common wheat speciation; nevertheless, whether glaucousness is an adaptive trait in wild wheat species remains unclear. Cuticular wax on plant surfaces plays an important role in reducing water loss under drought stress conditions for Arabidopsis and rice [1,4], and observations in these other species indicate that relationships between glaucousness and drought stress tolerance are tight. Presence of either
the *Iw1* or *Iw2* allele greatly reduces β-diketones in the wax components of plants, resulting in a non-glaucous phenotype [3,10]. Comparative study of glaucousness-related genes in near-isogenic lines (NILs) of a common wheat cultivar (S-615) (BC10F3 generation; [6]) demonstrates that *Iw* alleles had a negative impact on drought tolerance [3]. However, another study of *Iw1* in a NIL (BC2F3 generation) of common wheat did not detect an association between *Iw1* genotype and water-use efficiency [10].

In this study, we used a set of 210 accessions that represented the entire geographical range of *Ae. tauschii* to examine natural variation in wax production among *Ae. tauschii*, and found 20 glaucous accessions that were collected in the area that spans from Transcaucasia to the southern-eastern coastal region of the Caspian Sea (Figure 1, Additional file 2). In a previous study of 176 *Ae. tauschii* accessions collected from 105 different habitats throughout Afghanistan, Pakistan, and Iran, 17 glaucous accessions were found in this same area that spans from Transcaucasia to the southern-eastern coastal region [16]. Therefore, our findings were fully consistent with previous observations.

Most glaucous accessions belonged to the TauL2 lineage (Additional file 2). TauL2 accessions derived from geographically wide-spread sites throughout the Transcaucasia/Middle East region; these sites represented the western habitats of *Ae. tauschii* [19]. TauL1 accessions were collected from sites widely distributed throughout the species range, and most TauL1 accessions showed a non-glaucous phenotype. Notably, one TauL1 accession (IG127015), collected in Armenia, showed a glaucous phenotype, and the collection site was located in the middle of an area where glaucous TauL2 accessions were collected (Figure 1). Genotyping data suggested that IG127015 had an *Iw2* chromosomal region that was very similar to the *Iw2* chromosomal region of the glaucous TauL2 accessions. One possible explanation for this observation is that IG127015 acquired the *Iw2* chromosomal region from some glaucous individual of the TauL2 lineage. Such introgression could occur in the natural habitat where IG127015 was originally sampled and in experimental fields where the accession was propagated for several generations. Another explanation is that IG127015 became a wax producer through a *de novo* recessive mutation at the *Iw2* locus; this scenario, however, is unlikely because the molecular marker genotypes in the *Iw2* chromosomal region of IG127015 were largely identical to those in the *Iw2* chromosomal region of glaucous TauL2 accessions (Table 3).

Whether the glaucous phenotype of the exceptional TauL1 accession was due to introgression of a glaucous allele from a glaucous TauL2 plant may be difficult to discern. Genome-wide marker analyses using SNP array and diversity arrays technology (DArT) systems indicated that TauL2 was clearly distinct and genetically differentiated from TauL1 [18,19]. This high level of differentiation indicates that the two genealogical lineages have been reproductively isolated, and that, under natural conditions, inter-lineage hybridization seems to have occurred only rarely [17,18]. Nevertheless, the presence of a glaucous-type TauL1 accession indicated that the hybridization between TauL1 and TauL2 might have occurred, but the number of hybridizations seems to be quite small. Further detailed study is required to clarify the past occurrence of the TauL1-TauL2 inter-lineage hybridization in *Ae. tauschii*.

Causal loci for variation in glaucousness among *Ae. tauschii*

Previous studies show that, in *Ae. tauschii*, the causal gene for the glaucous/non-glaucous phenotypic difference is *Iw2*, and that the genotypes of glaucous and non-glaucous accessions were *W2W2Iw2Iw2* and *W2W2Iw2Iw2*, respectively [5,14]. The molecular markers tightly linked to *Iw2* were very closely associated with glaucous versus non-glaucous phenotypic difference among the 210 accessions of *Ae. tauschii* (Table 3). Thus, the allelic difference at the *Iw2* locus was the main contributor to the phenotypic difference between the glaucous and non-glaucous accessions of *Ae. tauschii* (Figure 2, Table 3). In common wheat, the markers derived from Bradi5g01160 are tightly linked to *Iw2* as well as *Iw1* [10,11]. Because the loci that control the glaucous versus non-glaucous phenotypic difference in *Ae. tauschii* mapped to the chromosome 2DS region where the common wheat *Iw2* gene resides (Figure 4), the same *Iw2* gene is likely involved in wax production in both *Ae. tauschii* and common wheat. Actually, although most SSR markers around the *Iw2* region were highly polymorphic among the *Ae. tauschii* accessions and Iranian wheat landraces, three markers co-localized with *Iw2* in *Ae. tauschii* S10812-12, S10812-14, and S10812-13; notably, each showed the KU-2126-type alleles in each of the 17 Iranian wheat landraces (Table 3). These results indicated that the Iranian wheat landraces, which exhibited the glaucous phenotype, had the *i2wiw2* genotype.

Marker order and gene order around *Ae. tauschii* *Iw2* was well conserved with those on barley chromosome 2HS and *Brachypodium* chromosome 5 (Figures 3 and 4). Similar chromosomal synteny between the *Iw1* region on 2BS and *Brachypodium* chromosome 5 was recently reported based on mapping with common wheat populations [10,11]. In *Ae. tauschii*, scaffold information derived from the draft genome data were available for detailed analysis of chromosomal synteny at the *Iw2* region. Chromosomal order of the selected scaffolds at *Iw2* revealed the occurrence of a local inversion during divergence between barley and *Ae. tauschii* (Figure 3). Moreover, information of predicted genes in the scaffolds showed that putative translocations occurred during
divergence between *Brachypodium* and *Ae. tauschii* (Figure 4). These results also indicated that several gaps existed between the *Ae. tauschii* scaffolds. Thus, colinearity among barley, *Brachypodium* and *Ae. tauschii* was observed in the *Iw2* syntenic region, as was reported recently [11], but further screening of *Ae. tauschii* BAC clones may be required for construction of the complete physical map at *Iw2*.

In the genotyping analysis with *Iw2*-linked markers, non-glaucous accessions with the *Iw2Iw2* genotype constituted the majority of all 210 accessions (Table 3). However, four non-glaucous accessions (KU-2074, KU-2079, KU-2104, and KU-2015) shared a genotype at *S10812-1* (the most tightly linked marker) with five glaucous accessions (IG127015, KU-2106, KU-2158, KU-2159, KU-2160), indicating that these four non-glaucous accessions may have the *Iw2Iw2* genotype in spite of the *S10812-1* genotypes. In fact, synthetic hexaploids from hybrids between Ldn, which has the glaucous genotype (*W1W1Iw1Iw1*) [10], and two of the four non-glaucous accessions, KU-2104 and KU-2105, exhibited the glaucous phenotype (Additional file 2). In contrast, the phenotypes of all synthetic hexaploids derived from the KU-2074 and KU-2079, were non-glaucous. Accordingly, KU-2074 and KU-2079 seemed to have the *Iw2Iw2* genotype even though they shared an *S10812-1* genotype with the five glaucous accessions. Taken together, all this evidence indicated that *Iw2* was the major gene that controls inhibition of wax production in *Ae. tauschii*.

As yet, no loss-of-function allele has been reported for *W2*, a major wax-producing gene in *Ae. tauschii*. In common wheat, however, some cultivars such as Chinese Spring and Salmon carry the recessive *w2* allele [5]. Similarly, non-glaucous-type accessions with the *w1* recessive allele have been discovered among wild emmer wheat [5]. Whether the recessive loss-of-function mutation occurred at the diploid level (i.e., in *Ae. tauschii*) or at the hexaploid level (i.e., in *T. aestivum*) is not known. Further studies are required to clarify the details of the genetic mechanism that underlies the wax production in *Ae. tauschii*.

**Implication of the *Iw2* variation in hexaploid wheat speciation**

Based on a comparative genic analysis among common wheat and its ancestral species, Tsunewaki [5] suggested that common wheat, which is hexaploid, is the product of a hybrid cross that took place between a glaucous cultivated emmer wheat with the genotype *W1W1Iw1Iw1* and a glaucous wild *Ae. tauschii* with the genotype the *W2W2Iw2Iw2* genotype in the mountainous region near the southwestern coastal part of the Caspian Sea. Here, we found that, of 210 *Ae. tauschii* accessions, only 20 had the glaucous phenotype (Additional file 2) and that a dominant allele at the *Iw2* locus were responsible for expression of the non-glaucous phenotype (Table 1). Furthermore, we found that, on the basis of the molecular-marker genotypes in the *Iw2* chromosomal region and the phenotypes of the synthetic common wheat lines, virtually all non-glaucous accessions had the *Iw2Iw2* genotype (Table 3, Additional file 2). A non-glaucous accession that had the *Iw2Iw2* genotype was not found among the 210 accessions. This finding was notable because the double recessive *w2w2Iw2Iw2* genotype, if present, would have also caused the non-glaucousness phenotype. The reason for the absence of any *Ae. tauschii* accession with the *w2w2 Iw2Iw2* genotype from this collection was not clear, but this fact may indicate that functional *W2* alleles confer some adaptive advantage under natural conditions. Taken together, the evidence from this study was consistent with the view that glaucous *Ae. tauschii* individuals that had the *W2W2Iw2Iw2* genotype were involved in the origin of hexaploid common wheat.

Previous evidence based on isozyme variations and DNA marker polymorphisms is consistent with the hypothesis that the birthplace of hexaploid wheat is within a broad area ranging from Armenia to southwestern Caspian Iran [18,40-42]. The geographic range of the parent populations of glaucous *Ae. tauschii* accessions was very consistent with the region postulated in this hypothesis (Figure 1). However, the *Ae. tauschii* subspecies-*strangulata* has been postulated to be the D-genome donor of common wheat [43]. Of the 210 *Ae. tauschii* accessions that we examined, only 12 accessions have markedly moniliform spikes, and each of these were originally collected in the southeastern coastal Caspian region [25,26]. Taxonomically, these accessions could be classified as *Ae. tauschii* Coss. subspecies *strangulata* (Eig) Tzvel. Our data demonstrated that all these *strangulata* accessions, which were not glaucous, had the *Iw2Iw2* genotype (Figure 1). On the assumption that the ancestral *Ae. tauschii* had the *W2W2Iw2Iw2* genotype, this finding may suggest that the southeastern coastal Caspian populations of *Ae. tauschii* subspecies *strangulata* do not represent the direct descendants of the ancestral populations that gave rise to hexaploid common wheat.

**Conclusions**

Analysis of the *Iw2* locus may contribute to improve our understanding of the evolution of hexaploid wheat. Of the 210 *Ae. tauschii* accessions, only 20 glaucous accessions were found in the area that spans from Transcaucasia to the southern coastal region of the Caspian Sea. Of the 82 synthetic wheat lines that we examined, 15 were glaucous, and each of the 67 non-glaucous lines was produced by crossing Ldn with a non-glaucous *Ae. tauschii* accession. Of the 15 glaucous lines, 13 were produced by crossing Ldn with glaucous *Ae. tauschii* accessions. The remained
two accessions seemed to have the Iw2w2 genotype according to the genotyping analysis with the Iw2-linked markers. Therefore, allelic differences at the Iw2 locus on the short arm of chromosome 2D were the main contributors to the phenotypic difference between the glaucous and non-glaucous accessions of Ae. tauschii. Some molecular markers, such as S10812-1, closely linked to Iw2 were significantly associated with variation in glaucousness in Ae. tauschii. These results suggest that the D-genome donor of common wheat could have been any Ae. tauschii variant that carried the recessive iw2 allele.

Availability of supporting data
The data sets supporting the results of this article are included within the article and its supplementary files.

Additional files

Additional file 1: List of primers developed in this study.
Additional file 2: List of all 20 Ae. tauschii glaucous accessions and wheat synthetics with the glaucousness phenotype in the 82 synthetic hexaploid wheat lines.

Abbreviations
CAPS: Cleaved amplified polymorphic sequence; HRM: High resolution melting; Ldn: Langdon; SNP: Single nucleotide polymorphism; SSR: Simple sequence repeat.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
RN carried out mapping and genotyping experiments and drafted the manuscript. XMI contributed to the bioinformatics-related analysis. VM surveyed the natural variation in Ae. tauschii and revised the manuscript. ST conceived the study, acquired the funding, examined the variation in wheat synthetics, and drafted and revised the manuscript. All authors have read and approved the final manuscript.

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