Mapping the ‘breaker’ element of the gametocidal locus proximal to a block of sub-telomeric heterochromatin on the long arm of chromosome 4S\textsuperscript{sh} of Aegilops sharobnensis

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Abstract

The ‘breaker’ element (GcB) of the gametocidal locus derived from Aegilops sharobnensis has been mapped to a region proximal to a block of sub-telomeric heterochromatin on chromosome 4S\textsuperscript{sh}L.

Introduction

The Sitopsis section of the genus Aegilops comprises five diploid species that carry the S genome, and among these species, Aegilops sharobnensis (2n = 14, S\textsuperscript{sh}S\textsuperscript{sh}) carries the S\textsuperscript{sh} genome which is closely related to the D genome of hexaploid wheat. Some S genome chromosomes carry gametocidal loci, which induce severe chromosome breakage in gametes lacking the gametocidal chromosome, and hence, result in gamete abortion. The preferential transmission of gametocidal loci could be exploited in wheat breeding, because linking gametocidal loci with important agronomic traits in elite wheat varieties would ensure retention of these traits through successive generations. In this study, we have mapped the breaker element of the gametocidal locus derived from Ae. sharobnensis to the region immediately proximal to a block of sub-telomeric heterochromatin on the long arm of chromosome 4S\textsuperscript{sh}.
chromosomes and chromatin bridges (Finch et al. 1984; King and Laurie 1993; Nasuda et al. 1998). As a consequence of these aberrations, the only gametes to survive are those carrying chromosomes with a gametocidal locus (Gc chromosomes), which is roughly 50 % of the pollen cells, thus ensuring the preferential transmission of these chromosomes to subsequent generations. This genetic phenomenon is of particular interest to breeders, because linking Gc loci with valuable traits in elite varieties would ensure maintenance of these traits in subsequent generations without the need for selection. This could be achieved by a GM approach; whereby the Gc locus would be present in a vector with the gene for the trait of interest (for example a disease resistance gene), which would then be inserted by transformation in the genome of the elite wheat variety. The progeny carrying this construct would always come through.

Studies of a range of different Gc chromosomes suggest that there are at least two phenotypes associated with the mechanism responsible for the preferential transmission of the Gc chromosome (Endo 1990; reviewed Tsujimoto 2005). The first phenotype is referred to as the ‘breaker’, and induces chromosome breakage. The second is an ‘inhibitor’ phenotype, which prevents chromosome breakage. Chromosome aberrations do not occur in gametes carrying both elements, as the gametocidal action is neutralised by the inhibitor. This hypothesis is supported by the observations that T. aestivum cv. Norin 26 possesses a suppressor of the 3C gametocidal locus derived from Ae. caudata (Tsujimoto and Tsunewaki 1985b), and that the development of a knock-out mutant of the breaker element Gc2 on Ae. sharonensis chromosome 4S\textsuperscript{sh}, has lost the chromosome fragmentation function, but has retained the inhibitor element (Friebe et al. 2003). The gametocidal loci within Sitopsis genomes are classified in groups according to their action and strength (Endo 1990). Ae. sharonensis carries gametocidal loci on chromosomes 2S\textsuperscript{sh} and 4S\textsuperscript{sh} (Endo 1985).

The Ae. sharonensis 4S\textsuperscript{sh} breaker element (GcB) has been previously mapped by C-banding to the distal end of the long arm (Endo 2007). The aim of the present study was to further map GcB on Ae. sharonensis chromosome 4S\textsuperscript{sh}. This chromosome is homoeologous to wheat group 4. Previously, two hexaploid wheat introgression lines (Brigand 8/2 and Brigand 8/9) had been developed, involving the translocation of the long arm of chromosome 4S\textsuperscript{sh} (4S\textsuperscript{sh}L) from Ae. sharonensis onto the long arm of chromosome 4D of T. aestivum (4DS-4DL-4S\textsuperscript{sh}L translocation) (King et al. 1996). The occurrence of this translocation is consistent with recent studies showing that the Ae. sharonensis genome is more closely related to the D genome than the B genome (Marcussen et al. 2014). The strategy for mapping GcB employed a similar approach as that used for the mapping of the Phl locus (Roberts et al. 1999; Griffiths et al. 2006). This involved developing irradiated populations of the introgression line, and screening these populations with DNA markers specifically designed to the 4S\textsuperscript{sh}L chromosome segment. Traditional mapping approaches such as development of NILs (near isogenic lines) were not possible, since the Phl locus present in polyploid wheat prevents recombination between wheat and homoeologous (e.g. alien) chromosomes. Irradiation of populations for mapping the Phl locus generated deletions alone, but irradiation of the introgression line populations induced both deletions and several translocations of differing sizes involving the 4S\textsuperscript{sh}L segment. These mutant lines were then phenotyped for the presence or absence of GcB. This phenotyping involved scoring the fertility of F\textsubscript{1} hybrids generated from crossing the deletion or translocation line with wild type wheat, as well as analysing chromosome fragmentation in first pollen mitosis. The sizes of new translocations and deletions were characterized using markers for the 4S\textsuperscript{sh}L segment, so that a minimum region covering the GcB element could be defined. Using this approach, the GcB element was delimited to a small region immediately proximal to a sub-telomeric heterochromatin block at the distal end of the chromosome.

Materials and methods

Plant material

Two T. aestivum cv. ‘Brigand’ introgression lines (Brigand 8/2 and Brigand 8/9), each carrying a 4DS-4DL-4S\textsuperscript{sh}L translocation from Ae. sharonensis accession 2170001 (King et al. 1996), were individually crossed with T. aestivum cv. ‘Huntsman’, using Huntsman as the pollen donor. Huntsman does not possess the 4DS-4DL-4S\textsuperscript{sh}L translocation, so the resulting F\textsubscript{1} seeds were hemizygous for the Ae. sharonensis introgression. Two mutant populations were created by gamma irradiation. A total of 1011 hybrid F\textsubscript{1} (M\textsubscript{0}) seeds were irradiated with 300 Grays (558 from the Brigand 8/2 × Huntsman cross and 453 from the Brigand 8/9 × Huntsman cross) and 586 hybrid F\textsubscript{1} (M\textsubscript{0}) seeds were irradiated with 350 Grays (395 from the Brigand 8/2 × Huntsman cross and 191 from the Brigand 8/9 × Huntsman cross). The M\textsubscript{1} seeds irradiated with 300 Grays and 350 Grays germinated at a rate of 87.8 % (personal communication, Surbhi Mehra) and 49 %, respectively. Spikes were semi-sterile, which confirmed the effect of the gametocidal locus in these hemizygous plants. M\textsubscript{1} plants were self-pollinated to give homozygous M\textsubscript{2} seeds. M\textsubscript{2} plants were screened for deletions and/or chromosomal translocations. Lines with deletions or chromosomal translocations
in the *Ae. sharonensis* introgression from the Y300 population were backcrossed onto *T. aestivum* cv. Huntsman to check for the presence or absence of the gametocidal effect. Deletion and translocations 5F8, 5D12, 7C12, 12A5, and 12H3 all originated from an initial cross between introgression line Brigand 8/2 and *T. aestivum* cv. Huntsman.

Marker development

Thirty-three markers were designed that showed polymorphism between *T. aestivum* cv. Huntsman and the introgressed *Ae. sharonensis* segment from accession 2170001 in Brigand 8/2 and Brigand 8/9. Nine of these markers were initially designed as SSCP–SNP markers (Bertin et al. 2005) from *Ae. sharonensis* accession 1644 sequences obtained from a transcriptome assembly by Bouyioukos et al. (2013). These were aligned to their corresponding wheat homologs based on a best BLAST hit to predict intron position and size (Brenchley et al. 2012). When wheat sequence was not available, the rice (*Oryza sativa*) homolog was used. These SSCP–SNP markers were later converted into high-throughput KASP™ markers (He et al. 2014), by designing triplets of FAM, VIC and common primers around SNPs between *T. aestivum*, cv. Huntsman and the *Ae. sharonensis* accession 2170001 allele. Each marker was given the prefix ‘KASP’ followed by the *Ae. sharonensis* cDNA read number from the transcriptome analysis.

Marker UTV39 was designed from nucleotide sequence of clone UTV39 (Giorgi et al. 2003), forward primer 5’GATCCGTTTGGTACATCAGGT and reverse primer 5’GATCTTTCCAGTAGTTCTCTAA. This marker was later converted into a KASP™ marker as for the SSCP–SNP markers and was designated as ‘KASP UTV3’ (Online Resource 1).

The remaining twenty-four markers were obtained from two analyses using the high density 90,000 SNP genotyping array (Wang et al. 2014). The first analysis identified KASP™ markers with polymorphisms between the genomic DNA of Brigand with no introgression and Brigand 8/2 amongst the 90,000 SNPs of the array. These markers identified polymorphisms between the *Ae. sharonensis* introgression and wheat group 4 chromosomes. The second analysis was carried out using genomic DNA from Huntsman and the Y300 translocation line 12A5 to identify further markers in the distal region of the *Ae. sharonensis* introgressed segment. These twenty-four markers were each designated with a unique eight digit number.

Twenty-two of the thirty-three KASP™ markers were further checked for specificity to the D genome, and redesigned when necessary to amplify the wheat D allele only (VIC primer) and the *Ae. sharonensis* allele (FAM primer). *Ae. sharonensis* sequence was obtained by BLASTing the SNP sequence to the *Ae. sharonensis* whole-genome database (Marcussen et al. 2014). The *Ae. sharonensis* sequence was then aligned to the corresponding A, B and D best BLAST hits obtained by BLASTing the SNP sequence to 4A, 4B, 4D and 5A on the URGI BLAST webpage (https://urgi.versailles.inra.fr/blast/blast.php) for each marker. These twenty-two D genome specific KASP™ markers were used to genotype M₂ plants (Online Resource 1).

KASP UTV39 could not be designed as a D specific marker as it is a repeat sequence specific to the S sh genome and has no significant BLAST hit to a D sequence in wheat. The best wheat BLAST hit is a 5BS contig.

Genotyping of Y300 M₂ population

DNA extractions were carried out on leaf material of M₂ plants at the 2–3 leaf stage, according to the Somers and Chao protocol http://maswheat.ucdavis.edu/PDF/DNA0003.pdf original reference in Pallotta et al. (2003).

PCR reactions for SSCP–SNP markers were carried out using a touchdown program: 95 °C for 15 min, then ten cycles of 95 °C for 30 s, 63 °C for 30 s (−0.5 °C per cycle) and 72 °C for 90 s, followed by 30 cycles of 95 °C for 30 s, 58 °C for 30 s and 72 °C for 90 s. NB: for marker UTV39, the annealing temperature was 50 °C instead of 58 °C. Markers were run on SSCP gels as described in Bertin et al. (2005).

All KASP™ amplifications were carried out using 2 µl of KASP™ master mix (LGC group, United Kingdom), 0.056 µl of primer mix (12 µl FAM primer at 100 µM + 12 µl of VIC primer at 100 µM + 30 µl of common primer at 100 µM + 46 µl of dH₂O) and 2 µl of DNA at 5 ng/µl, following the same program with an optimal cycle number varying from 30 to 40 for each marker (Online Resource 1): 94 °C for 15 min, 10 cycles of 94 °C for 20 s, 65 °C for 1 min and 94 °C for 20 s, then either 30, 35 or 40 cycles of 94 °C for 20 s and 57 °C for 1 min. Fluorescence was measured by a Tecan Safire plate reader. Results were analysed using the KlusterCaller software (LGC group, UK).

Fluorescence in situ hybridisation (FISH)

Root tip squashes from germinating seedlings of *T. aestivum* cv. Brigand 8/2 and M₃ seedlings of lines 12A5, 12H3 and 7C12 were prepared as described in Gill et al. (1991) with minor modifications.

Fluorescence in situ hybridisation (FISH) was carried out as described in Prieto et al. (2007). The probe used was a dUTP-digoxigenin labelled product generated by amplification of high quality DNA of *Ae. sharonensis* accession 1644 with *UTV39* primers. Labelling with dUTP-digoxigenin was carried out directly, with no nick translation.
being necessary due to the short length of the PCR product. Observations were made using a Nikon Eclipse E600 CCD microscope and images captured with a Hamamatsu ORCA-ER C4742-95 digital camera. Images were merged using ImageJ software.

Phenotyping

The phenotype of each M2 hybrid plant was determined by measuring spike fertility and by looking at chromosome fragmentation during first pollen mitosis. Spike fertility was measured as the percentage of florets that contained well-developed seeds. Only the two outermost (oldest) florets within each spikelet were counted. Undersized florets at the top and bottom of the spike were not included in the calculations. Hybrid plants were grown in three batches, with a 2 week interval between each sowing. One to four plants were grown for each line per batch and three to eight ears per plant were phenotyped. Parental plants of Huntsman and Brigand 8/2 were grown as controls alongside each batch so that environmental effects on spike fertility could be taken into account. The average percentage seed set was calculated from 102, 82 and 78 phenotyped ears of combined Huntsman and Brigand 8/2 plants for batches 1, 2 and 3, respectively. The average percentage of viable seeds was calculated from all phenotyped ears for each hybrid line [7C12, 12A5, 12H3 and whole segment translocation line (WST), backcrossed onto Huntsman], per batch. Each average was adjusted according to the average percentage of viable seeds for self-pollinated parental lines used as controls calculated per batch, to take into account any environmental factors. This was carried out by multiplying the mutant hybrid line average by 100 and then dividing it by the average for the controls. An average of approximately 50 % viable seeds indicated that GcB was present as it lead to abortion of around half of the gametes.

Observations of chromosome fragmentation at anaphase of first pollen mitosis were carried out on aceto-carmine stained anther squashes, using a Leica DM2000 microscope and a Leica DFC450 image capture system.

Results

Marker development

Thirty-three markers were designed to be polymorphic between hexaploid wheat and the 4SshL Ae. sharonensis segment to map the gametocidal breaker element (GcB) within this introgression. Twenty-two of these markers were further designed to be specific to the hexaploid wheat D genome and polymorphic with the orthologous 4Ssh sequence.

The terminal region of the 4SshL chromosome segment carries a large block of heterochromatin. Probe pAesKB52 has been shown to hybridise to major sub-telomeric heterochromatin sites in Ae. sharonensis (Ananthawat-Jönsson and Heslop-Harrison 1993) and its sequence shares 90.1 % identity to clone UTV39 (Giorgi et al. 2003). This indicated that the UTV39 repeat would constitute a good probe for the detection of the block of sub-telomeric heterochromatin present on 4SshL. This was confirmed by a FISH experiment using the UTV39 repeat as a probe on mitotic cells from root tip squashes of 4DS:4DL:4SshL Brigand 8/2 translocation line (Fig. 1a).

As expected, the probe hybridised to the distal end of two chromosomes in each cell (chromosomes 4D carrying the 4Ssh introgression). The UTV39 repeat could not be converted to a D genome specific marker for the genotype screens as it is specific to the S genome, but could be used as a presence/absence marker for the S copies of the UTV39 repeat. Orthologous sequences of this repeat are distributed throughout the hexaploid wheat genome, but these are in relatively small numbers compared to the estimated 40,000 copies present on four of the seven chromosomes of Ae. sharonensis (Giorgi et al. 2003). Fluorescence in situ hybridisation (FISH) experiments using the UTV39 repeat as a probe on hexaploid wheat root tips do not detect a signal (as shown in Fig. 1b: FISH using UTV39 probe on root tip squashes of Huntsman), but when UTV39 is used as an SSCP–SNP marker or in KASP™ technology, an amplified product is generated both for hexaploid wheat (e.g. cv. Huntsman) and Ae. sharonensis DNA. These products are polymorphic. However, only the Ssh copies of UTV39 are amplified, due to the extremely high copy number of UTV39 repeats, in translocation lines such as Brigand 8/2 and Brigand 8/9 where both the Ae. sharonensis (Ssh) copies and the wheat orthologous sequences are present.

Synteny between Triticeae chromosome group 4, Brachypodium distachyon chromosome 1, and rice chromosome 3 was used as a tool to provide an approximate gene content of this 4SshL terminal region. Approximately half the markers for the introgressed segment were identified by this approach. The remaining markers were derived from other sources (as described in “Materials and methods”). However, it was apparent that the order of genes is extensively rearranged within the terminal regions. This is consistent with the reported decrease in synteny levels in the distal regions of chromosomes (Akhunov et al. 2003). Consequently, the newly developed markers could not be ordered according to synteny. However, seven of the twenty-two KASP™ markers were converted into Ae. sharonensis specific KASP™ markers, designed around SNPs between accesions 1644 and 2189 (Bouyioukos et al. 2013). These
markers were used to screen an F6 recombinant population resulting from a cross between *Ae. sharonensis* accessions 1644 and 2189 (resistant and susceptible to race Ug99 of stem rust pathogen, respectively). This allowed the mapping of these seven markers relative to one another (highlighted in yellow in Online Resource 1). This mapping revealed KASP23047 as the most proximal of these seven markers. The order of all twenty-two KASP™ markers was established later in the study from the analysis of the presence or absence of these markers observed on the deleted and translocated chromosomes following irradiation (Fig. 2), with the UTV39 repeat being the most distal marker.

Screening of irradiated populations

A total of 3,764 M2 plants from the Y300 irradiated population and 3300 M2 plants from the Y350 irradiated population were genotyped using the twenty-two KASP™ markers and KASP UTV39 (Online Resource 1). Ninety-eight percent of the Y300 M2 plants and 99.6% of the Y350 M2 plants carried just the Brigand 4DS·4DL-4SshL translocated chromosome, possessing a homozygous S (*Ae. sharonensis* SNP allele) genotype. Thus, the Brigand 4DS·4DL-4SshL translocated chromosome is preferentially transmitted, while the 4D (Huntsman) chromosome lacks the introgression and therefore, the *Gc* locus is fragmented.

**Fig. 1** Fluorescence in situ hybridisation using probe UTV39 (green) on DAPI stained chromosomes of mitotic cells from root tip squashes of parental lines Brigand 8/2 (a) and Huntsman (b), translocation lines 12A5 (c), 12H3 (d) and deletion line 7C12 type 1 (e) and type 2 (f). Scale bar represents 10 µm
The H (Huntsman SNP) allele is not present in most M₂ plants (Fig. 3).

**Deletions lines resulting from the Y350 irradiation**

The genotype screening of M₂ plants derived from the Y350 irradiated M₁ population, identified three lines carrying deletions involving the 4S⁹L segment. However, genotyping revealed that the whole \textit{Ae. sharonensis} segment had been deleted in one of these lines, so \textit{GcB} could not be further mapped using this line. The remaining two deletion lines (5F8 and 5D12) derived from the Y350 population possessed a deletion encompassing six markers (130489–KASP25343), and thirteen markers (KASP23047–KASP25343), respectively within the introgressed segment (Fig. 2). Both these M₂ deletion lines are a homozygous S genotype for the remaining \textit{Ae. sharonensis} alleles. The absence of Huntsman alleles in these lines indicated that the Huntsman chromosome had been fragmented at the M₁ stage (Fig. 3), and so the \textit{Gc} locus is still present in both the non-deleted segments of the \textit{Ae. sharonensis} introgression, i.e. within the region from the telomere to marker KASP25343.

**Translocation lines resulting from the Y350 irradiation**

The genotype screening of M₂ plants from the Y350 irradiated M₁ population identified two lines with chromosomal rearrangements typical of translocations. However, genotyping revealed that translocations were far more common in the Y300 population, with fifty-two lines being identified. Forty-eight of the fifty-two lines derived from the Y300 population and the two translocation lines from the Y350 population carried the whole \textit{Ae. sharonensis} introgression translocated onto the Huntsman 4DL chromosome, such that each marker along this translocated chromosome amplified an H and an S sequence. These lines are designated as whole segment translocations (WST) (Fig. 2). The translocation did not result in the deletion of \textit{GcB}, as the percentage of viable seeds in hybrids resulting from the backcross of these M₂ lines onto wheat cv. Huntsman ranged from 35.9 to 40.5 % (Fig. 4), indicating that...
the \( GcB \) element was still present (percentage of seed set in self-pollinated spikes of parental Huntsman and introgression line Brigand 8/2 was also determined to take account of any environmental impact on seed viability). The reciprocal product of this event, a truncated Brigand 8/2 chromosome, would have been fragmented by the \( GcB \) element present in the other gametes and would therefore not have come through in the \( M_2 \) generation.

Of the remaining four translocation lines (12A5, 12H3, 12H4 and 12H5) derived from the Y300 population, three (12H3, 12H4 and 12H5) originated from the same \( M_1 \) ear and shared the same translocated genotype. Thus, only one of these lines, 12H3, is represented in Figs. 2, 3 and 4. Line 12A5 (Fig. 2) has a translocation of the distal region of the \( Ae. \ sharonensis \) segment (comprising markers 143315, KASP8170, 137535 and 124239) onto the end.

### Fig. 3
Diagram representing chromosomal rearrangements caused by gamma irradiation in lines 5D12 (and 5F8—similar segregation pattern but different size deletion), 7C12, 12A5 and 12H3 and their segregation from \( M_1 \) to \( M_2 \) plants. *Lines with no rearrangement* in the \( Ae. \ sharonensis \) introgression are also represented and constituted 98 and 99.6 \% of the Y300 and Y350 populations, respectively.
of the Huntsman chromosome resulting in the duplication of this terminal region. Thus, markers were genotyped as both H and S as in a hemizygous situation (Figs. 2, 3). Line 12H3 has undergone a similar translocation event, with a duplication of H and S segments for marker KASP8170 only. However, in 12H3 the two distal markers, 137535 and 124239, were genotyped as S only (Fig. 2), which may suggest that the equivalent region of the Huntsman chromosome was translocated onto the other chromosome carrying most of the *Ae. sharonensis* introgression. This translocated chromosome lacked the *Gc* locus and would have therefore been fragmented during pollen mitosis of the M1 self-pollination stage (Fig. 3). The two translocation lines 12A5 and 12H3 were phenotyped by backcrossing the M2 plants onto Huntsman and assessing the percentage of viable seeds in the resulting hybrid ears as described for the deletion lines. Hybrid ears exhibited 26.8–47.7 % viable seeds for the 12A5 backcross and 39–56.6 % viable seeds for the 12H3 backcross (Fig. 4), indicating the presence of the gametocidal effect in both lines. Further confirmation that the semi-sterile spike phenotype resulted from the presence of the *Gc* element in line 12H3 (which carried the smallest *Ae. sharonensis* segment) was obtained by observing fragmentation in half the pollen cells of hybrid plants resulting from the 12H3 backcross (Fig. 5). Numerous chromosome fragments were observed between separating chromatids at anaphase of first pollen mitosis in the 12H3 backcrossed onto Huntsman hybrids, typical of previous observations of gametocidal induced fragmentation (Finch et al. 1984; King and Laurie 1993; Nasuda et al. 1998). This provided evidence that the *GcB* element was still present in this line.

These results, and especially the analysis of the smallest introgression segment (line 12H3), further defined the *GcB* element as being between marker 143315 and the telomere including the heterochromatin block on the *Ae. sharonensis* segment (Fig. 2). Moreover, the 4S<sub>ab</sub>L segment in 12H3 may also contain the inhibitor element. This translocated segment causes fragmentation of chromosomes lacking the 4S<sub>ab</sub>L sub-telomeric segment but not of itself, suggesting that it also carries the inhibitor element. However, this would need to be confirmed by independent mapping of this element.

Deletions lines resulting from the Y300 irradiation

The genotype screening of M2 plants from the Y300 irradiated M1 population also identified nine lines carrying deletions involving the 4S<sub>ab</sub>L segment. As in the case of those identified in the Y350 irradiated M1 population, the whole *Ae. sharonensis* segment had been deleted in eight of these lines. However, the remaining line, 7C12, had a deletion of only part of the 4S<sub>ab</sub>L segment. Genotyping analysis of this line revealed that the M2 progeny segregated with a 1:2:1 inheritance ratio (Fig. 3). Genotyping identified 25 % of the progeny as having the Huntsman allele only (type 3) and 50 % as having both S and H alleles for most of the introgressed segment, with the region from 100738 to 124239 being Huntsman only (type 2). The lack of fragmentation of the Huntsman chromosome suggested that the
GcB element must have been deleted in the M1 line as a result of the irradiation. This gave rise to the three progeny types illustrated in Fig. 3. The remaining 25% of the M2 progeny for 7C12 (type 1) were homozygous for a partially deleted Brigand 4DS-4DL-4SshL translocated chromosome (deletion covering markers 100738–124239).

The presence of the GcB element was confirmed by backcrossing plants homozygous for the partially deleted Brigand 4DS-4DL-4SshL translocated chromosome (7C12 type 1) and hemizygous for the partially deleted Brigand 4DS-4DL-4SshL translocated chromosome and the Huntsman 4D chromosome (7C12 type 2) onto the Huntsman parent. Hybrid plants resulting from the backcrosses were left to mature and the percentage of viable seeds was calculated for each ear. Spikes from hybrid plants resulting from crosses between 7C12 M2 types 1 and 2 onto wheat cultivar Huntsman had a combined average of 84.1–100% viable seeds (Fig. 4), confirming the absence of fragmentation at the M1 stage.

Genotyping revealed that the 4SshL segment in 7C12 carried a deletion involving the very distal region from markers 100738 to 124239. However, surprisingly, genotyping also revealed that this line carried S copies of the UTV39 repeat sequence, suggesting that the introgressed segment may still carry the Ae. sharonensis heterochromatin block and telomere region. Confirmation of this required FISH analysis using the UTV39 marker as a probe, on root tip squashes derived from this line to confirm the presence of the heterochromatin block. KASP UTV39 and SSCP–SNP UTV39 genotyping revealed all six lines (5F8, 5D12, 7C12 types 1 and 2, 12A5 and 12H3) to carry S copies of the UTV39 repeat sequence (Fig. 2). FISH on mitotic cells from root tip squashes of lines 12A5, 12H3, 7C12 type 1 and 7C12 type 2 (Fig. 1c–f) mirrored the genotyping analysis, revealing the presence a hybridisation signal for probe UTV39 in all four lines. The hybridisation signal observed in cells of lines 12A5, 12H3 and 7C12 type 1 was identical to that observed in the Brigand 8/2 parental line (Fig. 1a) and in Giorgi et al.
(2003); i.e. a hybridisation signal at the telomeric end of two of the chromosomes (the 4D pair). Line 7C12 type 2 is the hemizygous genotype and therefore only has one chromosome carrying the S copies of UTV39, which is observed as a hybridisation signal at the end of only one chromosome (Fig. 1f). The presence of the heterochromatin block is expected for lines 5F8, 5D12, 12A5 and 12H3, as the distal markers in the 4SbL region are present. However, it is surprising that these same markers are deleted in 7C12, yet the heterochromatin block is retained. Thus, the analysis reveals that the GcB element must map in the region immediately proximal to the heterochromatin block.

Discussion

The GcB element of the Gc locus has thus been mapped to the region distal to marker 143315 on chromosome 4SbL (Fig. 2) and proximal to the heterochromatin block. Marker KASP24858 maps proximal to marker 143315 (Fig. 2), and is orthologous to rice gene LOC_Os03g02600 (Online Resource 1). Yet, this gene is present on wheat BAC_205008 which has been localised to the telomeric bin 4BL-10 (0.95–1.0) (Salina et al. 2009). This BAC also carries copies of the Spelt 52 repeat in two arrays in a “head-to-tail” orientation (Salina et al. 2009). Spelt52 is analogous to pAesKB52 (Salina et al. 2006) which in turn is analogous to UTV39 (Giorgi et al. 2003). This suggests that marker KASP24858 is sub-telomeric, along with the other five markers distal to it (100738–124239) down to the heterochromatin block in a similar structure to that of BAC_205008 (Salina et al. 2009). Consequently, the GcB element is defined to the region immediately proximal to the sub-telomeric heterochromatin block (represented in Fig. 2) and might constitute a very small region.

The localisation of the GcB element in the region proximal to the heterochromatin block will enable further mapping of the area and identification of the GcB element. One strategy would be to screen the wheat cultivar Chinese spring BAC library with markers 143315, KASP8170, 137535 and 124239 to physically contig this region, enabling its size to be estimated. The corresponding Ae. sharonensis sequences could then be identified from the Ae. sharonensis whole-genome database (Marcussen et al. 2014) and aligned to form an Ae. sharonensis contig from which the GcB element could be fine mapped. This strategy could be combined with undertaking RNA-Seq on pollen cells derived from the lines carrying different translocated and deleted chromosomes. Candidates for the locus would be anything expressed in the pollen cells from 5F8, 5D12, 12A5 and 12H3, but absent in 7C12.

Several hypotheses have been proposed for the mode of action of the gametocidal locus. These include a dual-function model (breaker and inhibitor), in which the locus is similar to a restriction-modification system in bacteria (Wilson and Murray 1991; Tsujimoto 2005). However, there is one hypothesis, which is consistent with the mapping of the GcB element to the sub-telomeric region. Tsujimoto and Tsunewaki (1985a) suggested a transposon theory based on hybrid dysgenesis in Drosophila melanogaster, caused by P elements. The symptoms include sterility, lethality, mutations, and chromosome breakage, which are all identical to gametocidal induced symptoms. P elements are mobile elements. They can produce the transposase required for their mobility, which makes them autonomous. The fragmentation of chromosomes in other gametes could simply result from the activation of mobile elements by a factor which is able to travel between the gametes, while the suppressor activity is unable to travel. P elements can be inhibited by P-repressors which can be located elsewhere in the genome. This is referred to as the trans silencing effect (TSE) (Ronsseray et al. 1998). TSE could be the means by which the gametocidal inhibitor suppresses the effects of the gametocidal breaker. This hypothesis would explain the suppression by T. aestivum cv. Norin 26 of the GcB effect from Ae. caudata (Tsujimoto and Tsunewaki 1985b). Ronsseray et al. (1998) reported a pair of autonomous P elements inserted at the telomere of an X chromosome in D. melanogaster, in telomeric associated sequences (TAS). This model is a good hypothesis for the gametocidal system: the breaker may be a transposon similar to these telomeric P elements and the inhibitor would be located close to the breaker in the sub-telomeric region. The mapping of the GcB to this chromosome region is consistent with this hypothesis for the mode of action of the gametocidal locus.

Author contribution statement

EK wrote the article, designed markers and carried out the cytological analyses. EK and TD designed the experiments, developed and processed plant populations and collected and analysed data. TD and GM revised the article. MM performed the analysis of the Ae. sharonensis transcriptome, developed a genetic map providing a core order of the markers and advised on marker design. AB, MDR and JS assisted with marker development, screen of M2 populations and phenotyping of deletion and translocation lines. SM produced the Y300 population and grew and harvested the M1. GM and IK designed the project and obtained funding. GM supervised experimental designs and data analyses.

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Conflict of interest The authors declare they have no conflict of interest.

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