DArTseq-based analysis of genomic relationships among species of tribe Triticeae

Edet, Offiong U.; Gorafi, Yasir S. A.; Nasuda, Shuhei; Tsujimoto, Hisashi

CITATION:
Edet, Offiong U. ...[et al]. DArTseq-based analysis of genomic relationships among species of tribe Triticeae. Scientific reports 2018, 8(1): 16397.

ISSUE DATE: 2018-11-06

URL: http://hdl.handle.net/2433/245792

RIGHT: This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.
DArTseq-based analysis of genomic relationships among species of tribe Triticeae

Offiong U. Edet1,2, Yasir S. A. Gorafi1,3, Shuhei Nasuda4 & Hisashi Tsujimoto1

Precise utilization of wild genetic resources to improve the resistance of their cultivated relatives to environmental growth limiting factors, such as salinity stress and diseases, requires a clear understanding of their genomic relationships. Although seriously criticized, analyzing these relationships in tribe Triticeae has largely been based on meiotic chromosome pairing in hybrids of wide crosses, a specialized and labourious strategy. In this study, DArTseq, an efficient genotyping-by-sequencing platform, was applied to analyze the genomes of 34 Triticeae species. We reconstructed the phylogenetic relationships among diploid and polyploid Aegilops and Triticum species, including hexaploid wheat. Tentatively, we have identified the diploid genomes that are likely to have been involved in the evolution of five polyploid species of Aegilops, which have remained unresolved for decades. Explanations which cast light on the progenitor of the A genomes and the complex genomic status of the B/G genomes of polyploid Triticum species in the Emmer and Timopheevi lineages of wheat have also been provided. This study has, therefore, demonstrated that DArTseq genotyping can be effectively applied to analyze the genomes of plants, especially where their genome sequence information are not available.

Triticeae is one of the most economically important tribes of the grass family, Poaceae, and includes such globally significant species as bread wheat (Triticum aestivum L.), barley (Hordeum vulgare L.), and rye (Secale cereale L.). Efforts to analyze the genomes of species constituting this tribe to uncover the evolutionary relationships among them have been on for decades1–5. These research efforts, beyond attempting to clarify the taxonomy of Triticeae, are largely propelled by the overwhelming role of wild species of this tribe as potential sources of essential alleles for the improvement of their cultivated relatives, especially bread wheat5–8.

Analysis of genome differences and phylogenetic relationships in Triticeae has mostly been conducted using cytogenetic approaches that rely on meiotic chromosome pairing in hybrids of wide crosses. However, chromosome pairing is affected by diverse factors, and the reliability of failed chromosome pairing as an indicator of genome dissimilarity has been questioned9–11. Molecular cytogenetic methods such as C-banding, fluorescence in situ hybridization and genomic in situ hybridization have also helped to generate useful information on genome differences and phylogenetic relationships in the Triticeae1–3,12,13. Other molecular approaches, including isozyme analysis, variations in low-molecular-weight glutenin subunit and DNA marker systems have provided some explanations on Aegilops-Triticum relationships, the origin and differentiation of Aegilops species, and intra- and inter-specific variations in the D and U genome clusters of Aegilops species14–17. Also, a combination of morphology, organelle and nuclear genes reportedly gave insights into the phylogenetic relationships among diploid taxa in Triticeae18. Nevertheless, diploid progenitors of Ae. crassa, Ae. vavilovii, Ae. juvenalis, Ae. columnaris and Ae. triariistata, and the exact progenitor of the B genome of hexaploid wheat and other polyploid Triticum species are still in dispute19–22. Also, there are opposing opinions regarding the donors of A genomes of polyploid species in the Emmer (AB group) and Timopheevi (AG group) lineages of wheat23–25. In the classification of Aegilops species, the justification for including Ae. speltoides in section Sitopsis is still under discussion26–32. To validate the results so far obtained and fill remaining gaps, more molecular data, especially those with a wide genomic coverage, are needed11.

1Arid Land Research Center, Tottori University, Tottori, 680-0001, Japan. 2United Graduate School of Agricultural Sciences, Tottori University, Tottori, 680-8553, Japan. 3Agricultural Research Corporation (ARC), P. O. Box 126, Wad Madani, Sudan. 4Laboratory of Plant Genetics, Graduate School of Agriculture, Kyoto University, Kyoto, 606-8502, Japan. Correspondence and requests for materials should be addressed to H.T. (email: tsujim@alrc.tottori-u.ac.jp)

Received: 2 July 2018
Accepted: 26 October 2018
Published online: 06 November 2018
DArTseq is one of the cheap and easy but efficient genotyping-by-sequencing platforms which allow genome-wide marker discovery through restriction enzyme-mediated genome complexity reduction and sequencing of the restriction fragments. In this study, we applied DArTseq genotyping to analyze the genomes of 34 species in the tribe Triticeae. DArTseq generates two types of data: SNP and SilicoDArT. The former is the nucleotide polymorphisms found in the tag sequences and the latter is the presence/absence variation (PAV) of the tag sequences. The choice of which data to use depends on the research objective. Using both types of data, we clarified the extent of genomic similarity among Aegilops species in different sections and clusters and the evolutionary relationships between diploid and polyploid species in Aegilops and Triticum species. We confirmed the already known genomic constitutions of some polyploid species of Aegilops and Triticum species, and provided evidence-based explanation for the origin of the unidentified (X) genomes in Ae. crassa, Ae. columnaris, Ae. vavilovii, Ae. triaristata, and Ae. juvenalis. The consistency of the outcomes of this study with previous reports and the flexibility of DArTseq genotyping make this marker system suitable for routine applications to analyze plant genomes.

Results
Diploid analyzers of polyploid Aegilops and Triticum species. To determine the putative progenitors of each of the polyploid species of Aegilops, SilicoDArT markers in the diploid genomes of all the Aegilops species were used as genome analyzers (Table 1). For each of the diploid species, species-specific markers were selected by filtering markers present in one species but absent in all the others. This made dominant SilicoDArT markers preferred in this analysis, as codominant SNP markers do not give information on PAVs. The progenitors of the polyploid species were estimated based on the proportions of diploid species’ SilicoDArT markers retained in the genomes of the polyploid species. Because the number of the species-specific markers is affected by genetic similarity among the diploid species, especially the Sitopsis species, the genomes of the polyploid species of Aegilops were first analyzed with all the markers in each diploid genome of Aegilops species (Fig. 1a) before being analyzed with diploid species-specific markers (Fig. 1b). This allowed us to determine the suitability of the species-specific markers in estimating the progenitors of the polyploid species. The use of species-specific markers as analyzers reduced the background noise produced by monomorphic markers among the diploid species (Fig. 1b).

Table 1. SilicoDArT markers of diploid analyzers of polyploid Aegilops species.

| Species         | Total No. of markers | Species-specific markers (%) |
|-----------------|----------------------|-----------------------------|
| Ae. mutica      | 12238                | 837 (6.84)                  |
| Ae. speltoides  | 9330                 | 699 (7.49)                  |
| Ae. longissima  | 18321                | 761 (4.15)                  |
| Ae. sharonensis | 18205                | 723 (3.97)                  |
| Ae. bicornis    | 16465                | 598 (3.63)                  |
| Ae. searsii     | 15402                | 1633 (10.60)                |
| Ae. tauschii    | 20288                | 7420 (36.57)                |
| Ae. caudata     | 19086                | 6514 (34.13)                |
| Ae. comosa      | 17377                | 3941 (22.68)                |
| Ae. uniaristata | 16719                | 4003 (23.94)                |
| Ae. umbellulata | 19523                | 6627 (33.94)                |

Figure 1. Estimation of the putative diploid progenitors of 12 polyploid Aegilops species based on the proportions of diploid species’ SilicoDArT markers retained in the genomes of the polyploid species. (a) Analysis based on all the markers of the diploid species. (b) Analysis based on the species-specific markers of the diploid species.
Having confirmed the adequacy of the species-specific markers of the analyzers, polyploid genomes of *Triticum* species were analyzed with only species-specific diploid analyzers (Table 2). Therefore, the conclusions made regarding the progenitors of the polyploid species (*Aegilops* and *Triticum*) are based on the species-specific markers of the analyzers. To select analyzers for the polyploid genomes of *Triticum* species, the genomes of 16 bread wheat-related diploid species were screened on the basis of the proportions of homoeology of their SilicoDArT markers to those of A, B and D genomes of bread wheat. The homoeology of the diploid genomes to each of the genomes of bread wheat was estimated based on the number of markers of the polyploid species assigned to each of the three genomes of bread wheat (Table 2). This estimation was possible because, in this study, we used DArTseq platform optimized for bread wheat. A diploid species with at least 10% homoeology to any of the three genomes of hexaploid wheat was selected as an analyzer for the corresponding genomes of each of the six polyploid *Triticum* species. With this criterion, a total of 13 analyzers were selected. Species-specific markers of the 13 selected diploid analyzers were then filtered for the analysis of the putative progenitors of the genomes of the polyploid species (Table 2).

**Table 2.** Selection of diploid analyzers of polyploid *Triticum* species with respect to homoeology of their SilicoDArT markers to those of A, B and D genomes of bread wheat. A species whose proportion of markers assigned to any of the three genomes of bread wheat is not less than 10% was selected (emboldened) for the analysis of the corresponding genomes of the polyploid species.

![Table 2](https://example.com/table2.png)

| Species            | A genome     | B genome     | D genome     |
|--------------------|--------------|--------------|--------------|
|                     | Shared | Specific | Total (%) | Shared | Specific | Total (%) | Shared | Specific | Total (%) |
| Reference genome (bread wheat) | 16901 | 4688 | 21,589 (100) | 13415 | 7227 | 20642 (100) | 30187 | 5167 | 35354 (100) |
| *T. ararbi*         | 6114 | 3672 | 9786 (45.3) | 1249 | 95  | 1344 (6.5)  | 2763 | 93  | 2856 (8.1)  |
| *T. boeoticum*      | 5492 | 753  | 6245 (28.9) | 1369 | 71  | 1436 (7.0)  | 2870 | 59  | 2929 (8.3)  |
| *Ae. mutica*        | 1824 | 1348 | 3172 (14.7) | 2331 | 1966 | 4297 (20.8) | 4690 | 2284 | 6974 (19.7) |
| *Ae. speltoides*    | 1139 | 1129 | 2268 (10.5) | 2507 | 4405 | 6912 (33.5) | 3335 | 1883 | 5218 (14.8) |
| *Ae. longissima*    | 2301 | 392  | 2693 (12.5) | 3513 | 755  | 4268 (20.7) | 6831 | 594 | 7425 (21.0) |
| *Ae. sharonensis*   | 2294 | 336  | 2630 (12.2) | 3487 | 769  | 4256 (20.6) | 6799 | 632 | 7431 (21.0) |
| *Ae. bicornis*      | 2018 | 379  | 2397 (11.1) | 3026 | 756  | 3782 (18.3) | 6258 | 638 | 6896 (19.5) |
| *Ae. secasi*        | 1915 | 156  | 2071 (9.6)  | 3610 | 495  | 4105 (15.0) | 5558 | 265 | 5823 (16.5) |
| *Ae. tauchii*       | 817  | 99   | 916 (4.2)   | 934  | 174  | 1108 (5.4)  | 8988 | 7568 | 16556 (46.8) |
| *Ae. caudata*       | 1849 | 1129 | 2969 (13.8) | 1955 | 1013 | 2968 (14.4) | 5321 | 1848 | 7169 (20.3) |
| *Ae. comosa*        | 1959 | 1022 | 2981 (13.8) | 2126 | 1102 | 3228 (15.6) | 5545 | 1904 | 7449 (21.1) |
| *Ae. uniaristata*   | 1805 | 672  | 2477 (11.5) | 2049 | 724  | 2773 (13.4) | 5155 | 1254 | 6409 (18.1) |
| *Ae. umbellulata*   | 2014 | 928  | 2942 (13.6) | 2055 | 1017 | 3072 (14.9) | 5174 | 1653 | 6827 (19.3) |
| *S. cereale*        | 699  | 23   | 722 (3.3)   | 959  | 64   | 1023 (5.0)  | 1840 | 30  | 1870 (5.3)  |
| *D. villum*         | 566  | 20   | 586 (2.7)   | 793  | 44   | 837 (4.1)   | 1538 | 16  | 1554 (4.4)  |
| *H. vulgare*        | 291  | 8    | 299 (1.4)   | 431  | 19   | 450 (2.2)   | 824  | 9   | 833 (2.4)   |

**Genomic differentiation and evolutionary relationships among polyploid and diploid species of *Aegilops*.** Before applying the genome analyzers to determine the progenitors of the polyploid species, we used a total of 28,264 polyploid species-specific SilicoDArT markers, ranging from 187 in *Ae. juvencalis* to 4,759 in *Ae. cylindrica* (Table 3), to confirm genomic difference among the 12 polyploid species of *Aegilops*. The polyploid species-specific markers were selected in the same manner as the diploid species-specific markers. The relatively low numbers of species-specific markers in the genomes of *Ae. crassa* and *Ae. juvencalis* is obviously because large proportions of their genomes (D and U genomes) are shared by the other species (Table 3). With the possibility of genomic adjustments during polyploidization and the assumption that the original progenitors of the polyploid species may be different from the accessions of the diploid species used in this study, only diploid analyzers with considerably higher proportions of monomorphism with the polyploid species were taken as the putative progenitors of the polyploid species. Our analysis confirmed the putative diploid progenitors of *Ae. ventricosa* (D*Vv*), *Ae. cylindrica* (C*D*), *Ae. kotschyi* (S*UgM*), *Ae. biuncialis* (U*Me*), *Ae. triuncialis* (UC), *Ae. ovata* (U*M0*), and *Ae. variabilis* (S*U*U*P*) (Fig. 1). Noteworthy is that the proportions of the markers of the three Sitopsis species (Ae. bicornis, Ae. longissima, and Ae. sharonensis) retained in the genomes of the two polyploid species with S-related genomes (*Ae. secasi* and *Ae. variabilis*) were not reasonably different. This made it difficult to decide which of the Sitopsis species donated the S-related genomes to *Ae. kotschyi* and *Ae. variabilis*, although *Ae. bicornis* and *Ae. longissima*, respectively, seem to be the most likely candidates. This observation confirms the likelihood of a common ancestry of the Sitopsis species. Therefore, the original progenitor of the S-related genomes of the polyploid species may have been an ancient relative of the Sitopsis species, which is probably extinct. Although we adopted the polyploid species-specific markers (Table 3) to differentiate *Ae. kotschyi* (S*UgM*) from *Ae. variabilis* (S*U*U*P*) and *Ae. biuncialis* (U*Me*) from *Ae. ovata* (U*M0*), these pairs of species have identical genomic constitutions (same progenitors; Fig. 1), and therefore may be considered to be variants/subspecies of the same species in each case.
Table 3. Species-specific SilicoDArT markers of 12 polyploid species of *Aegilops*.

| Species            | Ploidy | Reported genomic formula | Proposed genomic formula |
|--------------------|--------|--------------------------|--------------------------|
| *Ae. speltoides*   | 4x, 6x | D^4X^4, D^4D^2X^4        | DT^4, D^4T^4             |
| *Ae. vavilovii*    | 6x     | D^4X^4S^4                | DT^4S^4                  |
| *Ae. juvenalis*    | 6x     | XD^4U                    | UD^4T                    |
| *Ae. triaristata*  | 4x, 6x | U^4X^4, U^4X^4N^4         | UN, UN^T                 |
| *Ae. columnaris*   | 4x     | U^4X                        | UT^4                     |
| **Total**          | —      | —                         | 28264                    |

Table 4. Proposed modifications in the genomic representations of five polyploid *Aegilops* species. T^5: joint representation of *Ae. speltoides* and *Ae. mutica*.

| Species            | Ploidy | Genomic representations of the five polyploid *Aegilops* species. |
|--------------------|--------|------------------------------------------------------------------|
| *Ae. speltoides*   | 6x     | D^4D^2X^4S^4                                                      |
| *Ae. vavilovii*    | 6x     | D^4X^4S^4                                                        |
| *Ae. juvenalis*    | 6x     | D^4X^4                                                           |
| *Ae. triaristata*  | 4x, 6x | U^4X^4, U^4X^4N^4                                                 |
| *Ae. columnaris*   | 4x     | U^4X                                                           |

Cluster analysis of diploid and polyploid *Aegilops* species. A phylogenetic tree (Fig. 2a) constructed with 15,512 frequently called SNP markers separated the diploid *Aegilops* species into the already reported sections, except that *Ae. speltoides* did not cluster with other species in the section Sitopsis, which has been reported by other researchers. Aegilops *umbellulata* (section Aegilops) seemed more distant from the others, whereas *Ae. speltoides* (section Sitopsis) appeared closer to *Ae. mutica* (section Amblyopyrum), and relatively distant from other species of section Sitopsis. Among Sitopsis species, *Ae. longissima* and *Ae. sharoneensis* appeared genomically more proximal to each other than to others. The polyploid species of *Aegilops* formed two clusters clearly based on the putative common diploid progenitors, *Ae. tauschii* (D cluster) and *Ae. umbellulata* (U cluster) (Fig. 2b). *Aegilops juvenalis*, bearing both D and U genomes, clustered closely with *Ae. crassa* and *Ae. vavilovii* in the D cluster, indicating a possible evolutionary link between its (*Ae. juvenalis*) genome and the two species in the D cluster. This again suggests the likelihood of the presence of a diploid genome, perhaps T^5, common to *Ae. crassa*, *Ae. vavilovii* (Fig. 1).

Genomic and evolutionary relationships in the *Aegilops-Triticum* species. We used species-specific SilicoDArT markers of 13 bread wheat-related diploid species (Table 2) to determine the elementary donors of the A, B and D genomes in six polyploid *Triticum* species. As described for the estimation of the progenitors of the
polyploid species of *Aegilops*, the proportions of species-specific markers of the diploid species shared with the genomes of the polyploid species enabled the determination of the progenitors of the genomes of the polyploid *Triticum* species (Figs 3–6).

The genome of *T. urartu* was the closest to the A genomes of all the polyploid species analyzed (Fig. 3), suggesting that *T. urartu* is the most likely donor of the A genome in each of them. The considerable similarity between the A genome of each of the polyploids and *T. boeoticum* – another A genome species – suggests a common ancestry of *T. boeoticum* and *T. urartu*. Similarly, *Ae. searsii* seems to be the most closely related to the B/G genomes of the polyploid species (Fig. 4). However, the proportion of *Ae. speltoides* markers assigned to the reference B genome is higher than those of every other diploid species analyzed (Table 2). This strongly suggests

---

**Figure 2.** Reconstruction of the evolutionary relationships among *Aegilops* species on the basis of 15,512 SNP markers. (a) 11 diploid species. (b) 12 polyploid species. The sections of the diploids and two main clusters of the polyploids are labelled in brown\cite{11,12}.

**Figure 3.** Determination of the donor of the A genomes of six polyploid *Triticum* species using 11 diploid species-specific SilicoDArT markers assigned to the A genome of bread wheat.

**Figure 4.** Determination of the donor of the B/G genomes of six polyploid *Triticum* species using 10 diploid species-specific SilicoDArT markers assigned to the B genome of bread wheat.
an evolutionary link between the genome of *Ae. speltoides* and the B/G genomes of the polyploids. This link is further supported by an almost equal similarity of the genomes of the two diploid species to the G genome of *T. araraticum* (Fig. 4). Using the species-specific markers of *Ae. speltoides* and *Ae. searsii* as analyzers, we found that chromosome 4S of each of the diploid species are almost equally similar to chromosome 4B/G of each of the polyploids (Fig. 5). But chromosomes 2S, 3S and 7S of *Ae. speltoides* appeared to be more similar to the corresponding chromosomes of *T. araraticum* than those of *Ae. searsii* are. These observations give the impression that the B/G genomes of polyploid *Triticum* species are likely to be recombinant genomes with varying contributions from *Ae. speltoides* and *Ae. searsii*. Analysis of the D genomes of the three hexaploid species unambiguously traced them to *Ae. tauschii* as the sole donor (Fig. 6).

A further analysis using 66, 434 SNP markers consistently called in the six polyploid genomes (see online Supplementary Table S2 for information of the markers) indicated 72% similarity (monomorphism) across their A genomes, B/G genomes and the combined AB/AG genomes. However, higher similarity was observed among the AB genomes: hexaploids, 94%; tetraploids, 90%; hexaploid and tetraploid genomes combined, 84%. The slight differences in the proportions of monomorphic markers in the different groups of the AB genomes suggest that the AB genomes of the hexaploid species originated from the same tetraploid species, whereas those of the tetraploid species may have evolved from different accessions of the elementary A and B genome progenitors (*T. urartu* and *Ae. speltoides/Ae. searsii*, respectively). The lower similarity (84% as compared to 94%) across the hexaploid and tetraploid AB genomes may reflect further modification of AB genomes in hexaploid species resulting from their interaction with the D genome.

**Discussion**

The clustering patterns of *Aegilops* species were largely consistent with the established classifications. Diploid species separated on the basis of their known sections in the genus; polyploid species were delineated following the presence of common diploid progenitor genomes (D and U genomes) among them. However, as reported previously, *Ae. speltoides* appeared distant from other species in the section Sitopsis; hence, its inclusion in the section needs to be reconsidered. Markers specific to each of the 12 polyploid species clearly showed considerable polymorphisms among these genomes, including the genomes of species which arose from the same diploid progenitors. This suggests that genetic modifications, such as chromosomal alterations, may have occurred during independent evolutionary events of those species with identical progenitors. Therefore, without these specific markers, it would be difficult to genomically differentiate *Ae. kotschyi* (S’U’) from *Ae. variabilis* (S’U’P) and *Ae. biuncialis* (U’M’) from *Ae. ovata* (U’M’) because, from the stand point of our result (Fig. 1) and previous studies, the species in each pair evolved from the same progenitors. Although each of the species in these two sets are recognized as independent, on the basis of differences in cytoplasmic progenitors and/or nuclear genome variation, this classification does not seem to be clearly justified. Therefore, in our opinion, each pair should be regarded as variants/subspecies of the same species.

The reported unknown diploid genomes, initially represented as modified M genome and later changed to X, in the genomes of *Ae. triaristata*, *Ae. crassa*, *Ae. juvenalis*, *Ae. vavilovii*, and *Ae. columnaris*, is traceable to *Ae. mutica* or *Ae. speltoides*. The small proportions (<10%) of *Ae. comosa*-specific markers shared with the five
polyploid species (Fig. 1b) is insufficient to infer the existence of remnants of *Ae. comosa* genome in the polyploid genomes. Assuming *Ae. comosa* was originally involved in the evolution of the polyploids, species-specific elements from other progenitors may have spread and eventually masked *Ae. comosa*-specific elements. Our data suggest that ancient or ancestral forms of *Ae. speltoides* or *Ae. mütica*, which are probably extinct, donated the unidentified genomes to the five polyploid species. From our analysis, it appears that all the polyploid species originally had a genome of such an ancient species (Fig. 1). This observation agrees with the hypothesis that *Ae. mütica* (*syn. Amblyoppyrum müticum*) and *Ae. speltoides*, both allogamous species with ancestral traits, diverged earlier than other *Aegilops* species and may therefore be the ancestors of the other *Aegilops* species. Therefore, each diploid *Aegilops* species may have retained a substantial portion of the common ancestral genome (*Ae. speltoides/Ae. mütica* or their ancestor). The difference in the representation of the common progenitor in each of the polyploid species can result from the peculiar evolutionary event(s) of each species. Polyploids that arose from the hybridization of the common diploid ancestor with other diploid species should have larger portions of the genome of the common ancestor than those that did not directly evolve from the common ancestor.

We validated the putative diploid progenitors of the A and D genomes of polyploid *Triticum* species to be *T. urartu* and *Ae. tauschii*, respectively. We have also provided information that may help to explain the complex nature of the B/G genomes. The genomes of both *Ae. speltoides* and *Ae. searsii* are similar to the B/G genomes, especially to that of *T. araraticum* (Fig. 4), a relatively less advanced tetraploid genome; thus, the B/G genomes of polyploid *Triticum* species may have evolved from an ancestral genome that later differentiated into those of *Ae. speltoides* and *Ae. searsii*. Alternatively, the B/G genome may have arisen from the hybridization of *Ae. speltoides* and *Ae. searsii* before the emergence of the AB/AG genome at different times. The above considerations support earlier postulations that the B genome is the most modified of the three genomes of hexaploid wheat, whereas the A and D genomes are substantially similar to those of *T. urartu* and *Ae. tauschii*, respectively. The previously suggested origin of the A genome of *T. araraticum* from *T. boeoticum* is probably invalid (Fig. 3). Our results agree with the hypothesis that both Emmer and Timopheevi lineages of polyploid wheat have the same sources of elementary A and B/G genomes. However, a common ancestry of the A-genome species cannot be ruled out and the A genomes of polyploid *Triticum* species may have evolved from a common ancestor of *T. urartu* and *T. boeoticum* before the differentiation of the two species. Although no karyotypic differences have been detected between these diploid A-genome species, low fertility of interspecific F1 hybrid plants of these two species has been reported. The latter study, consistent with our result, confirms that the two species are genomically different. As previously documented, our study suggests that the A and G genomes in *T. araraticum* are less modified than the A and B genomes in the Emmer lineage.

This study has demonstrated that DArTseq genotyping can be applied to conduct a large scale analysis of plant genomes, mostly because it allocates markers to individual chromosomes, which can be easily extracted and analyzed. This genomic sequence-based platform ensures a wide genomic coverage and is not subject to criticism associated with the factors that affect meiotic chromosome pairing in hybrids of distant crosses, which forms the main anchor of cytogenetic systems of genome analysis. Also, the number of informative markers generated by DArTseq outstrips what is possible with conventional DNA marker procedures, making it more robust and reliable. Genotyping of all the available accessions of species in tribe Triticeae using this platform would clarify the genomic relationships between the cultivated and wild species. This information would make the use of the available gene pools for breeding much more precise and would also help to clarify Triticeae taxonomy. As polyploidy and interspecific hybridization are key events in the evolution of higher plants, this genome analysis approach would be useful in other groups of plant species, especially polyploids with unclear phylogeny.

**Methods**

**Plant materials.** All 23 *Aegilops* species, eight *Triticum* species, and three distant relatives of wheat were analyzed. Except bread wheat, represented by two cultivars ‘Chinese Spring’ (CS) and ‘Norin 61’ (N61), each species was represented by one accession (Table 5). Seedlings were raised in Greenhouses of the Arid Land Research Center, Tottori University and Laboratory of Plant Genetics, Kyoto University, Japan. Depending on growth rate and plant size, fresh leaves were harvested from each 2–4-week-old seedlings and genomic DNA samples were isolated and purified using the cetyl trimethyl ammonium bromide method. Quality check, quantification
Table 5. List of plant materials. Names and genome symbols of relatives of wheat can be found in Hagras et al.56. The high marker number generated by this system gives it an edge over previous molecular marker procedures applied for genomic analysis of Triticeae species 14–17. It, therefore, serves as a cheap alternative to genome analysis by whole genome sequencing, where the sequence information of genomes intended to be analyzed are not available. Two types of data are generated by DArTseq: SNP and SilicoDArT. SNP markers are nucleotide polymorphisms present in the restriction fragments, while SilicoDArT markers represent PAV of the restriction fragments. Therefore, codominant SNP markers are scored “0” (reference allele homozygote), “1” (SNP allele homozygote) and “2” (heterozygote: presence of both reference and SNP alleles), while dominant SilicoDArT markers are scored in a binary fashion, with “1” representing presence of the restriction fragment with the marker sequence and “0” designating its absence.

| ID     | Species       | Subspecies/cultivar | Ploidy | Genome | Source     |
|--------|---------------|---------------------|--------|--------|------------|
| KU–12007 | Ae. mutica Boiss. | —                   | 2x     | T      | NBRP       |
| KU–2–5 | Ae. speltoides Tausch | typica             | 2x     | S      | NBRP       |
| KU–4–1 | Ae. longissima Schweinl. et Muschl | typica           | 2x     | S'     | NBRP       |
| KU–5–3 | Ae. sharonensis Eig | typica            | 2x     | S'^4   | NBRP       |
| KU–4–6 | Ae. searsii Feldman et Kile ex Hammer | —             | 2x     | S'     | NBRP       |
| KU–3–1 | Ae. bicornis (Forssk.) Jaub. et Sp. | typica        | 2x     | S^5    | NBRP       |
| KU–2159 | Ae. tauschii Coss. | typica            | 2x     | D      | NBRP       |
| KU–5860 | Ae. caudata L. | polyaethera       | 2x     | C      | NBRP       |
| KU–17–1 | Ae. comosa Sibth. et Sm. | comosa       | 2x     | M      | NBRP       |
| KU–19–3 | Ae. uniaristata Vis. | typica      | 2x     | N      | NBRP       |
| KU–8–2 | Ae. umbellulata Zhuk. | typica  | 2x     | U      | NBRP       |
| KU–7–1 | Ae. cylindrica Host. | typica    | 4x     | CD     | NBRP       |
| KU–22–1 | Ae. ventricosa Tausch | comosa   | 4x     | D'N    | NBRP       |
| KU–9–1 | Ae. ovata L. | vulgaris       | 4x     | U'M^4  | NBRP       |
| KU–11–1 | Ae. columnaris Zhuk. | typica  | 4x     | U'M^4  | NBRP       |
| KU–13–6 | Ae. kotschyi Boiss. | leptostachya | 4x     | S'U^a  | NBRP       |
| KU–15–1 | Ae. triuncialis L. | typica       | 4x     | C'T^a  | NBRP       |
| KU–13–1 | Ae. varsubii Eig | intermedia    | 4x     | S'L^a  | NBRP       |
| KU–12–1 | Ae. biuncialis Vis. | typica    | 4x     | U'M^3  | NBRP       |
| KU–10–1 | Ae. triaristata Willd. | vulgaris  | 6x     | U'M'N^4| NBRP       |
| KU–21–1 | Ae. cerasa Boiss. | typica       | 6x     | D'M^4, D'D'M^4 | NBRP |
| KU–21–7 | Ae. varslavii (Zhuk.) Chennav. | palaestina | 6x     | D'M^8 | NBRP       |
| KU–23–3 | Ae. juvenalis (Thell.) Eig | typica  | 6x     | D'M'M^4 | NBRP |
| KU–199–11 | T. urartu Thum. Ex. Gandil. | nigrum  | 2x     | A^a    | NBRP       |
| KT001–001 | T. boeoticum Boiss. | boeoticum  | 2x     | A^a    | NBRP       |
| KT099–17 | T. darcum Desf. | Langdon KU  | 4x     | AB     | NBRP       |
| KU–491 | T. dicoccum (Schrank) Schuebl. | —        | 4x     | AB     | NBRP       |
| KU–108–1 | T. dicoccoides (Koern. Ex Aschers & Graebn.) Schweif | kotschyssum | 4x     | AB     | NBRP       |
| KU–196–1 | T. aratiticum Jakubz. | tumaniani | 4x     | AG     | NBRP       |
| KT020–003 | T. aestivum L. | Chinese Spring | 6x     | AB     | NBRP       |
| KU–260 | T. aestivum L. | Norin 61   | 6x     | AB     | NBRP       |
| KT018–002 | T. macha Dekapr. & Menabde | palaeomereticum | 6x     | AB     | NBRP       |
| TACBOW0071 | S. cereale | Pektas  | 2x     | R      | NBRP       |
| TACBOW0119 | D. villosum | —      | 2x     | V      | NBRP       |
| TACBOW0116 | H. vulgare | Betzes  | 2x     | H      | NBRP       |

and concentration adjustment for sequencing and genotyping were accomplished with NanoDrop2000C Spectrophotometer (ThermoScientific). The concentration of each sample was adjusted to 50 ng/μL.

Genotyping-by-sequencing and data analysis. Purified DNA samples (1 μg for each sample) were sent to Diversity Arrays Technology Pty Ltd (http://www.diversityarrays.com/) for sequencing and marker identification. Sequences of the genomic representations were aligned to the wheat_ChineseSpring10 reference genome and wheat_ConsensusMap_version_4. This is because our analysis is based on DArTseq platform optimized for hexaploid wheat. DArTseq is a genotyping-by-sequencing system which utilizes Next-Generation-Sequencing platforms (HiSeq. 2500 in our case) to sequence the most informative representations of genomic DNA samples to aid marker discovery. In comparison to the array version of DArT, DArTseq results in higher marker densities58. The high marker number generated by this system gives it an edge over previous molecular marker procedures applied for genomic analysis of Triticeae species14–17. It, therefore, serves as a cheap alternative to genome analysis by whole genome sequencing, where the sequence information of genomes intended to be analyzed are not available. Two types of data are generated by DArTseq: SNP and SilicoDArT. SNP markers are nucleotide polymorphisms present in the restriction fragments, while SilicoDArT markers represent PAV of the restriction fragments. Therefore, codominant SNP markers are scored “0” (reference allele homozygote), “1” (SNP allele homozygote) and “2” (heterozygote: presence of both reference and SNP alleles), while dominant SilicoDArT markers are scored in a binary fashion, with “1” representing presence of the restriction fragment with the marker sequence and “0” designating its absence.
Frequently called SNP markers (>0.9 call rate) were used for phylogenetic tree constructions and differentiation of the genomes of polyploid species of *Aegilops*, whereas SilicoDArT markers (>0.7 call rate) were used for the determination of putative progenitors of the polyploid *Aegilops* and *Triticum* species. This reduction in call rate was made to accommodate more markers, ensure wider genomic coverage and reduce bias. To estimate the phylogenetic relationships among the 11 diploid and 12 polyploid *Aegilops* species, the raw genotypic data of the two sets (diploid and polyploid) were subjected to cluster analysis. Pearson’s correlation coefficient (r) was used as similarity index, and the genetic distances among the species were estimated by transforming the r values to distance values, using $d = 100(1 – r)$ (http://genomes.urv.cat/UPGMA/) [57]. Species-specific SilicoDArT markers of the polyploid species of *Aegilops* were used to differentiate their genomes, while species-specific SilicoDArT markers of diploid species of *Aegilops* were used to estimate the diploid-polyploid evolutionary relationships among all the *Aegilops* species. Diploid *Triticum* and *Aegilops* species whose total SilicoDArT markers showed at least 10% homology to the total SilicoDArT markers in any of the three genomes of hexaploid wheat were selected as analyzers to determine the putative progenitors of the corresponding genomes of each polyploid *Triticum* species. Species-specific SilicoDArT markers of these selected diploid species were used as analyzers to determine the putative progenitors of each polyploid *Triticum* species. In determining the progenitors of all the polyploid species (*Aegilops* and *Triticum*), the proportions of the species-specific markers of the diploid analyzers retained in the genomes of the polyploid species were used as a basis to draw conclusions on genomic proximity and evolutionary relationships among the species. Species-specific markers of *Ae. speltoides* and *Ae. searsii* were further used to examine the relationship between the seven B/G-genome chromosomes of each of the polyploid *Triticum* species and those of the two diploid species. The two diploid species were chosen based on the close proximity of their genomes to the B/G genomes of the polyploid species.

### Availability of Materials and Data

The data on which are conclusions are based are included within the article and supplementary files and plant materials can be sourced from KOMUGI database maintained by the National BioResource Project – Wheat, Japan (https://shigen.nig.ac.jp/wheat/komugi/).

### References

1. Badaeva, E. D. Evaluation of phylogenetic relationships between five polyploid *Aegilops* L. species of the U-genome cluster by means of chromosomal analysis. *Genetika* 38, 799–811 (2002).

2. Badaeva, E. D. et al. Genome differentiation in *Aegilops*. 4. Evolution of the U-genome cluster. *Plant Syst Evol* 246, 45–76, https://doi.org/10.1007/s00686-003-0072-4 (2004).

3. Badaeva, E. D. et al. Genetic classification of *Aegilops columnaris* Zhuk. (2n = 4x = 28, U(c)(U(c))(X)(c)(X)(c)) chromosomes based on FISH analysis and substitution patterns in common wheat × *Ae. columnaris* introgressive lines. *Genome* 61, 131–143, https://doi.org/10.1139/gen-2017-0186 (2018).

4. Friebe, B., Badaeva, E. D., Kammer, K. & Gill, B. S. Standard karyotypes of *Aegilops uniaristata*. *Genome* 53, 176–183, https://doi.org/10.1007/s10045-009-9318-7 (2010).

5. Molnár-Láng, M., Ceoñón, C. & Dolezel, J. Alien introgression in wheat: cytogenetics, molecular biology, and genomics. 385 (Springer, Switzerland, 2015).

6. Zhang, R. Q. et al. Development of VNTR chromosome alterations and physical mapping of molecular markers specific to *Dasypyrum villosum*. *Mol Breeding* 37, https://doi.org/10.1007/s11032-017-0671-3 (2017).

7. Riar, A. A., Kaur, S., Dhaliwal, H. S., Singh, K. & Chhuneja, P. Introggression of a leaf rust resistance gene from *Aegilops caudata* to bread wheat. *J Genet* 91, 155–161 (2012).

8. Ali, N. et al. Introggression of chromosome segments from multiple alien wheat species in bread wheat lines with wheat streak mosaic virus resistance. *Hereditas* (Edinb) 117, 114–123, https://doi.org/10.1080/00170496.2016.136 (2016).

9. Baum, B., Estes, J. & Gupta, P. Assessment of the genomic system of classification in *Triticum*. *Am J Bot* 74, 1388–1395 (1987).

10. Farooq, S., Jibaj, N. & Shah, T. M. Promotion of homoelogous chromosome-pairing in hybrids of *Triticum-aestivum×Aegilops variabilis*. *Genome* 33, 825–828, https://doi.org/10.1139/g90-124 (1990).

11. Seberg, O. & Petersen, G. A critical review of concepts and methods used in classical genome analysis. *Bot Rev* 64, 372–417 (1998).

12. Molnar, I. et al. Flow sorting of C-genome chromosomes from wild relatives of wheat *Aegilops marshallii*, *Ae. triuncialis*, and *Ae. cylindrica*, and their molecular organization. *Ann Bot* 116, 189–200, https://doi.org/10.1093/aob/mcv073 (2015).

13. Coriton, O. et al. Assignment of *Aegilops variabilis* Eish chromosomes and translocations carrying resistance to nematodes in wheat. *Genome* 52, 338–346, https://doi.org/10.1007/s10049-009-0119 (2009).

14. Wang, S. et al. Phylogenetic analysis of C, M, N, and U genomes and their relationships with *Triticum* and other related genomes as revealed by LMW-GS genes at Glu-3 loci. *Genome* 54, 273–284, https://doi.org/10.1007/s10049-010-1119 (2011).

15. Goryunova, S. V., Chikuda, N. N. & Kochieva, E. Z. RAPD analysis of the intraspecific and interspecific variation and phylogenetic relationships of *Aegilops* L. species with the U genome. *Genetika* 46, 945–959 (2010).

16. Goryunova, S. V., Kochieva, E. Z., Chikuda, N. N. & Pukhal’skii, V. A. Phylogenetic relationships and intraspecific variation of D-genome *Aegilops* L. as revealed by RAPD analysis. *Genetika* 40, 642–651 (2004).

17. Mizuno, N., Yamazaki, M., Matsuoka, Y., Kawahara, T. & Takumi, S. Population structure of wild wheat D-genome progenitor *Aegilops tauschii* Coss.: implications for intraspecific lineage diversification and evolution of common wheat. *Mol Ecol* 19, 999–1013, https://doi.org/10.1111/j.1365-2940.2010.04537.x (2010).

18. Seberg, O. & Petersen, G. Phylogeny of *Triticum* (Poaceae) based on three organelle genes, two single-copies nuclear genes, and morphology. *Aliso: A J Syst Evol Bot* 23, 362–371, https://doi.org/10.5642/aliso.20072301.29 (2007).

19. Dvorak, J. Genome analysis in the *Triticum-Aegilops* alliance. In: *Proc 9th Intl Wheat Genet Symp*. (ed A. E. Slinkard) 8–11 (University Extension Press: University of Saskatchewan, 1998).

20. Resta, P., Zhang, H. B., Dubcovsky, J. & Dvorak, J. The origins of the genomes of *Triticum bicincture*, *T. ovatum*, *T. neglectum*, *T. columnare*, and *T. rectum* (Poaceae) based on variation in repeated nucleotide sequences. *American Journal of Botany* 83, 1556–1565, https://doi.org/10.2307/2445829 (1996).

21. Badaeva, E. D., Friebe, B., Zoschuk, S. A., Zelenin, A. V. & Gill, B. S. Molecular cytogenetic analysis of tetraploid and hexaploid *Aegilops* crassa. *Chromosoma* 66, 629–637, https://doi.org/10.1007/A10095273291 (1998).

22. Feldman, M. & Levy, A. Origin and evolution of wheat and related *Triticum* species. In: *Alien introgression in wheat: cytogenetics, molecular biology, and genomics* (eds M. Molnár-Láng, C. Ceoñón, & J. Dolezel) 21–76 (Springer, 2015).

23. Dorofeev, V., Filatenco, A. & Migushova, E. Wheat. In: *Flora of cultivated plants* Vol. 1 (eds V.F. Dorofeev & O.N. Korovin) (1979).

24. Migushova, E. F. & Komarev, A. V. Genetic heterogeneity of wild *Triticum dicoccoides* from Iraq. *J. Agricul. Sc.* 9, 18–19 (1975).
25. Golovnina, K. A., Kondratenko, E. Y., Blinov, A. G. & Goncharov, N. P. Phylogeny of the A genomes of wild and cultivated wheat species. Russ J Genet 45, 1360–1367, https://doi.org/10.1134/S1022795409110106 (2009).
26. Gornicki, P. et al. The chloroplastic view of the evolution of polyploid wheat. New Phytol 204, 704–714, https://doi.org/10.1111/nph.12931 (2014).
27. Chantret, N. et al. Molecular basis of evolutionary events that shaped the hardiness locus in diploid and polyploid wheat species (Triticum and Aegilops). Plant Cell 17, 1033–1045 (2005).
28. Kilian, B. et al. Independent wheat B and G genome origins in outcrossing Aegilops progenitor haplotypes. Mol Biol Evol 24, 217–227, https://doi.org/10.1093/molbev/msl151 (2007).
29. Giorgi, D., D'Ovidio, R., Tanzarella, O. A., Ceoloni, C. & Poredu, E. Isolation and characterization of S genome specific sequences from Aegilops sect. sitopsis species. Genome 46, 478–489, https://doi.org/10.1177/001667350204600402 (2003).
30. van Slageren, M. Wild wheats: a monograph of Aegilops L. and Amblyopyrum (Jaub.&Spach) Eig., 512 (Wageningen Agric. Univ. and ICARDA, 1994).
31. Kilian, B. et al. Aegilops. In: Wild Crop Relatives: Genomic and Breeding Resources - Cereals (ed C Kole) 1–76 (Springer, 2011).
32. Goryunova, S. V., Chikida, N. N. & Kochieva, E. Z. Molecular analysis of the phylogenetic relationships among the diploid Aegilops species of the section Sitopsis. Russ J Genet 44, 115–118, https://doi.org/10.1134/S1022795408010146 (2008).
33. Andrews, K. R., Good, J. M., Miller, M. R., Luikart, G. & Hohenlohe, P. A. Harnessing the power of RADseq for ecological and evolutionary genomics. Nat Rev Genet 17, 81–92, https://doi.org/10.1038/nrg.2015.28 (2016).
34. Davey, J. W. et al. Genome-wide marker discovery and genotyping using next-generation sequencing. Nat Rev Genet 12, 499–510, https://doi.org/10.1038/nrg3102 (2011).
35. Melville, J. et al. Identifying hybridization and admixture using SNPs: application of the DArTseq platform in phylogeographic research on vertebrates. Roy Soc Open Sci 4, https://doi.org/10.1098/roso.160106 (2017).
36. Feldman, M. & Levy, A. A. Genome evolution due to allopolyploidization in wheat. Genetics 192, 763–774, https://doi.org/10.1534/genetics.114.164316 (2012).
37. Cabi, E., Dogan, M., Ozler, H., Akaydin, G. & Karagoz, A. Taxonomy, morphology and palynology of Aegilops variabilis (Zhek.) Chernov (Poaceae: Triticaceae). AIP J Agr Res 5, 2841–2849 (2010).
38. Dvorak, J. & Zhang, H. B. Reconstruction of the phylogeny of the Genus Triticum from variation in repeated nucleotide sequences. Theor Appl Genet 84, 419–429, https://doi.org/10.1007/BF00229502 (1992).
39. Golovnina, K. A. et al. Molecular phylogeny of the genus Triticum L. Plant Syst Evol 264, 195–216, https://doi.org/10.1007/s00606-006-0478-x (2007).
40. Goryunova, S. V., Chikida, N. N. & Kochieva, E. Z. AFLP, RAPD, and ISSR analysis of intraspecific polymorphism and interspecific differences of allotetraploid species Aegilops kotschyi Boiss. and Aegilops variabilis Eig. Russ J Genet 53, 568–575, https://doi.org/10.1134/S1022795417050040 (2017).
41. Ananthawat-Jonsson, K. Molecular cytogenetics of Leymus: Mapping the Ns genome-specific repetitive sequences. J Syst Evol 52, 716–721, https://doi.org/10.1111/jse.12106 (2014).
42. Ling, H. Q. et al.Draft genome of the wheat A-genome progenitor Triticum urartu. Nature 496, 87–90, https://doi.org/10.1038/ nature12020 (2013).
43. Daud, H. M. & Gustafson, J. P. Molecular evidence for Aegilops tauschii. Genome 39, 543–548, https://doi.org/10.1139/g96-069 (1996).
44. Lelley, T., Stachel, M., Grausgruber, H. & Vollmann, J. Analysis of relationships between Aegilops tauschii and the D genome of wheat utilizing microsatellites. Genome 43, 661–666, https://doi.org/10.1139/gen-43–4-661 (2000).
45. Luo, M. C. et al. Genome sequence of the progenitor of the wheat D genome Aegilops tauschii. Nature 551, 498, https://doi.org/10.1038/nature24486 (2017).
46. Odinotsno, T. I. et al. Analysis of Triticum boeoticum and Triticum urartu seed defensins: To the problem of the origin of polyploid wheat genomes. Biochimie 90, 939–946, https://doi.org/10.1016/j.biochi.2008.02.023 (2008).
47. Miyashita, N. T., Mori, N. & Tsumewaki, K. Molecular variation in chloroplast DNA regions in ancestral species of wheat. Genetics 137, 883–889 (1994).
48. Tsumewaki, K. et al. Use of RFLP analysis for wheat germplasm evaluation. In: Biodiversity and wheat improvement (ed A B Damania) 17–31 (John Wiley & Sons, 1993).
49. Giorgi, B. & Bozomi, A. Karyotype Analysis in Triticum IV: Analysis of (Ae. speltoides x T. boeoticum) amphiploid and a hypothesis on the evolution of tetraploid wheats. Caryologia 22, 289–306 (1969).
50. Fricano, A. et al. Crossability of Triticum urartu and Triticum monococcum wheats, homoeologous recombination, and description of a panel of interspecific introgression lines. G3-Genes Genom Genet 4, 1931–1941, https://doi.org/10.1534/g3.114.013623 (2014).
51. Kihara, H. Genomanalyse bei Triticum und Aegilops. Cytologia 1, 263–284 (1930).
52. Dewey, D. R. The genomic system of classification as a guide in intergeneric hybridization with the perennial Triticeae. In: Gene manipulation in plant improvement (ed J Gustafson) 209–279 (Plenum Publishing Corporation, 1984).
53. Love, A. Conspectus of the Triticeae. Feddes Repert 95, 423–521 (1984).
54. Moore, G. Early stages of meiosis in wheat and the role of Ph1. In: Genetics and genomes of wheats (eds G J Muehlihauber & C Feuillet) 237–252 (Springer, 2009).
55. Alik, K., Gerard, P. R., Schwarzer, T. & Heslop-Harrison, J. S. Polyploidy and interspecific hybridization: partners for adaptation, speciation and evolution in plants. Ann Bot 120, 184–194 (2017).
56. Kilian, A. et al. Diversity arrays technology: a generic genome profiling technology on open platforms. Methods Mol Biol 888, 67–89, https://doi.org/10.1007/978-1-61779-870-2_5 (2012).
57. García-Valverde, S., Palau, J. & Romeu, A. Horizontal gene transfer in glycosyl hydrolases inferred from codon usage in Escherichia coli and Bacillus subtilis. Mol Biol Evol 16, 1125–1134, https://doi.org/10.1093/oxfordjournals.molbev.a026203 (1999).
58. Bernhardt, N. Taxonomic treatments of Triticaceae and wheat genus Triticum. In: Alien introgression in wheat: cytogenetics, molecular biology, and genomics (ed M Molnár-Láng, Ceoloni, C. Doležel, J.) 1–20 (Springer, Switzerland, 2015).
59. Zhang, P. et al. Wheat–Aegilops introgressions. In Alien introgression in wheat: cytogenetics, molecular biology, and genomics (eds M. Molnár-Láng, C. Ceoloni, & J. Doležel) 221–244 (Springer, Switzerland, 2015).
60. Hagaras, A. A. A., Kishii, M., Sato, K., Tanaka, H. & Tszumitomo, H. Extended application of barley EST markers for the analysis of alien chromosomes added to wheat genetic background. Breeding Sci 55, 335–341, https://doi.org/10.1276/jsbbs.55.335 (2005).

Acknowledgements
The authors are grateful to the Marginal Region Agriculture Project of Tottori University, Japan (No. 29D2001) for funding this study. We thank Dr. Miyuki Nitta of NBRP-Wheat, and Laboratory of Plant Genetics, Kyoto University, Japan for her excellent technical assistance.
Author Contributions
All the authors conceived and designed the study. S.N. grew all the *Aegilops* and 7 *Triticum* species and isolated DNA, OUE managed and extracted DNA samples from *Hordeum vulgare*, *Secale cereale*, *Elymus ciliaris* and the two cultivars of *T. aestivum*, genotyped and analyzed all the genomes using markers generated by Diversity Arrays Technology Pty Ltd, and interpreted the data. H.T. supervised the study. All the authors wrote different parts of the manuscript, and jointly read and approved the final version.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-34811-y.

Competing Interests: The authors declare no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018