RESEARCH PAPER

A carbohydrate-binding protein, B-GRANULE CONTENT 1, influences starch granule size distribution in a dose-dependent manner in polyploid wheat

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Abstract

In Triticeae endosperm (e.g. wheat and barley), starch granules have a bimodal size distribution (with A- and B-type granules) whereas in other grasses the endosperm contains starch granules with a unimodal size distribution. Here, we identify the gene, BGC1 (B-GRANULE CONTENT 1), responsible for B-type starch granule content in Aegilops and wheat. Orthologues of this gene are known to influence starch synthesis in diploids such as rice, Arabidopsis, and barley. However, using polyploid Triticeae species, we uncovered a more complex biological role for BGC1 in starch granule initiation: BGC1 represses the initiation of A-granules in early grain development but promotes the initiation of B-granules in mid grain development. We provide evidence that the influence of BGC1 on starch synthesis is dose-dependent and show that three very different starch phenotypes are conditioned by the gene dose of BGC1 in polyploid wheat: normal bimodal starch granule morphology; A-granules with few or no B-granules; or polymorphous starch with few normal A- or B-granules. We conclude from this work that BGC1 participates in controlling B-type starch granule initiation in Triticeae endosperm and that its precise effect on granule size and number varies with gene dose and stage of development.

Keywords: Aegilops, B-type starch granule content, crop breeding, FLOURY ENDOSPERM 6, granule size distribution, polymorphous starch, starch granule initiation, TILLING mutant, Triticeae, wheat grain.

Introduction

Triticeae species, such as bread wheat (Triticum aestivum L.), are unusual among grasses in having two types of starch granule in their endosperm called A- and B-type granules. These originate from two starch granule initiation events that are separated in time and space. The first event gives rise to a single large A-type granule per plastid and takes place early in endosperm development in the main body of the plastid. The second granule initiation event gives rise to small B-type granules and takes place several days after the first event during endosperm development at least partly within the plastid stromules. Thus, endosperm plastids in wheat each contain one large A-type granule and several small B-type granules.

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The endosperms of almost all species of Triticeae, including the domesticated cereals wheat, barley (Hordeum vulgare L.), and rye (Secale cereale L.), and also wild grasses such as Aegilops, contain both A- and B-type starch granules. However, a few species of wild Triticeae have normal A-granules, but reduced numbers of B-granules such as Aegilops peregrina (Hack.) which was shown to be B-granule-less (Stoddard, 1999; Stoddard and Sarker, 2000). To understand the control of B-granule content, the impact of B-granules on starch functional properties and the mechanisms involved in starch granule initiation in wheat (and in plants generally), we created a population of Aegilops with varying B-granule content by crossing the B-less tetraploid Ae. peregrina with a synthetic tetraploid Aegilops called KU37 that has B-granules (Howard et al., 2011). KU37 has the same genome composition (SSUU) as Ae. peregrina but was derived from a cross between the diploids A. sharonensis (SS) and A. umbelillata (UU) (Tanaka, 1955, 1983). Using this population, we identified a quantitative trait locus (QTL) on the short arm of chromosome 4S that accounted for 44% of the control of B-granule content (Howard et al., 2011). As a result, we hypothesized that other Triticeae species, such as bread wheat, have a gene controlling B-granule content, BGC1 (B-GRANULE CONTENT 1), in a syntenous position on the group 4 chromosomes. We tested this hypothesis in the bread wheat cultivar Paragon by selecting and combining together in one plant, large chromosomal deletions spanning the putative BGC1 regions of chromosomes 4A and 4D (Chia et al., 2017). The single-deletion mutants had normal starch, but the double-deletion mutant line was found to lack B-type starch granules. Thus, we successfully transferred the B-granule-less trait seen in Aegilops to bread wheat, and showed that the BGC1 gene responsible for granule initiation must be one of the 240 genes that were common to both the 4A and 4D deletions (Chia et al., 2017).

Here, we describe the identification of the BGC1 gene in Aegilops and wheat. First, we mapped BGC1 in progeny derived from the Ae. peregrina x KU37 cross that had been used for the initial QTL mapping. Surprisingly, we found that the BGC1 gene is orthologous to FLOURY ENDOSPERM6 (FLO6), a known player in starch synthesis in cereal endosperm from work on mutants of rice (Oryza sativa), a known player in starch synthesis in cereal endosperm (Seung et al., 2009). In Arabidopsis (Arabidopsis thaliana L.) chloroplasts, the FLO6 orthologue PTST2 is known to be involved in starch granule initiation (Seung et al., 2017). Using TILLING (targeting induced local lesions in genomes) mutants of both tetraploid and hexaploid wheat (Krasileva et al., 2017) with different dosages of functional BGC1, we discovered that this gene in the Triticeae not only controls B-type granule content, but that depending on the stage of grain development it can contribute to either promoting or suppressing granule initiation.

Materials and methods

Plant material

The sources of the Aegilops and wheat deletion mutants, and methods for growth in the glasshouse and the field, were as described previously (Howard et al., 2011; Chia et al., 2017) except that Aegilops grown in the glasshouse were not vernalized. For fine mapping, in addition to the F2- and F3-derived (KT) lines from the cross between Ae. peregrina and the synthetic Aegilops, KU37 (Howard et al., 2011), a set of backcrossed (BC) lines were developed from a cross between one F2 line, KU1-4, which has B-type starch granules, and the natural B-granule-less parent, Ae. peregrina. One BC1F1 plant was backcrossed to Ae. peregrina and the BC2F1 grains were grown and screened with markers flanking the BGC1 region (between the QTL peak marker 4G and the telomere) (Supplementary Table S1 at JXB online) to identify 10 homozygous recombinant plants. These were grown and self-pollinated to give BC3F2 grains which were screened again as before; heterozygous recombinant lines were self-pollinated, and homozygous recombinant lines were selected.

TILLING mutants with polymorphisms in the genes of interest in the tetraploid wheat Kronos and the hexaploid wheat Cadenza (Krasileva et al., 2017) were identified from http://www.wheat-tilling.com/ and http://plants.ensembl.org/Triticum_aestivum/, and obtained from www.seedstor.ac.uk. The mutations were chosen based on their likelihood to affect the function of the encoded protein; that is, they were nonsense (stop) or missense mutations, or mutations that affect splicing sites of the mRNA (to remove introns).

Kronos and Cadenza TILLING mutants were grown in speed-breeding conditions (21–22 °C; 22 h day/17–18 °C; 2 h night; Watson et al., 2018) in Levington’s M2 compost (LBS, Colne, Lancashire, UK) in either a glasshouse where natural lighting was supplemented by high-pressure sodium lamps or an Adaptis 1000 growth chamber with a red/white LED canopy (Convirion, Winnipeg, Canada). Spikes were harvested 4 weeks post-anthesis and dried at 30 °C in an oven prior to sowing.

DNA and RNA preparation

DNA was extracted from seedling leaves as described by Fulton et al. (1995). RNA was extracted from flag leaves and developing grains both harvested 19–21 d after ear emergence (mid grain development). RNA was extracted using an RNeasy Plant mini kit (Qiagen.com) according to the manufacturer’s instructions. Samples of RNA from Ae. peregrina grain, Ae. peregrina leaves, KU37 grains, and KU37 leaves were pooled to give four samples with a total of 5 µg of total RNA per sample (at a minimum concentration of 25 ng µl⁻¹). These were submitted for sequencing to The Genome Analysis Centre (now the Earlham Institute), Norwich, UK. After sample quality control, Illumina barcoded RNA TruSeq libraries were constructed and the four libraries were sequenced over two lanes, in pools of two, on the Illumina HiSeq 2000 platform. Using 100 bp paired-end reads, at least 100 million reads per lane were generated. After data quality control, base calling, and formatting, the Aegilops sequence data were aligned to a T. aestivum reference genome (either pseudo-chromosomes that were organized using wheat–Brachypodium synteny or the NimbleGen wheat exome capture probe set). When it became available, the sequence data were re-organized to Refseq v1.0 (International Wheat Genome Sequencing Consortium (IWGSC), 2018) with HISAT-2.0.5 (Kim et al., 2015). The Refseq v1.0 alignments where sorted and the candidate regions were extracted (chr4A:589084002-591920577, chr4B:20715580-23835481, and chr4D:10926756-13253764) using samtools-1.4.1 (Li et al., 2009). All RNA sequencing (RNA-Seq) data were visualized and compared using IGV software (Robinson et al., 2011) to identify single nucleotide polymorphisms (SNPs) suitable for the design of molecular markers for use in genetic mapping. All raw RNA-Seq reads have been deposited in the Sequence Read Archive (SRA) under accession ERR83409626.

Aegilops genotype analysis

To distinguish the parental and heterozygous plants within the mapping population, KASP primers were designed to polymorphisms in genes within the region of interest. These were tested on a subset of the population to see whether or not they amplified genes linked to BGC1 using the PCR-based KASP genotyping assay as described by the manufacturer (https://www.lgcgroup.com). As the Aegilops plants used here are tetraploid and BGC1 lies on subgenome S but not on U, only half of the successful KASP assays would be expected to be linked to BGC1, and this
was found to be the case (data not shown). The \textit{BGC1}-linked markers on chromosome 4S and their corresponding genes in wheat are given in Supplementary Table S1.

**Phenotype analysis**

Starch granules were extracted from single or half-grains as described previously (Chia et al., 2017) except that the method was scaled down from 10 grains per sample to one half-grain per sample. Starch granule size distribution was measured using either (i) a manual image analysis method described previously (Chia et al., 2017); (ii) an automated image analysis method using a cell counter (Biorad TC20); or (iii) using a Coulter counter (Beckman Coulter Multisizer 4e) fitted with a 70 µm aperture tube, with Isoton II diluent (Beckman Coulter). Methods (i) and (ii) both involve microscopic examination of starch followed by image analysis to define the proportion of granules <10 µm in diameter. B-type starch granules in Triticeae species are generally found to be smaller than 10 µm in diameter whereas A-type granules are 5–40 µm in diameter. Thus, the sizes of A- and B-type granules overlap and the <10 µm category contains both B-granules and some small A-granules. For clarification, these measurements will therefore be referred to as measurements of small granules rather than measurements of B-granules.

Method (iii), the Coulter counter, estimates particle volume from the transient drop in electrical resistance caused by the passage of a particle through an aperture. For samples containing both A- and B-type granules, the relative total volume of B-type starch granules was estimated by fitting a mixed Gaussian distribution. A minimum of 50,000 particles were measured per sample.

**Cloning and sequencing of \textit{Aegilops} BGC1**

Primers were designed to \textit{BGC1} genes from \textit{Ae. tauschii} or \textit{T. aestivum} and used to amplify the \textit{Aegilops} \textit{BGC1} genes in overlapping fragments (Supplementary Table S2A). Suitable fragments were gel purified and cloned into \textit{Escherichia coli}, and individual clones were isolated and multiplexed using a Clonejet PCR cloning kit and Genejet plasmid kit (both from Thermofisher.com), according to the manufacturer’s instructions. Multiple (>3) clones for each gene were sequenced, and the trimmed consensus sequences were assembled into contigs using Geneious software (www.geneious.com). The intron/exon positions were inferred by comparison with the rice \textit{FLO6} cDNA sequence (LOC_Os03g48170).

We were unable to amplify fragments of \textit{BGC1}-S spanning exons 1–7 from B-granule-less \textit{Ae. peregrina} using our cloning primers. To test this region further, several more 4S-specific primer pairs (and various controls) were designed and used to amplify genomic DNA from \textit{Aegilops} and wheat (Chinese Spring). The primer sequences, PCR conditions, and associated amplicon information are given in Supplementary Table S2B and Supplementary Fig. S2.

**Genotype analysis of TILLING mutants of wheat**

Primers (Supplementary Fig. S3) were designed to distinguish mutant and wild-type sequences, and the primary plants and their progeny were genotyped using the PCR-based KASP™ genotyping assay as described by the manufacturer (https://www.lgcgroup.com). Phenotyping was performed as described above for \textit{Aegilops} grains [method (ii)].

**Identification of triple \textit{BGC1} mutants in hexaploid wheat**

The Paragon double-deletion mutant (with deletions of the \textit{BGC1} regions of chromosomes 4A and 4D; Chia et al., 2017) was crossed to a Cadenza TILLING mutant Cadenza1730 with a nonsense mutation in \textit{BGC1}-B (Q184*). The \textit{F2} grains were allowed to self-pollinate, the resulting \textit{F2} grains were cut in half, and the embryo halves were grown. DNA was extracted from the \textit{F2} seedling leaves and this was genotyped to identify triple-mutant \textit{F2} plants as follows. Plants homozygous for the TILLING mutation in \textit{BGC1}-B were identified using a KASP genotyping assay (Supplementary Table S3). Plants homozygous for either the A-genome or the D-genome deletions were identified by the absence of PCR amplification with homoeologue-specific primers to \textit{BGC1}-A and \textit{BGC1}-D. As a positive control, DNAs from putative deletion mutants were also tested by PCR with primers specific for \textit{BGC1}-B and unaffected by the TILLING mutation. Starch was extracted from the \textit{F2} half-grains identified as triple mutants, and examined by microscopy.

**Scanning electron microscopy**

Samples of purified starch were suspended in water and 2 µl was applied to a glass cover slip mounted on the surface of an aluminium pin stub using double-sided adhesive carbon discs (Agar Scientific Ltd, Stansted, Essex, UK). The water was allowed to evaporate. Mature grains were fractured in mid-section using a scalpel blade and mounted directly on double-sided adhesive carbon discs attached to stubs. The stubs were sputter coated with ~15 nm gold particles in a high-resolution sputter coater (Agar Scientific Ltd), transferred to a FEI Nova NanoSEM 450, and viewed at 3 kV.

Fractured and etched samples of Franubet barley starch were prepared for SEM as follows. Extracted starch (0.4 g) was suspended in 1 ml of dH2O, frozen in liquid nitrogen, and ground with a pestle and mortar pre-cooled with liquid nitrogen until the starch slurry began to thaw. The mortar was refilled with liquid nitrogen and the starch slurry was ground again. This procedure was repeated once more. The ground starch slurry was transferred to 1.5 ml tubes, centrifuged for 2 min at 5700 g, and the supernatant was discarded. The pooled pellets were resuspended in a total of 3 ml of dH2O, and aliquots of 0.5 ml were added to 0.5 ml of 100 mM sodium acetate (pH 5.5) containing 0 (control) or 89 U of α-amylase (sigmaldrich.com). After incubation at 37 °C for 0 (control) or 2–3 h and centrifugation at 5700 g for 2 min, the supernatant was discarded and 0.5 ml of cold (−20 °C) acetone was added. The samples were allowed to settle for 12 h and to air-dry prior to SEM preparation and observation as above.

**Results**

**Fine mapping \textit{BGC1} in \textit{Aegilops}**

Progeny from the \textit{Ae. peregrina}×KU37 cross were genotyped to identify plants with recombination in the \textit{BGC1} region on chromosome 4S (Howard et al., 2011). Genetic markers were designed to polymorphisms between the two parents, \textit{Ae. peregrina} and KU37, that were identified in RNA-Seq data from grains and leaves (Supplementary Table S1). Selected recombiant plants were allowed to self-fertilize, and homozygous recombinant lines were identified, phenotyped, and genotyped to generate a high-density genetic map across the \textit{BGC1} interval (Fig. 1).

The homozygous recombinant lines were grown and analysed in multiple years and locations to identify lines with different granule size distributions (Fig. 2A). The genetic and phenotypic data mapped \textit{BGC1} between two groups of co-segregating markers, KT113/TC36/TC35 and KT117/KT70 (Fig. 1).

A gene encoding a PTST-family protein is a strong candidate for \textit{BGC1}

We used the \textit{Aegilops} genetic map to establish the syntenic interval in the reference hexaploid bread wheat genome of accession Chinese Spring. Our previous analysis of bread wheat deletion mutants showed that \textit{BGC1} is present in at least two genomes (A and D; Chia et al., 2017). We therefore simplified the synteny analysis by ignoring genes in the region of interest.
that are present in only one genome (Fig. 3). Comparison of gene order between the wheat A, B, and D subgenomes and the Aegilops S subgenome showed only minor differences, with local re-arrangements. From this synteny analysis, we identified 13 genes as candidates for BGC1 (Supplementary Table S4). Examination of their annotations showed that one gene, known in rice as FLOURY ENDOSPERM 6 (FLO6) (Gene 7 in Fig. 3 and Supplementary Table S4), has a known involvement in starch synthesis. In rice and barley, mutations disabling FLO6 cause disruption of starch granule structure (Suh et al., 2017). The FLO6 and PTST2 proteins are homologous to proteins in higher plants called PROTEIN TARGETING TO STARCH (PTST). All members of PTST contain two types of domain: one or more coiled-coil domains (thought to be involved in protein–protein interactions; Mason and Arndt, 2004), and a C-terminal carbohydrate-binding domain of the CBM48 class (Seung et al., 2017).

### Sequencing the BGC1 candidate gene in Aegilops

To test whether the candidate PTST-encoding gene in Aegilops is BGC1, we cloned and sequenced the gene from the mutant Ae. peregrina and the wild-type KU37 (Supplementary Table S2A). Two homologous sequences representing the U and S subgenome homoeologues of BGC1 (BGC1-U and BGC1-S) were expected for each of these tetraploid Aegilops. Partial genomic sequences of the Aegilops genes were obtained and have been deposited in GenBank (accession nos MK848198, MK848199, MK848200, and MK848201). The S- and U-genome sequences were differentiated by the SNP sequence in the S-genome gene that was used to map it to the BGC1 interval (marker TC73; Fig. 1), which was shown previously to lie on chromosome 4S (Howard et al., 2011). Compared with rice FLO6 (Fig. 4; Supplementary Fig. S1), the KU37 and Ae. peregrina U-genome orthologues lack the 5' end of exon 1 that encodes the transit peptide and the first 9–13 amino acids of the mature protein. These two partial sequences vary,particularly in the intron sequences, but they encode identical proteins. We also cloned a truncated S-genome orthologue from KU37 that lacked exons 1–3 and most of the 3'-untranslated region (UTR). However, we were unable to clone any part of the S-genome ortholog from Ae. peregrina except for the 3' end encoding exon 8 and 9, and the 3'-UTR. This raised the possibility that in Ae. peregrina, the S-genome BGC1 candidate gene is almost completely deleted at the 5' end. To explore this further, primers were designed to amplify various regions of the BGC1 candidate genes of Aegilops and wheat (Supplementary Table S2B; Supplementary Fig. S2A). No amplification was seen with primers binding to intron 3, exon 4–5, or intron 5 of the S-genome homoeologue of Ae. peregrina, although products (Supplementary Fig. S2A; P4, P5, and P6) were obtained with these primers for the KU37 S-genome homoeologue. This suggests that the B-granule-less Ae. peregrina may contain a functional U-genome BGC1 candidate gene but that the other homoeologue, on the S-genome, is probably dysfunctional due to severe truncation.

### Testing the BGC1 candidate gene in wheat

Sequencing in Aegilops suggested that BGC1 is an orthologue of the genes that encode PTST in rice and Arabidopsis, FLO6.
BGC1 influences starch granule size distribution in wheat and PTST2, respectively. To test whether mutations in the wheat BGC1 candidate gene result in reduction or elimination of B-type starch granules, we selected TILLING lines of the *T. durum* wheat cultivar, Kronos (a tetraploid with genome composition AABB) with ethylmethane sulfonate (EMS)-induced mutations in either the A-genome (*BGC1*-A) or B-genome (*BGC1*-B) homoeologue (Krasileva et al., 2017; www.wheat-tilling.com; Supplementary Fig. S3). Both of the *BGC1*-A lines (Kronos2244 and Kronos3145) have induced nonsense mutations resulting in premature stop codons. All of the *BGC1*-B lines (Kronos3239, Kronos0456, and Kronos3889) have missense mutations in the region encoding the BGC1 CBM48 domain.

Starch from homozygous single mutant grains was subjected to image analysis to quantify the small-granule content (B-granules plus small A-granules) (Fig. 2B). All of the single mutant lines had a normal small-granule content (Student’s *t*-test; *P >*0.05). One of the *BGC1*-A nonsense lines, K2244, was crossed to each of the three *BGC1*-B missense lines and homozygous double mutant lines (and their corresponding wild-type segregant lines) were selected. One of the double mutants (K2244×K3889) had a normal small-granule content, but the other two (K2244×K3239 and K2244×K0456) had significantly lower (54–59%) small-granule contents than the wild types and the single mutants. When observed microscopic-ally, the grains and extracted starch from these two double mutants had very few B-granules (e.g. Fig. 5A; *T. durum* aabb; K2244×K3239). The small-granule content of these two tetraploid (Kronos) wheat double mutants (Fig. 2B) was very similar to that seen for the hexaploid (Paragon) double-deletion mutant (--BB--; Fig. 2C) and the natural B-granule-less species *Ae. peregrina* (Aep; Fig. 2A) that were described previously (Howard et al., 2011; Chia et al., 2017).
The barley mutant Franubet forms compound starch granules

Despite being closely related taxonomically, the starch phenotype of the barley bgc1 mutant, Franubet, is very different from that of other Triticeae with mutations in BGC1 (tetraploid wheat, hexaploid wheat, and Ae. peregrina), all of which have low or zero B-granule contents. In Franubet barley, both A- and B-type starch granules are largely absent and in their place are starch granules that appear to be fractured or fragmented (DeHaas et al., 1983).

To investigate this apparent discrepancy, we examined the grains and starch of the Franubet mutant of barley (Fig. 5A; H. vulgare lhh). Franubet starch was clearly distinct from that of B-granule-less wheat. The starch granules were highly heterogeneous in morphology, as described previously (DeHaas et al., 1983; Saito et al., 2018; Verhoeven, 2019). There were few, if any, normal A-type or B-type granules in Franubet starch. Some abnormally large A-type granules, irregular or lobed granules, and granules that appear to be compound or fractured were seen.

Compound granules form from multiple, separately initiated granules that become compressed together within a plastid to form polygonal shapes. The presence of compound granules in Franubet in place of single A-type granules would indicate an increase in granule number per plastid rather than a decrease as seen in B-granule-less wheat. We therefore investigated the structure of these starch granules further. Each subgranule in a compound granule has its own set of growth rings (unlike fractured granules that form when a single granule breaks apart late in grain development). Cracking and partly digesting extracted Franubet and Nubet (the wild-type parent of Franubet) starch confirmed that Franubet contains compound starch granules with individual ring structures (Fig. 5B). Furthermore, examination of some of the larger than normal, A-type granules in Franubet showed that, although they appear simple from the outside, when cracked and etched, these granules contain within them separately initiated subgranules. Such granules are called semi-compound granules and have been observed in other plant species such as the bulbs of Scilla ovatifolia Baker (family Hyacinthaceae) (Badenhuizen, 1965). However, as far as we are aware, semi-compound granules have not been observed previously in cereal endosperm.

Generation and analysis of wheat entirely devoid of functional BGC1

The apparent discrepancy between the phenotypes of wheat/Aegilops and barley BGC1 mutants could be due to incomplete elimination of functional BGC1 protein in the former species. Unlike barley, which is diploid, the wheat and Aegilops B-granule-less mutants are polyploid. We hypothesized that not all of the BGC1 homoeologues in these polyploid mutants are completely defective. To investigate this, we crossed the Paragon double-deletion mutant (BGC1 genotype –BB--) to a hexaploid wheat (Cadena) TILLING mutant (Cadenza1730) with a nonsense mutation in the B-genome homoeologue of BGC1 (genotype AABBDD). We selected from the F2 progeny a triple mutant line (genotype --bb--) unable to make any BGC1 protein due to deletion of BGC1-4A and BGC1-4D, and a nonsense (stop) mutation in BGC1-4B.

The starch of this bgc1 triple mutant (--bb--), when viewed microscopically, looked very different from that of the B-granule-less Ae. peregrina, the Paragon double deletion mutant (--BB--), and the Kronos double mutants (aabb). Rather than B-granule-less, the starch resembled that in Franubet barley (Fig. 5A; T. aestivum; --bb--; and Fig. 5C). However, there were more simple granules in the triple mutant wheat
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These could be A-type granules or semi-compound granules. No, or very few, B-type granules were observed. The starch granules in the wheat bgc1 triple mutant were very heterogeneous in shape and size. SEM of grains showed that the granule size and shape varied greatly between cells, and between plastids within the same cell. There were many compound granules, but these varied in the number of constituent subgranules. Some had just a few large subgranules, sometimes arranged in a line or in an irregular shape (these were also observed in Franubet starch). Some of the compound granules were composed of many polygonal subgranules, similar in size to B-granules, and some had many, even smaller, subgranules.

Coulter counter analysis of starch granule size distribution

To further characterize the size distribution of the various wheat starches, we used a Coulter counter. We compared starches from wild-type Paragon (BGC1 genotype AABBDD), B-granule-less Paragon double-deletion mutant (--BB--), and the triple mutant with polymorphous starch (--bb--). The starches in all three genotypes had different starch granule size distributions (Fig. 6A, B). The wild-type starch had a bimodal size distribution (Fig. 6A). By fitting a mixed Gaussian distribution to the data, we deconvoluted the two overlapping peaks of A- and B-type granules, which showed that the B-granule...
content of Paragon starch was $4.2\pm0.8\%$ (v/v). The double deletion mutant starch had a unimodal size distribution as it entirely lacked the B-granule peak. The polymorphous starch in the triple mutant also had a unimodal distribution, but this was wider than that of the B-granule-less starch, and included many small granules of <5 µm in diameter (Fig. 6B).

Discussion

Fine mapping in *Aegilops* revealed a gene underlying the *BGC1* locus controlling B-type starch granule content. The *BGC1* gene is an orthologue of the *FLOURY ENDOSPERM 6* (*FLO6*) genes in rice and barley (Peng et al., 2014; Saito et al., 2018) and the *PTST2* gene in Arabidopsis (Seung et al., 2017) that encode proteins in the PROTEIN TARGETING TO STARCH family. Cloning and sequencing of the candidate gene in *Aegilops* supported the identification of *BGC1*. Whilst a near full-length *BGC1*-S gene that would encode a protein with no obvious defects was found in KU37 which has normal starch, we were unable to amplify a full-length *BGC1*-S from the natural B-granule-less species *Ae. peregrina*. We suggest that the *BGC1*-S gene in *Ae. peregrina* may have been partially deleted, or substantially modified from normal. The candidate gene was further tested by selecting a tetraploid wheat mutant with disrupted *BGC1* genes. The mutant was found to have few, if any, B-granules, like *Ae. peregrina* and a hexaploid bread wheat (Paragon) double mutant with deletions of the *BGC1* regions of chromosomes 4A and 4D (Chia et al., 2017). This result confirms the identification of *BGC1* and shows that *BGC1* controls B-granule content in wheat and *Aegilops*.

All of the B-granule-less wheat and *Aegilops* described here are polyploids, and all are, or could be, partial *BGC1* mutants. First, we found that *Ae. peregrina* has one *BGC1* homoeologue that is probably truncated and therefore dysfunctional and a second apparently normal *BGC1* homoeologue. Secondly, the B-granule-less hexaploid wheat double mutant lacks *BGC1*-A and *BGC1*-D but has *BGC1*-B. Thirdly, the two B-granule-less
tetraploid wheat double mutants each has a BGC1-A nonsense mutation and a BGC1-B missense mutation. We cannot be sure that the two different missense mutations used here eliminate BGC1 functionality: they may just reduce it. One other B-genome BGC1 missense mutation (K3889) when combined with the same A-genome BGC1 nonsense mutation had no impact on starch granule phenotype. We assume therefore that the K3889 BGC1 missense mutation has little or no impact on BGC1 functionality.

The starch phenotypes of the flo6 mutants of rice and barley, which are diploid species, differ from those of the bgc1 mutants of Aegilops and wheat. Rice and barley flo6 mutants have very abnormal granules of diverse size and shape. We have referred to this type of starch as ‘polymorphous’ starch. Unlike the B-granule-less BGC1 partial mutants, these diploids are not B-granule-less. Thus, early in grain development, it has the opposite effect: BGC1 promotes the initiation of many B-type granules per plastid. However, such mutants can have either increased or decreased numbers of initiations depending on the species, and granule morphology can also vary between neighbouring endosperm cells and even between plastids in the same cell. In rice, which normally has compound granules, flo6 mutants have at least some compound granules with more and smaller subgranules than normal, suggesting that in wild-type rice endosperm, FLO6 restricts the number of granule initiations per plastid. In the Triticeae, bgc1 mutants entirely lacking BGC1 have compound and semi-compound granules instead of the single A-type granule per plastid that normally forms early in grain development. Thus, one of the functions of BGC1 in wheat grains may be to limit the number of granule initiations that occur in a plastid so that a single A-type granule can form in the main body of the plastid. However, reduction but not elimination of BGC1 in the Triticeae polyploids results in the loss of B-type starch implying that BGC1 stimulates B-granule initiation in the stromules. Thus, early in grain development in wild-type Triticeae, BGC1 restricts granule initiation to one A-type granule per plastid in but then later in development, it has the opposite effect: BGC1 promotes the initiation of many B-type granules per plastid.

In some growth conditions, wheat is known to accumulate a third type of starch granule, C-type, late in grain development (Bechtel et al., 1990; Zhang et al., 2010). However, in our experiments, we were not been able to distinguish this third granule class. Our starch granule size distributions were bimodal (Fig. 6). Therefore, it remains to be discovered what, if any, impact BGC1 has on the initiation of C-type granules.

**Fig. 6.** Starch granule size distribution in hexaploid wheat. Starches were purified from mature grains of Paragon (AABBDD, wild-type control), the Paragon double deletion mutant (–BB–), and the triple mutant selected from the progeny of a cross between the Paragon double-deletion mutant with a Cadenza TILLING mutant with a nonsense mutation in BGC1-B (AAbbDD). Starches were analysed using a Coulter counter. Values for Paragon and the Paragon double mutant are means for three samples of starch, each from a different grain. Values for the triple mutant are means for six samples of starch each from a different F2 grain. Data are presented in two ways: as % total granule volume (A) and as % total number of granules (B).
At present we cannot explain the apparently contradictory effects of BGC1 on granule initiation at different developmental stages, or the great diversity of starch granule morphology that is observed in the absence of BGC1. The answers may lie in understanding the timing of expression of BGC1 during grain development, the location of BGC1 within plastids, and the nature of its interactions with starch, glucans, and other proteins. Several interaction partners have been identified in Arabidopsis leaves for the BGC1 orthologue PTST2, including starch synthase 4, which is also required for proper starch granule initiation (Roldán et al., 2007; Seung et al., 2017). PTST2 also co-purified in immunoprecipitation experiments with two other plastidial coiled-coil proteins: MRC/PIH1 and MFP1 (Seung et al., 2018). MRC/PIII is a direct interaction partner of SS4, while MFP1 is a thylakoid-associated protein that localizes PTST2 to discrete patches in the chloroplast (Seung et al., 2018; Vandromme et al., 2019). The location of PTST2 in these patches was proposed to restrict granule initiation events to defined areas of the plastid. Wheat has orthologues of all of these proteins, but their role in the endosperm has not been studied. Interestingly, all these proteins appear to promote granule initiation in Arabidopsis leaves, rather than repress it. Future work may determine whether any of these interactions or localization patterns are conserved in amyloplasts of the Triticeae, and other cereals.

We conclude from this work that BGC1 participates in controlling B-type starch granule initiation in Triticeae endosperm but that its precise effect on granule size and number varies with gene dose and stage of development. It is likely that the production of B-granule-less starch involves a delicate balance between the amount of BGC1 and that of one or more of its interacting partners. This may be easier to achieve in a polyploid species than in a diploid.

Data deposition
Partial genomic sequences of the Ae. peregrina BGC1 genes (accession numbers MK848198, MK848199, MK848200, and MK848201). NCBI GenBank. www.ncbi.nlm.nih.gov/genbank/

Supplementary data
Supplementary data are available at JXB online.
Table S1. Ae. peregrina molecular markers.
Table S2. PCR primers for cloning and analysis of BGC1 genes.
Table S3. Triple mutant genotyping primers.
Table S4. Genes in the region containing BGC1.
Fig. S1. Ae. peregrina BGC1 sequence alignment.
Fig. S2. Analysis of BGC1 genes in T. aestivum, Ae. peregrina, and KU37.
Fig. S3. BGC1 TILLING lines of wheat.

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Author contributions
KT, CU, and DS were responsible for the design of the research; the bulk of the practical work was done by TC, BS, and KT, with contributions from MC and TV; JS was responsible for making, growing, and selecting the back-crossed Ae. peregrina lines used for mapping; MT and RRG were responsible for Ae. peregrina RNA-Seq analysis, and together with RK, for wheat bioinformatic analysis; KT drafted the paper with major contributions from CU, DS, and RRG, and minor contributions from TC, BS, and TV. Intermediate authors are listed in alphabetical order.

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