Multiple wheat genomes reveal global variation in modern breeding

Advances in genomics have expedited the improvement of several agriculturally important crops but similar efforts in wheat (*Triticum* spp.) have been more challenging. This is largely owing to the size and complexity of the wheat genome, and the lack of genome-assemble data for multiple wheat lines. Here we generated ten chromosome pseudomolecule and five scaffold assemblies of hexaploid wheat to explore the genomic diversity among wheat lines from global breeding programs. Comparative analysis revealed extensive structural rearrangements, introgressions from wild relatives and differences in gene content resulting from complex breeding histories aimed at improving adaptation to diverse environments, grain yield and quality, and resistance to stresses. We provide examples outlining the utility of these genomes, including a detailed multi-genome-derived nucleotide-binding leucine-rich repeat protein repertoire involved in disease resistance and the characterization of Sm1, a gene associated with insect resistance. These genome assemblies will provide a basis for functional gene discovery and breeding to deliver the next generation of modern wheat cultivars.

Wheat is a staple food across all parts of the world and is one of the most widely grown and consumed crops. As the human population continues to grow, wheat production must increase by more than 50% over current levels by 2050 to meet demand. Efforts to increase wheat production may be aided by comprehensive genomic resources from global breeding programs to identify within-species allelic diversity and determine the best allele combinations to produce superior cultivars.

Two species dominate current global wheat production: allotetraploid (AABB) durum wheat (*Triticum turgidum* ssp. *durum*), which is used to make couscous and pasta, and allohexaploid (AABBDD) bread wheat (*Triticum aestivum*), used for making bread and noodles. A, B and D in these designations correspond to separate subgenomes derived from three ancestral diploid species with similar but distinct genome structure and gene content that diverged between 2.5 and 6 million years ago. The large genome size (16 Gb for bread wheat), high sequence similarity between subgenomes and abundance of repetitive elements (about 85% of the genome) hampered early wheat genome-assembly efforts. However, chromosome-level assemblies have recently become available for both tetraploid and hexaploid wheat. Although these genome assemblies are valuable resources, a list of affiliations appears at the end of the paper.
they do not fully capture within-species genomic variation that can be used for crop improvement, and comparative genome data from multiple individuals is still needed to expedite bread wheat research and breeding. Until now, comparative genomics of multiple bread wheat lines have been limited to exome-capture sequencing\(^1\), low-coverage sequencing\(^2\) and whole-genome scaffolds assemblies\(^3\). Here we report multiple reference-quality genome assemblies and explore genome variation that, owing to past breeder selection, differs greatly between bread wheat lines. These genome assemblies usher a new era for bread wheat and equip researchers and breeders with the tools needed to improve bread wheat and meet future food demands.

Global variation in wheat genomes

To expand on the genome assembly of wheat for Chinese Spring\(^1\), we generated ten reference-quality pseudomolecule assemblies (RQAs) and five scaffold-level assemblies of hexaploid wheat (Supplementary Note 1, Supplementary Tables 1–3). For each RQA, we performed de novo assembly of contigs (contig N50 > 48 kb) that were combined into scaffolds (N50 > 10 Mb) spanning more than 14.2 Gb (Supplementary Note 1). The completeness of the genomes was supported by a universal single-copy orthologue (BUSCO) analysis that identified more than 97% of the expected gene content in each genome (Supplementary Note 1). More than 94% of the scaffolds were ordered, oriented and curated using 10X Genomics linked reads and three-dimensional chromosome conformation capture sequence (Hi-C) to generate 21 pseudomolecules, as done previously for wheat\(^1\) and barley (Hordeum vulgare)\(^5\). The size and structure of the genomes were similar to that of Chinese Spring, and we observed high collinearity between the pseudomolecules (Extended Data Fig. 1). We also independently validated the scaffold placement and orientation in the pseudomolecule assembly of CDC Landmark by Oxford Nanopore long-read sequencing (Extended Data Fig. 2a, Supplementary Note 2). To complement the RQAs, we generated scaffold-level assemblies of five additional bread wheat lines (Supplementary Note 1). To determine the global context of the 15 assemblies, we combined our data with existing datasets\(^6\),\(^7\),\(^8\) (Fig. 1a, Supplementary Table 4). The genetic relationships were in agreement with those reported in previous studies\(^9\),\(^10\) and reflected pedigree, geographical location and growth habit (that is, spring versus winter type). There was also a clear separation between the newly assembled genomes and Chinese Spring, supporting that they capture geographical and historical variation not represented in the Chinese Spring assembly.

Polyploidy and CNV drive gene diversification

Single-nucleotide polymorphisms (SNPs), insertions or deletions (indels), presence/absence variation (PAV) and gene copy number variation (CNV) influence agronomically important traits. This is particularly true for polyploid species such as wheat, in which gene redundancy can buffer the effect of genome variation\(^12\). To assess gene content, we projected around 107,000 high-confidence gene models from Chinese Spring\(^1\) onto the RQAs (Supplementary Note 3). The total number of projected genes exhibited a narrow range, between 118,734 and 120,967 (Supplementary Table 5). We identified orthologous groups among projected genes and used the alignment of the orthologous groups to examine SNPs in coding sequences (Supplementary Note 3). The peak positions of nucleotide diversity across the three subgenomes were highly similar to those reported in previous studies\(^13\),\(^14\), supporting a strong representation of breeding diversity within the RQAs (Extended Data Fig. 3a, b). The correlation of synonymous nucleotide diversity \(\pi\) \((r = 0.11–0.29)\) and Tajima’s \(D\) \((r = 0.02–0.06)\) between homoeologous was low (Supplementary Tables 6–8). This suggested that polyploidization increased the number of targets of selection and contributed to broad adaptation of bread wheat, as in wild polyploid plant species\(^15\),\(^16\).

Further investigation of orthologous groups indicated that 88.1% were unambiguous (clusters containing at most one member in each cultivar) (Extended Data Fig. 3c, Supplementary Table 5). Orthologous groups comprising exactly one gene in each line (‘complete’) were the most frequent (approximately 73.5% of genes per cultivar), suggesting strong retention of orthologous genes within the ten RQAs. The residual genes represented either singleton genes with no reciprocal best BLAST hits or genes located in complex clusters in at least one cultivar. Roughly 12% of genes showed PAVs, and their clustering resulted in relationships (Fig. 1b) that were consistent with SNP-based phylogenetic similarities (Fig. 1a). In addition, approximately 26% of the projected genes were found in tandem duplications, indicating that CNV is a strong contributor of genetic variation in wheat.

To provide an example of gene expansion on emerging breeding targets, we performed a more detailed analysis of the restorer of fertility (Rf) gene families (Supplementary Note 4). Rf genes are involved in restoring pollen fertility in hybrid breeding programs\(^11\), and we identified a previously undescribed clade within the mitochondrial transcription termination factor (mTERF) family (Supplementary Table 9), which has recently been implicated in fertility restoration in barley\(^14\). Of note, this clade shows evolutionary patterns similar to those of Rf-like pentatricopeptide repeat (PPR) proteins, representatives of which are associated with Rf3, a major locus used in hybrid wheat breeding programs (Extended Data Fig. 4). Although wheat is currently not a hybrid crop, there is substantial interest in Rf genes and their potential application in hybrid wheat production systems\(^15\). To our knowledge, no Rf genes have been cloned in wheat and our analysis of Rf genes in multiple RQAs and identification of an Rf clade in wheat is an important step forward in tackling the challenges of hybrid wheat breeding.

The wheat NLR repertoire

To further exemplify the use of multi-genome comparisons for characterizing agricultonically relevant gene families, we examined gene expansion in nucleotide-binding leucine-rich repeat (NLR) proteins, which are major components of the innate immune system and are often causal genes for disease resistance in plants\(^16\),\(^17\),\(^18\). We performed de novo annotation of loci that contain conserved NLR motifs (NB-ARC–leucine-rich repeat) and identified around 2,500 loci with NLR signatures in each RQA (Supplementary Tables 10, 11). A redundancy analysis showed that only 31–34% of the NLR signatures are shared across all genomes, and the number of unique signatures ranged from 22 to 192 per wheat cultivar. We estimated the number of unique NLR signatures that can be detected by incrementally adding more wheat genomes to the dataset; this revealed that 90% of the NLR complement is reached at between 6 (considering 95% sequence identity) and 11 wheat lines (considering 100% protein sequence identity) (Fig. 1c). The total NLR complement of all wheat lines consisted of 5,905 (98% identity) to 7,780 (100% identity) unique NLR signatures, highlighting the size and complexity of the repertoire of receptors involved in disease resistance.

Transposon signatures identify introgressions

Transposable elements make up a large majority of the wheat genome and a critical role in genome structure and gene regulation. We characterized the overall transposable element content (81.6%) and its composition (69% long terminal-repeat retrotransposons (LTR) and 12.5% DNA transposons) in the RQAs (Supplementary Table 5). Across all RQAs, we annotated 1.22 × 10⁶ full-length (fL)-LTRs, which clustered lines into the same groups we observed from our analysis of PAV and SNPs (Fig. 1a, b, Extended Data Fig. 3d). Generally, unique fL-LTRs (147,450) were young (median of 0.9 million years) and were enriched in the highly recombing, more distal chromosomal regions (Fig. 1d). By contrast, shared fL-LTRs were older (median of 1.3 million years) and were more evenly distributed across the pericentric regions (Fig. 1d). The
RLC-Angela fl-LTRs were the most abundant (21,000–27,000 full-length copies per genome) and analysis of variant patterns identified several chromosomal segments that contained numerous unique or rare retrotransposon insertions (Extended Data Fig. 5), which, on the basis of breeding history, we hypothesize to represent introgressions. For example, the LongReach Lancer RQA revealed two unique regions, a pericentric region on chromosome 2B and a segment on the end of chromosome 3D (Extended Data Fig. 5). We used pedigree analysis to postulate the source of the introgressions and performed whole-genome sequencing of multiple accessions of putative donors. LongReach Lancer carries the stem rust resistance gene Sr36, derived from an introgression from *Tritium timopheevii*, and the resistance genes *Lr24* (leaf rust) and *Sr24* (stem rust), derived from tall wheatgrass *Thinopyrum ponticum*. We generated whole-genome sequence reads from multiple accessions (Supplementary Table 1) followed by comparison of chromosomes with the RQAs confirmed that Jagger, Mace, SY Mattis and CDC Stanley carry the 2N′ S introgression, which spans about 33 Mb on chromosome 2A (Fig. 2c, Extended Data Fig. 6a). We annotated the coding genes within this region and identified 535 high-confidence genes; more than 10% were predicted to be associated with disease resistance, including genes that encode putative NB-ARC and NLRs (Extended Data Fig. 6b, Supplementary Tables 17, 18). Furthermore, we used genotyping by sequencing to detect the 2N′ S segment in three wheat panels and discovered that its frequency has been increasing in breeding germplasm and its presence is consistently associated with higher grain yield (Extended Data...
The centromeres are vital for cell division and chromosome pairing during meiosis. In plants, functional centromeres are defined by the epigenetic placement of the modified histone CENH3. We therefore used CENH3 chromatin immunoprecipitation and sequencing (ChIP-seq) to determine the positions and sizes (about 7.5–9.6 Mb) of the centromeres for each RQA (Supplementary Tables 21, 22), which were consistent with previous estimates for wheat. Furthermore, all chromosomes showed a single active site, implying that previous reports of multiple active centromeres in Chinese Spring were artefacts of misoriented scaffolds. However, we found examples in which the relative position of the centromere was shifted owing to several pericentric inversions, including inversions on chromosomes 4B and 5B (Extended Data Fig. 7a, b). We also observed one instance in which the centromeric position changed, but was not associated with a structural event. Specifically, on chromosome 4D in Chinese Spring, the centromere is shifted by around 25 Mb relative to the consensus position (Fig. 2d). This shift was previously recognized by cytology but was hypothesized to result from a pericentric inversion. However, the high degree of collinearity between genomes supports the hypothesis that Cen4D in...
Chinese Spring has shifted to a non-homologous position; this shifting of centromeres to non-homologous sites has also been reported in maize. By characterizing the centromere positions for these diverse wheat lines, we provide strong evidence for changes in centromere position caused by structural rearrangements and centromere shifts.

Large-scale structural variation between genomes

Structural variants are common in wheat, and impact genome structure and gene content. We characterized large structural variants using pairwise genome alignments (Extended Data Fig. 1), changes in three-dimensional topology of chromosomes revealed by Hi-C conformation capture directionalities along the genome, (Extended Data Fig. 8, Supplementary Table 23), which were confirmed by Oxford Nanopore long-read sequencing (Extended Data Fig. 7c, Supplementary Table 24, Supplementary Note 6). The most prominent event was a translocation between chromosomes 5B and 7B, observed in Arina LrFor, SY Mattis (Fig. 2e–g) and Claire. Normally, chromosomes 5B and 7B are approximately 737 and 762 Mb long, respectively, and we estimated that the recombinated chromosomes are 488 Mb (5BS/7BS) and 993 Mb (7BL/5BL) long, making 7BL/5BL the largest wheat chromosome (Extended Data Fig. 9a). In Arina LrFor and SY Mattis, the 7BL/5BL breakpoint resides within an approximately 5-kb GAA microsatellite, which we were able to span using polymerase chain reaction (PCR) (Extended Data Fig. 9b, c). By contrast, the breakpoint on 5BS/7BS was less syntenic, and we detected polymorphic fluorescence in situ hybridization signals between Arina LrFor and SY Mattis on the 5BS portion of the translocated chromosome segment, suggesting that the regions adjacent to the translocation events differ on 5BS/7BS (Supplementary Note 6). To determine the stability of the translocation in breeding, we genotyped for the translocation event in a panel of 538 wheat lines that represent most of the UK wheat gene pool grown since the 1920s. The translocation occurred in 66% of the lines and was selectively neutral (Supplementary Note 7). Notably, the Ph1 locus on chromosome 5B, which controls the pairing of homeologous chromosomes during meiosis, is near the translocation breakpoint, but remained highly syntenic between translocation carriers and non-carriers. Genetic mapping and analysis of short-read sequencing data indicated that the 5B/7B translocated chromosomes recombine freely with 5B and 7B chromosomes (Extended Data Fig. 9d), suggesting that chromosome pairing is not affected by the translocation.

Haplotype-based gene mapping

To develop improved wheat cultivars, breeders shuffle allelic variants by making targeted crosses and exploiting the recombination that occurs during meiosis. These alleles, however, are not inherited independently, but rather as haplotype blocks that often extend across multiple genes that are in genetic linkage. We quantified haplotype variation along chromosomes across the assemblies, and developed visualization software to support its utility (Supplementary Note 8). We used these haplotypes to characterize a locus that provides resistance to the orange wheat blossom midge (OWBM, Sitodiplosis mosellana Gehin), one of the most damaging insect pests of wheat, which is endemic in Europe, North America, west Asia and the Far East. Upon hatching, the first-instar larvae feed on the developing grains and damage the kernels (Fig. 3a). Sm1 is the only gene in wheat known to provide resistance to OWBM. CDC Landmark, Robigus and Paragon are all resistant to the OWBM, and all three carry the same 7.3-Mb haplotype within the Sm1 locus on chromosome 2B (Fig. 3b). To identify Sm1 gene candidates, we used high-resolution genetic mapping and refined the locus to a 587-kb interval in the CDC Landmark RQA (Fig. 3c, Extended Data Fig. 10a, Supplementary Table 25).
Through extensive genotyping of diverse breeding lines, we found an OWB susceptible cultivar, Waskada, that displayed a resistant haplotype except near one gene, which we annotated in CDC Landmark to encode a canonical NLR with kinase and major sperm protein (MSP) integrated domains (Fig. 3c). Oxford Nanopore long-read sequencing further confirmed the structure of the gene in CDC Landmark (Extended Data Fig. 10b). By contrast, the remaining assemblies (susceptible to OWB) lacked the NB-ARC domain, but the kinase and MSP domains remained intact (Fig. 3c). We sequenced the Waskada allele and found it contains the NB-ARC domain, but an alternative haplotype within the kinase domain (Fig. 3c. Extended Data Fig. 10c). This gene is expressed in wheat kernels and seedlings of Sn1 carrier lines, and the lack of cDNA amplification of the NB-ARC domain for non-carrier lines further supported an alternative gene structure (Extended Data Fig. 10c). We generated two knockout-mutant lines of this candidate gene in the Sn1 carrier line Unity41, and both were consistently resistant as susceptible to OWB (Supplementary Table 26). Sequencing of the candidate gene in these two mutants revealed a single point mutation in each line: a G>A mutation resulting in a Gly→Arg (G182R) amino acid substitution in the NB-ARC domain, and a G>A mutation, resulting in a stop codon (W98*) before the NB-ARC domain, encoding sensitivity to the necrotrophic effector ToxA produced by P. nodorum and Pyrenophora triticii-repentis42; however, both Rpq5 and Sn1 lack the MSP domain. To our knowledge, this is the first report of an NB-ARC-LRR-kinase-MSP coding gene associated with insect resistance. Additional research is needed to functionally validate these domains and their putative role in OWB resistance using tools such as gene editing. Nevertheless, we developed a high-throughput and low-cost competitive allele-specific PCR marker (KASP) that discriminates between OWB-susceptible and OWB-resistant lines with perfect accuracy (Extended Data Fig. 10d, Supplementary Table 27). Our analyses, along with the haplotype and synteny viewers (https://kiranjand.github.io/10wheatgenomes/, http://10wheatgenomes.plantinformatics.io/ and http://www.crop-haplotypes.com/), laid the foundation for identifying haplotypes for Sn1. Haplotypes can now be generated in breeding programs using single-marker or high-throughput-sequencing-based approaches, which can integrate desirable genes into improved cultivars more efficiently.

Discussion

We have built on the genome-sequence resources available for wheat and related species to produce ten RQAs and five scaffolded assemblies that represent hexaploid wheat lines from different regions, growth habits and breeding programs. We have identified and characterized SNPs, PAV, CNV, centromere shifts, large-scale structural variants and introgressions from wild relatives of wheat that can be used to identify and characterize important breeding targets. This was complemented by a transposable-element-analysis approach to identify candidate introgressions from wild relatives of wheat, for which we provided high-quality assemblies of segments already used in global breeding programs. Together, these RQAs present an opportunity for breeders and researchers to perform high-resolution manipulation of genomic segments and pave the way to identifying genes responsible for in-demand traits, as we demonstrated for resistance to the insect pest OWB. Functional gene studies will also be facilitated by comparative gene analyses, as exemplified by our analyses of orthologous groups, Rf genes and NLR immune receptors. Finally, we highlight haplotype blocks, which will facilitate marker development for applied breeding. Equipped with multiple layers of data describing variation in wheat, we now have powerful tools to increase the rate of wheat improvement to meet future food demands.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information, details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41586-020-2961-x.
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Methods

No statistical methods were used to predetermine sample size. The field experiments were randomized, but the wheat lines sequenced and assembled were not selected at random. The investigators were not blinded to allocation during experiments and outcome assessment.

Assemblies and annotation

**Genome assemblies.** We assembled the genomes of 15 diverse wheat lines using two approaches (Supplementary Table 1). The RQA approach used the DeNovoMAGIC v.3.0 assembly pipeline, previously used for the wild emmer wheat\(^1\), durum wheat\(^1\) and Chinese Spring RefSeqv1.0 assemblies. In brief, high-molecular-weight DNA was extracted from wheat seedlings as described previously\(^18\). Illumina 450-bp paired-end (PE), 800-bp PE and mate-pair (MP) libraries of three different sizes (3 kb, 6 kb and 9 kb) were generated. Sequencing was performed at the University of Illinois Roy J. Carver Biotechnology Center. 10X Genomics Chromium libraries were prepared and sequenced at the Genome Canada Genome Innovation Centre using the manufacturers’ recommendations to achieve a minimum of 30× coverage. Hi-C libraries were prepared using previously described methods\(^40\). For cultivars assembled to a scaffold level, we used the W2RAP-contigger using k = 200 (Supplementary Note 1). Two MP libraries (10 kb and 13 kb) were produced for each line except Weebill1, for which two additional MP libraries were used. Mate pairs were processed, filtered and used to scaffold contigs as described in the W2RAP pipeline (https://github.com/bioinfologies/w2rap). Scaffolds of less than 500 bp were removed from the final assemblies. Additionally, we performed Oxford Nanopore sequencing of CDC Landmark using R9 flow cells and the GridION sequencing technology (Supplementary Note 2).

**Nucleotide diversity analysis.** The variant call format data files from two wheat exome-capture studies\(^34\) were retrieved, combined, and filtered to retain hexaploid accessions and polymorphisms detected in both studies. The 10X Genomics Chromium sequencing data for each of the RQA lines were aligned to Chinese Spring RefSeqv1.0 using the LongRanger v.2.1.6 software. Alignment files from the assemblies assembled here and 16 Bioplatforms Australia lines\(^39\) with alignments obtained from the DAWN project\(^32\) were then used for variant calling by GATK v.3.8 at the same genomic positions identified by exome-capture sequencing. The variant files from the exome-capture studies, DAWN project and 10X Wheat Genomes lines were then merged and subjected to principal component analysis (PCA) using the prcomp function in R v.3.6.1.

**Gene projections.** We used the previously published high-confidence gene models for Chinese Spring to assess the gene content in each assembly. Representative coding sequences of each informant locus were aligned to pseudomolecules of each line separately using BLAT\(^33\) v.3.5 with the ‘fine’ parameter and a maximal intron size of 70 kb. BLAT matches seeded an additional alignment by exonerate\(^38\) in the genomic neighbourhood encompassing 20 kb upstream and downstream of the match position. Exonerate alignments required a minimal and maximal intron sizes of 30 bp and 20 kb, respectively. A linear regression of colocalized matches with complete alignments of the informant were computed for 10,000 such pairs to derive a normalization function and to render comparable scoring schemes for both methods. Subsequently, we selected the top-scoring match for each mapping pair as the locus for the gene projection. Projections were then filtered by alignment coverage (Supplementary Note 3), the open reading frame (ORF) contiguity, the observed mapping frequency of the informant, coverage of start and stop codons, and the orthology or potential dislocation of the match scaffold relative to its informant chromosome.

Identification of orthologous groups was analogous to the approach used previously\(^5\). Reciprocal best BLAST hit (RBH) graphs were derived from pairwise all-against-all BLASTn v2.8 transcript searches (minimal e-value ≤ 1 × 10\(^{-30}\)). Hits were assigned to homologous groups on the basis of gene models of Chinese Spring following a previously described homeologue classification\(^7\). Multiple sequence alignments for the population genetics analysis were performed using MUSCLE v.3.8 with default parameters (Supplementary Note 3). Using the gene projections, we quantified average pairwise genetic diversity (\(\pi\)), polymorphism (Watterson’s \(\theta_w\), and Tajima’s \(D\) using compute and polydNdS in the ibissequence v.1.0.3-1 package\(^56\). We retained diversity estimates for genes that were in all of the genomes and had ≤100 segregating sites. PAV was determined from the orthologous groups limited to one-to-one relations where there was no match in at least one genome.

**Analysis of the Rf-like gene family.** For Rf genes, the genome sequences were scanned for ORFs in six frame translations with the getorf program of the EMBOSS v.6.6.0 package. ORFs longer than 89 codons were searched for the presence of PPR motifs using hmmsearch from the HMMER v.3.2.1 package (http://hmmmer.org) and the hidden Markov models defined previously. The Pf02356 profile from the Pfam v32.0 database (http://pfam.xfam.org) was used to screen for ORFs carrying mTERF motifs. Downstream processing of the hmmsearch results followed the pipeline described previously\(^57\). ORFs with low hmmsearch scores were removed from the analysis as they are unlikely to represent functional PPR proteins. Only genes encoding mTERF proteins longer than 100 amino acids were included in the analysis. Rf-PPR motifs were identified as described\(^33\). The phylogenetic analyses were performed as described previously\(^33\). Conserved, non-PPR genes delimiting the borders of analysed Rf clusters were identified in the Chinese Spring RefSeqv1.0 reference genome and used to search for syntenic regions in the remaining wheat accessions with BLAST v.2.8. See Supplementary Note 4 for more details.

**NLR repertoire.** NLR signatures were annotated using NLR-Annotator\(^36\) (https://github.com/steuernib/NLR-Annotation) with the option -a. We estimated redundancy of NLR signatures between genomes at different thresholds of identity: 95%, 98% and 100%. For the 165 amino acids in the consensus of all NB-ARC motifs, this translates to 8, 3 and 0 mismatches of a concatenated motif sequence. To calculate the overall redundancy in all genomes, we counted the number of LR signatures added to a non-redundant set by adding genomes iteratively. This was done for 1 million random permutations.

**Repeat annotation.** Transposons were detected and classified by a homology search against the REdat_9.7 Poaceae section of the PGSB transposon library\(^64\) using vmatch (http://www.vmatch.de) with the following parameters: identity ≥ 70%, minimal hit length 75 bp, seedlength 12 bp (exact command line: -d -p -l 75 -identity 70 -seedlength 12 -exdrop 5). To remove overlapping annotations, the output was filtered for redundant hits via a priority-based approach in which higher-scoring matches where assigned first and lower-scoring hits at overlapping positions were either shortened or removed if there was ≥90% overlap with a priority hit or if <50 bp remained. Tandem repeats where identified with TandemRepeatFinder v.4.0.9 under default parameters and subjected to overlap removal as described above. Full-length LTR retrotransposons were identified withLTRharvest (http://genometools.org/documents/ltrharvest.pdf). All candidates were subsequently annotated for Pfam signatures using HMMER v.3.0 and filtered to remove false positives, non-canonical hybrids and gene-containing elements.

The inner domain order served as a criterion for the LTR retrotransposons which are identical upon insertion. The genetic distance was

The phylogenetic analyses were performed as described previously\(^33\). Conserved, non-PPR genes delimiting the borders of analysed Rf clusters were identified in the Chinese Spring RefSeqv1.0 reference genome and used to search for syntenic regions in the remaining wheat accessions with BLAST v.2.8. See Supplementary Note 4 for more details.
calculated with EMBOSS v.6.6.0 distmat (Kimura2-parameter correction) using a random mutation rate of $1.3 \times 10^{-6}$.

**Analysis of centromeric regions.** For each line with a RQA, ChiP was performed according to previous methods\(^a\) with slight modification using a wheat-specific CENH3 antibody.\(^b\) An antigen with the peptide sequence RKHPAVRTKALPK, corresponding to the N terminus of wheat CENH3, was used to produce an antibody using the custom-antibody production facility provided by Thermo Fisher Scientific. The customized antibody was purified and obtained as pellets. The antibody pellet (0.396 mg) was dissolved in 2 ml PBS buffer, pH 7.4, resulting in a working concentration of 198 ng µl\(^-1\). Nuclei were isolated from 2-week-old seedlings, digested with micrococcal nuclease and incubated overnight at 4°C with 3 µg of antibody or rabbit serum (control). Antibodies were captured using Dynabeads Protein G and the chromatin eluted using 100 µl of 1% sodium dodecyl sulfate, 0.1 M NaHCO\(_3\), preheated to 65°C. DNA isolation was then performed using ChiP DNA Clean & Concentrator Kit, and ChiP–seq libraries were constructed using TruSeq ChiP Library Preparation Kit and sequenced with a NovoSeq S4, which generated 150-bp paired-end reads.

For Chinese Spring, we used two datasets, SRR1686799\(^a\) (dataset 1) and the dataset generated in this study (dataset 2). Sequence reads were de-multiplexed, trimmed and aligned to each of the respective RQAs using HISAT2 v.2.1.0\(^a\). Alignments were sorted, filtered for minimum alignment quality of 30, counted in 100-kb bins using samtools 1.10 and BEDTools v.2.29, and visualized in R v.3.6.1. To define the midpoint of each centromere, we identified the highest density of CENH3 ChiP–seq reads using a smoothing spline in R v.3.6.1 with smooth.spline function as the centre of the respective centromere for a given chromosome. To compare centromeric positions of different genomes, the CENH3 ChiP–seq density was plotted along with MUMmer v.4.0 chromosome alignments. To determine the overall size of wheat centromeres, we considered each 100-kb bin with CENH3 ChiP–seq read density that was greater than three times the background (genome average) level of read density to be an active centromeric bin. The number of enriched bins for each genome were counted and averaged to a total of 21 chromosomes. This calculation included counting of unanchored bins.

**Analysis of introgressions**

**Identification of full-length RLC-Angela retrotransposons.** Retrotransposon profiles were created for each genome using the RLC-Angela family\(^a\) and consensus sequences obtained from the TREP database (www.botinst.uzh.ch/en/research/genetics/thomasWicker/trep-db.html). First, BLASTn was used to compare the ~1,700-bp LTR of RLC-Angela to each genome. Matching elements and 500 bp of flanking sequences were aligned to identify precise LTR borders as well as different sub-families and/or sequences variants. We then used BLASTn to compare the 18 consensus LTR sequences against each genome and then screened for pairs of full-lengthLTRs that are found in the same orientation within a window of 7.5–9.5 kb (RLC-Angela elements are ~8.7 kb long). These initial candidate full-length elements were screened for the presence of RLC-Angela polyprotein sequences by BLASTx, as well as for the typical 5-bp target-site duplications. We allowed a maximum of two mismatches between the two target-site duplications. All identified full-length RLC-Angela copies were then aligned to a RLC-Angela consensus sequence with the program Water from the EMBOSS v.6.6.0 package (www.ebi.ac.uk/Tools/emboss/). These alignments were used to compile all nucleotide polymorphisms into a single file. The variant call file was then used for PCA using the snpgdsPCA function in the R package SNPRelate v.3.11.

**Sequencing of the tertiary gene pool of wheat.** Genomic DNA (gDNA) was extracted and purified from young leaf tissue collected from multiple accessions of *T. timopheevii*, *A. ventricosa* and *T. ponticum* (Supplementary Table I2) following a standard CTAB–chloroform extraction method. Yield and integrity were evaluated by fluorometry (Qubit 2.0) and agarose gel electrophoresis. Paired-end libraries were prepared following the Nextera DNA Flex protocol. In brief, 500 ng gDNA from each accession was fragmented and amplified with a limited-cycle PCR. Each library was uniquely dual-indexed with a distinct 10-bp index code (IDT for Illumina Nextera DNA UD) for multiplexing, and quantified by qPCR (Kapa Biosystems). Final average library size was estimated on a Tapestation 2200. Libraries were normalized and pooled for sequencing on an Illumina NovaSeq 6000 S4 to generate ~5× coverage per genotype. Sequencing data were de-multiplexed and aligned to appropriate RQAs (Supplementary Table I2) in semi-perfect mode using the BBMap v.38 short-read alignment software (https://sourceforge.net/projects/bbmap/).

**Structural variation**

We karyotyped the lines using mitotic metaphase chromosomes prepared by the conventional acetocarmine–squash method. Non-denaturing fluorescence in situ hybridization (ND-FISH) of three repetitive sequence probes, Oligo-pSc119.2.1, Oligo-pTa535 and Oligo-pTa713, was performed as described\(^b\) (Supplementary Note 6). Chromosomes were counterstained with DAPI. Chromosome images were captured with an Olympus BX61 epifluorescence microscope and a CCD camera DP80. Images were processed and pseudocoloured with ImageJ v.1.31n in the Fiji package. For karyotyping, at least four chromosomes per accession were examined and compared to the karyotype of Chinese Spring as described previously\(^a\). Hierarchical clustering of karyotype polymorphisms was performed using the Ward method in R v.3.0.2, which was used to estimate distance. Next, we applied Hi-C analysis for inversion calling as described previously\(^a\). In brief, adapters were removed and reads were mapped to Chinese Spring using minimap2 v.2.10\(^b\) as we have done previously\(^a\). The raw Hi-C link counts were calculated in 1 Mb non-overlapping sliding windows and then normalized as described in our previous work\(^a\). Finally, the normalized Hi-C link matrix was subjected to inversion calling using R.

We performed flow cytometry of wheat cultivars Arina and Forno as previously described\(^a\), except that we used a FACSAria SORP flow cytometer and cell sorter (Becton Dickinson). The SB/7B translocation breakpoints were identified by comparison of chromosomes SB and 7B from Arina/LRFor and Julius. Sequence collinearity between Arina/LRFor and Julius was detected by BLASTN searches of 1,000-bp sequence windows every 100 kb along the chromosomes. Once an interruption of synteny was detected, sequence segments at the positions of synteny loss were extracted and used for local alignments to determine the precise breakpoint positions. PCR amplification of the SBS/7BS and 7BL/5BL translocation sites was performed using standard PCR cycling conditions.

**Characterization of haplotypes**

**Development of a wheat genome haplotype database.** To identify haplotypes, pairwise chromosome alignments were performed between the RQA using MUMmer v.4.0, which were combined with pairwise nucleotide BLASTN analyses of the genes ±2,000 bp using custom scripts in R v.3.6.1 (https://github.com/Uauy-Lab/pangenome-haplotypes)\(^a\) (Supplementary Note 8). The resultant haplotypes were uploaded to an interactive viewer (http://www.crop-haplotypes.com/). Pairwise BLASTn comparisons of the genes were also used to identify structural variants, and were uploaded into AccuSyn (https://accusyn.usask.ca/) and SynVisio (https://synvisio.github.io/#/) to create a wheat-specific database (https://kiranandi.github.io/10wheatgenomes/). Pretzel (https://github.com/plantinformatics/pretzel) was also used to visualize and compare the RQA and the projected gene annotations (http://10wheatgenomes.plantinformatics.io/).
Characterization of Sm1. Sm1-linked markers were located in RQAs using BLAST v.2.8.0. Two high-resolution mapping populations were developed, 99B60-EJ2D/Thatcher and 99B60-EJ2G/Infinity. Progeny heterozygous for crossover events near Sm1 were identified in the F2 generation, and the crossovers were fixed in the F3 generation. The resulting F2-derived F3 families were analysed with KASP markers within the Sm1 region and tested for resistance to OWBM in field nurseries to identify markers associated with Sm1. Ethyl methanesulfonate was used to develop knockout mutants in the Sm1 gene. Approximately 3,200 seeds of the Canadian spring wheat variety Unity (an Sm1 carrier) were soaked in a 0.2% (v/v) aqueous ethyl methanesulfonate solution for 22 h at 22 °C. The seed was then rinsed in distilled water and sown in a field nursery. The M1 seed was grown to maturity and bulk harvested. Approximately 6,000 M2 seeds were space planted in two field nurseries located in Brandon and Glenlea, Manitoba, Canada. Spikes were collected on a per-plant basis at maturity and were classified as resistant, susceptible or undamaged as done previously42,49. Putative Sm1 knockout mutants were re-tested for OWBM resistance in indoor cage tests23 in the M2 and M3 generations. M3-derived families were tested for resistance to OWBM in field nurseries (randomized complete block design, six environments, and eight replicates per environment).

Candidate genes were identified between Sm1 flanking markers on the CDC Landmark assembly using the projected gene annotations and FGENESH v.2.6 (http://www.softberry.com/), which were compared to the projected genes of non-carriers. Both 5′ and 3′ rapid amplification of cDNA ends (5′ and 3′ RACE) were used to verify the transcription initiation and termination sites of the gene candidate, whose structure was predicted by FGENESH v.2.6. In brief, RNA was extracted from the leaves of Unity (Sm1 carrier) seedlings (using the Qiagen RNeasy kit), RACE PCR performed (Invitrogen GeneRacer kit), and the PCR product cloned (Invitrogen TOPO TA Cloning kit for sequencing) and sequenced by Sangen sequencing. Prediction of the conserved domains was done using the NCBI Conserved Domain Search tool (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) and PROSITE (release 2020_01; https://prosite.expasy.org/). The LRR domain was defined on the basis of the presence of 2–42 LRR motif repeats of 20–30 amino acids each. LRR motifs were manually annotated23. Prediction of transmembrane regions and orientation was performed using the program TMPred NCBI Conserved Domain Search tool (https://embre.teenitol-it.ch/software/TMPREDForm.html).

To study the expression of Sm1, total RNA was extracted from four biological replicates from two wheat genotypes (Unity, CDC Landmark, Waskada and Thatcher) from two different tissues; seedling leaves and developing kernels (five days post anthesis) using NucleoSpin RNA Plant kit (Macherey-Nagel) according to the manufacturer’s instructions. RNA was treated with RNase-free DNase (RNase) (Macherey-Nagel) and reversed transcribed into cDNA using SuperScript IV Reverse Transcriptase kit (Invitrogen) according to the manufacturer’s instructions and the NB-ARC domain amplified by PCR.

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data availability
All sequence reads assemblies have been deposited into the National Center for Biotechnology Information Sequence read archive (SRA) (see Supplementary Table I for accession numbers). Sequence reads for the RQAs, T. ponticum, A. ventricosa and T. timopheevii have been deposited into the SRA (accession no. PRJNA544491) and ChIP-seq short read-data used for centromere characterization is deposited under accession no. PRJNA625537. All Hi-C data have been deposited in the European Nucleotide Archive (Supplementary Table I). The RQAs are available for direct user download at https://wheat.ipk-gatersleben.de/. All assemblies and projected annotations are available for comparative analysis at Ensembl Plants (https://plants.ensembl.org/index.html). Comparative analysis viewers are also online for synteny (https://kiranbandi.github.io/10wheatgenomes/, http://10wheatgenomes.plantinformatics.io/) and haplotypes (http://www.crop-haplotypes.com/). Seed stocks of the assembled lines are available at the UK Germplasm Resources Unit (https://www.seedstor.ac.uk/).

Code availability
Code for custom genome visualizers have been deposited in the public domain for haplotype viewer (https://github.com/Uayu-Lab/pangenome-haplotypes), Pretzel (https://github.com/plantinformatics/prezel), AccuSyn (https://github.com/jorgenunesziri/accusyn) and SynVisio (https://github.com/kiranbandi/synvisio). Additional scripts used for ChIP-seq analysis of the centromeres are provided at https://github.com/wheatgenetics/centromere.

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Extended Data Fig. 1 | Chromosome-scale collinearity between the RQA. Genomes were aligned chromosome by chromosome using MUMmer and are represented as dot plots. The introgression on chromosome 2B of LongReach Lancer (red rectangles) and 5B/7B translocation in SY Mattis and ArinaLrFor (purple rectangles) are indicated.
Extended Data Fig. 2 | Evaluation of the CDC Landmark RQA using Oxford Nanopore Long Reads. a, Scaffold-scaffold long read contact map showing shared read IDs between scaffold ends along the ordered scaffolds in the CDC Landmark pseudomolecules. The diagonal pattern indicates that adjacent scaffolds share the same long reads and are therefore properly ordered and oriented by Hi-C in the RQA. b, Characterization of inversion events on chromosomes 2A, 3A, and 3D. The directionality biases estimated from alignments of Hi-C data against Chinese Spring (left, top), and chromosome alignment of the inversion events between CDC Landmark and Chinese Spring RQAs (left, bottom) are shown. Long reads spanning the inversion events and magnified views of the reads aligning to the left and right boundaries of the inversions (right) are provided.
Extended Data Fig. 3 | Diversity of genes and TEs. 

a, Average pairwise genetic diversity of the homeologues (coding sequences only) of the A, B and D subgenomes. The mode of the A, B and D subgenome is 0.00057, 0.00082, and 0.0002, respectively. 

b, Tajima’s D estimates of coding sequences for each wheat subgenome. The lower and upper range of the boxplot hinges correspond to the first and third quartiles (the 25th and 75th percentiles). Boxplots show centre line, median; box limits, upper and lower quartiles; whiskers, 1.5 × interquartile range.

c, Total gene counts and orthologues for the RQA. Genes in orthologous groups with exactly one gene for each line (Complete; dark brown), genes contained in unambiguous orthologous groups missing an orthologue for at least one line, that is, PAV (2-10 Lines; light brown), and genes with ambiguous orthologues or CNV (Other; pink) are indicated.

d, Per cent of pairwise shared syntenic fl-LTRs between wheat lines.
Extended Data Fig. 4 | Evolutionary relationships among PPR and mTERF gene sequences. a, The RFL clade is in blue and all remaining P-class PPRs are in green. b, Clustered mTERF sequences are in blue and the remaining mTERFs are shown in green. The scale bar represents number of substitutions per site. c, Sequence inversions and copy number variation at the Rf3 locus on chromosome 1B. RFL genes are shown as light pink triangles above the chromosome scale. Conserved non-PPR genes used as syntenic anchors are shown on the chromosome scale as coloured triangles. The total number (T) and the number of putatively functional RFL genes with 10 or more PPR motifs (F) are indicated on the right side of each panel.
Extended Data Fig. 5 | Identification of alien introgressions from wheat relatives. A feature of foreign chromosomal introgressions is that they contain unique patterns of TE insertions. Shown are stretches of >20 Mb containing multiple polymorphic RLC-Angela retrotransposons that are found only in one or a few (≤4) of the sequenced lines. One representative chromosome for each wheat subgenome is shown. Individual polymorphic retrotransposons are indicated as coloured vertical lines. Colours correspond to the number of cultivars a foreign segment is found in. Regions of particular interest are indicated by black rectangles. These include the 2N'S alien introgression from *A. ventricosa* at the end of chromosome 2A in Jagger, Mace, SY Mattis and CDC Stanley, as well as introgression in the central region of chromosome 2B from *T. timopheevi* in LongReach Lancer, and introgression at the end of chromosome 3D from *T. ponticum* in LongReach Lancer.
Extended Data Fig. 6 | Detailed characterization of the 2NvS introgression from *A. ventricosa*. 

a, Pairwise alignments of the first 50 Mb of chromosome 2A. The black arrow indicates a possible unique haplotype within spelt. 

b, Orthologous genes between the 2NvS introgression from *A. ventricosa* in Jagger and the genes on chromosomes 2A, 2B, and 2D in Chinese Spring. 

c, Frequency of 2NvS introgression carriers in North American datasets from CIMMYT, Kansas State, and the USDA Winter Wheat Regional Performance Nursery (RPN) over time. 

d, Percent yield difference in lines that carry the 2NvS introgression. Two sided t-tests were performed to test for the significance of the impact of the 2NvS introgression. **P < 0.01; ***P < 0.001.
Extended Data Fig. 7 | Centromere positions and karyotype variation.
Functional centromere positions in the RQA have undergone structural and positional rearrangement. Chromosome alignments showing collinearity (black scaffolds in same orientation, grey scaffolds in opposite orientation) with relative density of CENH3 ChIP–seq mapped to 100 kb genomic bins for Chinese Spring (blue) and a representative genome of comparison (red) for chromosome 4B of CDC Stanley (a), and chromosome 5B of Julius (b). c, Detailed list and clustering of cytological features carried by each wheat line (Supplementary Note 6). Features that are identical (dark grey) or have a gain (black) or loss (light grey) relative to Chinese Spring are indicated.
Extended Data Fig. 8 | Hi-C validates inversions identified from pairwise chromosome alignments. Pairwise alignments of chromosome 6B from the RQA and Chinese Spring are shown. Above each alignment dot plot, the directionality biases estimated from alignments of Hi-C data against Chinese Spring are shown. Boundaries of diagonal segments are indicative of inversions and coincide with inversion boundaries identified from the chromosome alignments.
Extended Data Fig. 9 | Characterization of a translocation involving小麦 chromosomes 5B and 7B. a, Cytogenetic karyotypes of Forno (left) and Arina (right), the parents of ArinaLrFor. Note that the large recombinant chromosome 7B is represented by a distinct peak. b, Sequence of the translocation breakpoint on chromosome 7B of ArinaLrFor. Note that the exact breakpoint lies in a sequence gap (stretch of Ns). The bp positions are indicated at the left. Forward PCR primers are shown in red and reverse primers in blue. The overlap of the two reverse primers is shown in purple. The outer primer pair was used for PCR, while the inner pair was used for a nested PCR. c, PCR amplification of the fragment spanning the translocation breakpoint. The nested PCR yielded a ~5 kb fragment that spanned the translocation breakpoint and its identity was confirmed by sequencing. Both PCR and nested PCR were performed in duplicate; both replicates of the nested PCR were sequenced using the Sanger method. For gel source data, see Supplementary Fig. 1. d, Mapping of Illumina reads from the cultivars Arina and Forno on to the pseudomolecules of ArinaLrFor. Sequence derived from Forno is shown in blue, while sequenced derived from Arina is in red. Note that chromosomes 5B and 7B are derived from both parents, indicating that these parental chromosomes can recombine freely, despite the presence of a large 5B/7B translocation in Arina.
Extended Data Fig. 10 | See next page for caption.
Extended Data Fig. 10 | Confirmation of gene expression and gene structure for Sm1. a, Critical recombinants from the 99B60-EJ2G/Infinity and 99B60-EJ2D/Thatcher populations used to fine map Sm1. The 99B60-EJ2G/Infinity cross had 5,170 F2 plants, while 99B60-EJ2D/Thatcher cross had 5,264 F2 plants; only recombinant haplotypes between orange wheat blossom midge resistant (R) and susceptible (S) genotypes are shown. b, Oxford Nanopore long read confirmation of the Sm1 gene candidate in the CDC Landmark RQA (left), and alternative haplotype in Chinese Spring (right). Vertical coloured lines indicate sequence variants. c, Amplification of cDNA for the NB-ARC domain of the Sm1 gene candidate (top) and actin control (bottom) derived from RNA isolated from developing kernels (left) and wheat seedlings (right). Unity and CDC Landmark are carriers of Sm1. Waskada carries an alternative haplotype and does not carry Sm1 (see main text). Thatcher was used as a susceptible parent for fine mapping of Sm1 and does not contain the associated NB-ARC domain. The experiment was replicated on four independent biological samples for each condition. d, Distribution of an Sm1 allele-specific PCR marker in a diverse panel of >300 wheat lines.
Reporting Summary

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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

No software was used to collect data for this study.

Data analysis

A multitude of software and databases were used in this study, all of which have been listed, cited, or provided. These include: DeNovoMAGIC v3.0, W2RAP (no versions, https://github.com/bioinfologics/w2rap), LongRanger v2.1.6, GATK v3.8, R v3.6.1 and v3.0.2, BLAST v3.5, BLAST v2.8, MUSCLE v3.8, libsequence v1.8.3, EMBoss v6.6.0, HMMER 3.1b2, PFAM v32.0, NLR-annotator (no version, https://github.com/steuernb/NLR-Annotator), Vmatch v2.3.0, TandemRepeatFinder v4.07b, LTRharvest genometools-1.5.9, HMMER v3.0, MUMmer v3.23 (haplotype database) and v4 (all other analyses), HISAT v2.1.0, SNPrelate v3.11, BBTools/BBMap v38, ImageJ v1.51n, minimap2 v2.13, FGENESH v2.6, NCBI Conserved Domain Search tool (no version, https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb), PROSITE release 2020_01, TMPred v25, STAR v2.6.0b., AUGUSTUS v3.2.3., GMAP v2017-06-20, EvidenceModeler v1.1.1, AHRD v1.6, MCScanX v2.0, samtools v1.10, BEDtools v2.29, and custom data scripts (https://github.com/Uauy-Lab/pangenome-haplotypes; http://people.beocat.ksu.edu/~jpoland/centromeres/).

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

All sequence reads have been deposited into the National Center for Biotechnology Information sequence read archive (SRA) [see Supplementary Table 1 for accession numbers]. Sequence reads for the RQAs, Th. ponticum, Ae. ventricosa and T. timopheevii have been deposited into the SRA (no. PRJNA544491) and ChiP-
Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

- Life sciences
- Behavioural & social sciences
- Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size
No statistical methods were used to establish sample size. The samples that were sequenced were selected to represent modern breeding material from different continents that had known differences in pedigree and were known to carry different genes/traits/chromosomal segments of interest.

Data exclusions
All sequencing data generated was used in the genome assembly and analyses. Whenever possible, all data was included in the supporting analyses. Data exclusion applies only to some of the subsequent supporting analysis, which was pre-established based on limitations in the data. For example, we excluded the scaffolded assemblies from some analyses because the analyses required chromosome pseudomolecules. We performed diversity analysis both with the spelt genome but also excluding the spelt genome because it is a different species and is much more diverged and biased the results.

Replication
In all analyses that support the genome assemblies, the number of replicates or iterations are indicated in materials and methods or supplemental tables. In each case, all replications were successful and were used. The genome assemblies themselves were validated using multiple methods (i.e. BUSCO, genetic maps, HiC, 10x Genomics, cytology, and comparisons to Chinese Spring). The CDC Landmark assembly was further validated using Oxford Nanopore long read sequencing. This helped validate the other approaches.

Randomization
Randomization does not directly apply to the genome sequencing and assembly; however it applies to some of the supporting analyses. In these cases, the group design and data seeding for computational analysis are described in the materials and methods and adhere to widely accepted standards. For example, analysis of NLRs (Fig. 1c), 1 million random permutations were used. For the field experiments established for phenotyping of Sm1, all samples were replicated and randomized using appropriate experimental designs.

Blinding
Blinding does not apply to this study, as the study focuses on genome sequencing. This study focuses on plants genomics and the results of the study are not impacted by the concealment of treatment, data, or groups.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| □   | Antibodies            |
| □   | Eukaryotic cell lines |
| □   | Palaeontology         |
| □   | Animals and other organisms |
| □   | Human research participants |
| □   | Clinical data         |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| □   | ChIP-seq              |
| □   | Flow cytometry        |
| □   | MRI-based neuroimaging |

Antibodies

Antibodies used
Chromatin immunoprecipitation (ChIP) was performed using wheat CentH3 antibody (Koo et al., 2015). A antigen with the peptide sequence ‘RTKHPAVRTKALPKK’ corresponding to the N-terminus of wheat CENH3 was used to produce antibody utilizing the custom-antibody production facility provided by the Thermo Fisher Scientific, Illinois, USA (abs@thermofisher.com). A 0.396 mg of the antibody pellet was dissolved in 2 ml of PBS buffer, pH 7.4 resulting in 198 ng/ul of the working concentration.

Validation
In the manuscript, we validate the antibody according to a previous study of Chinese Spring (Koo et al., 2015) and achieved near
ChIP-seq

Data deposition

- Confirm that both raw and final processed data have been deposited in a public database such as GEO.
- Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

| Data access links | The data for the project has been deposited at NCBI: PRJNA625537 and analysis files are available for download: http://people.beocat.ksu.edu/~jpoland/centromeres/ |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------|
| Files in database submission | BED files, delta files (MUMmer), data analysis scripts |
| Genome browser session (e.g. UCSC) | Data for visualization is available at http://people.beocat.ksu.edu/~jpoland/centromeres/ |

Methodology

| Replicates | NA. Samples were obtained from 2-week-old seedlings. |
|------------|---------------------------------------------------|
| Sequencing depth | Paired-end reads were generated at varying levels of read depth, data was deposited at NCBI (PRJNA625537). |
| Antibodies | Wheat CenH3 antibody - see: Koo DH, Sehgal SK, Friebe B, Gill BS (2015) Structure and stability of telocentric chromosomes in wheat. PLoS One 10: e0137747. |
| Peak calling parameters | Reads mapped per 100kb bin were counted for each sample using BEDtools and output as a bed file. Scripts for data analysis are provided at http://people.beocat.ksu.edu/~jpoland/centromeres/. Unlike studies involving transcription factors, CENH3 ChIP-seq provides clear distinct peaks that are ~100 fold greater than background. |
| Data quality | SAM output files from HISAT2 were converted to BAM, sorted and filtered for minimum alignment quality of 30 using SAMtools. |
| Software | Reads for each sample were aligned to each of the respective genome assemblies using HISAT2. Reads mapped per 100kb bin were counted for each sample using BEDtools and output as a bed file. Scripts for data analysis are provided at http://people.beocat.ksu.edu/~jpoland/centromeres/ |