

Durum wheat genome highlights past domestication signatures and future improvement targets

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The domestication of wild emmer wheat led to the selection of modern durum wheat, grown mainly for pasta production. We describe the 10.45 gigabase (Gb) assembly of the genome of durum wheat cultivar Svevo. The assembly enabled genome-wide genetic diversity analyses revealing the changes imposed by thousands of years of empirical selection and breeding. Regions exhibiting strong signatures of genetic divergence associated with domestication and breeding were widespread in the genome with several major diversity losses in the pericentromeric regions. A locus on chromosome 5B carries a gene encoding a metal transporter (*TdHMA3-B1*) with a non-functional variant causing high accumulation of cadmium in grain. The high-cadmium allele, widespread among durum cultivars but undetected in wild emmer accessions, increased in frequency from domesticated emmer to modern durum wheat. The rapid cloning of *TdHMA3-B1* rescues a wild beneficial allele and demonstrates the practical use of the Svevo genome for wheat improvement.

Durum wheat (DW), *Triticum turgidum* L. ssp. *durum* (Desf.) Husn., genome BBAA, is a cereal grain mainly used for pasta production and evolved from domesticated emmer wheat

(DEW), *T. turgidum* ssp. *dicoccum* (Schrank ex Schübl.) Thell. DEW itself derived from wild emmer wheat (WEW), *T. turgidum* ssp. *dicoccoides* (Körn. ex Asch. & Graebn.) Thell., in the Fertile Crescent

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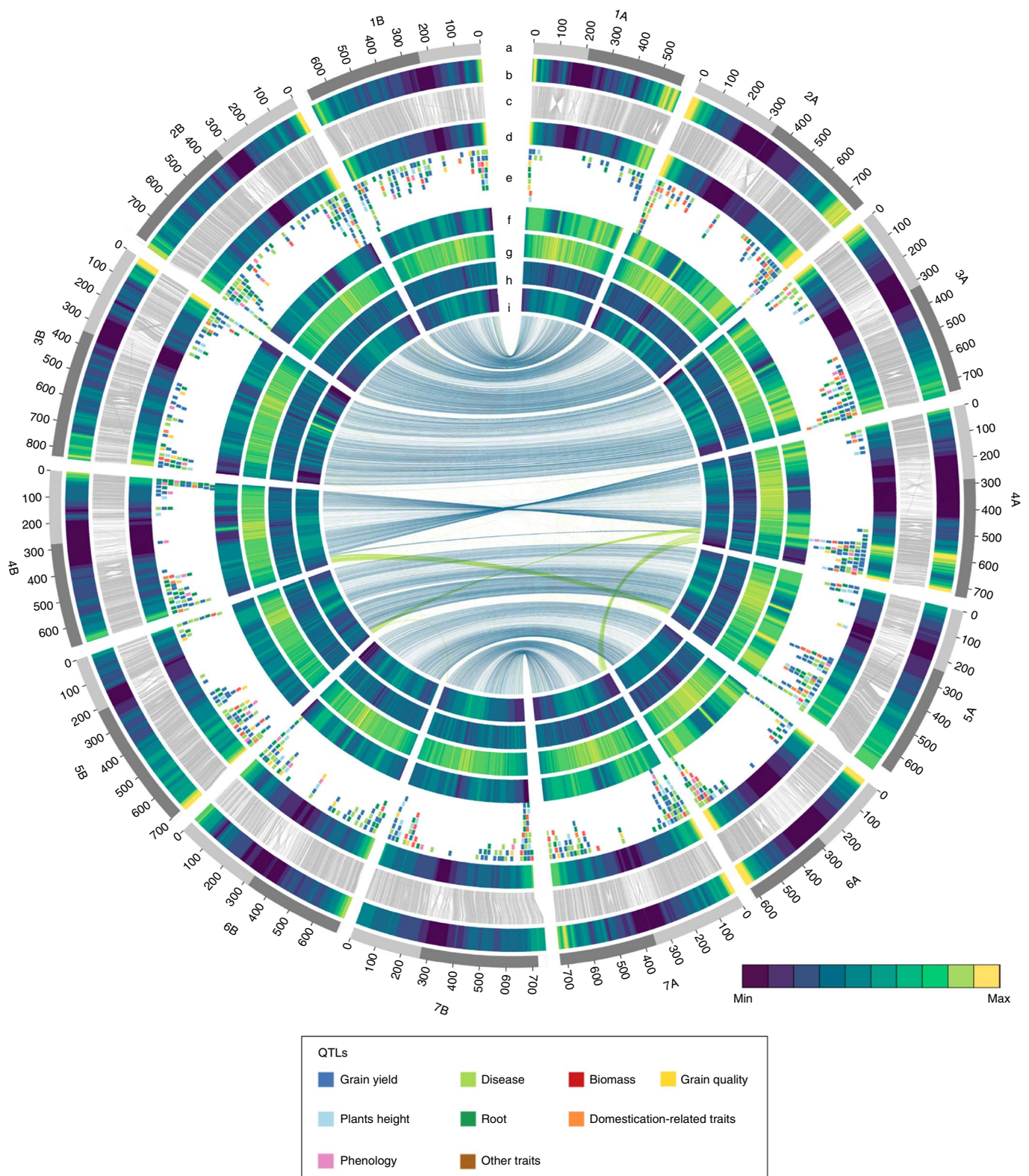


Fig. 1 | Structural, functional and conserved synteny landscape of the DW genome. Tracks from outside to inside. **a**, Chromosome name and size (100 Mb tick size, arms differentiated by gray shading). **b**, Density of WEW HC gene models (HC; 0–25 genes per Mb). **c**, Links connecting homologous genes between WEW and DW. **d**, Density of DW HC gene models (0–22 genes per Mb). **e**, Location of published QTLs. **f**, *k*-mer frequencies. **g**, Long terminal repeat (LTR)-retrotransposon density. **h**, DNA transposon frequency. **i**, Mean expression of HC genes calculated as $\log(\text{FPKM} + 1)$ of the mean expression value of all conditions (range 1.6–8.2). Links in center connect homoeologous genes between subgenomes; blue links between homoeologous chromosomes and green links between large translocated regions.

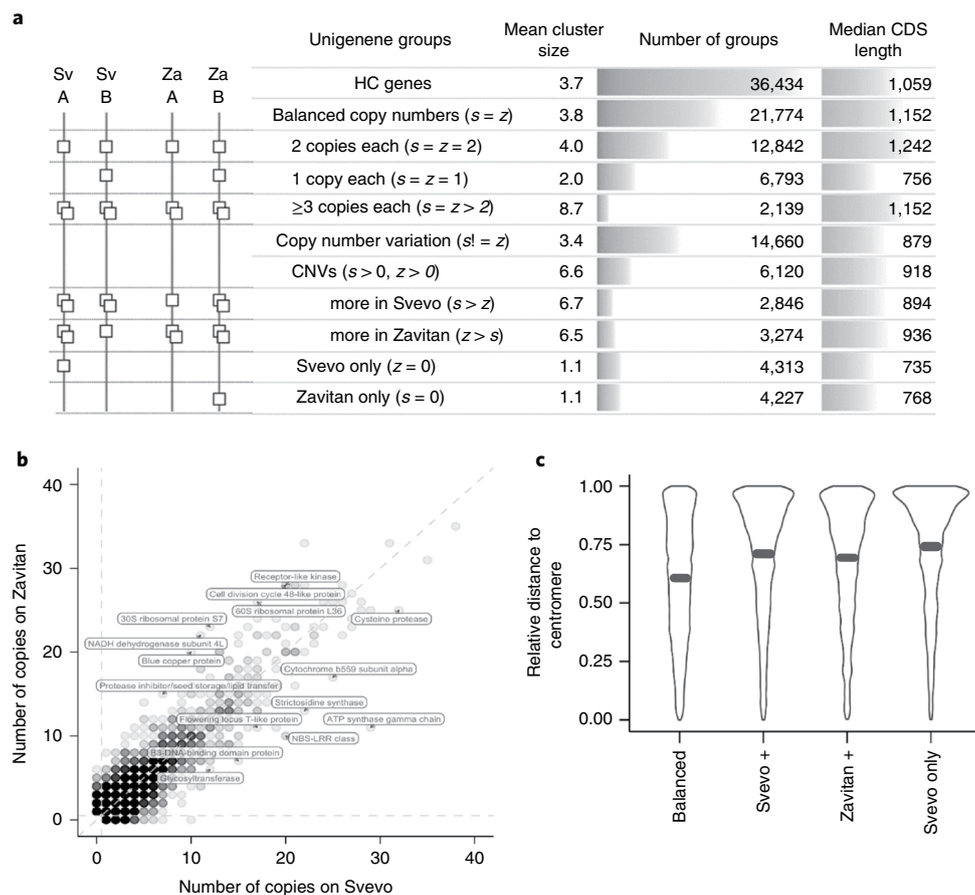


Fig. 2 | Comparison of the Svevo and Zavitan gene space. a, Main unigene group scenarios from co-clustering of Svevo and Zavitan HC genes. The diagram depicts the most common or typical scenarios, only HC intact genes were considered (CNV, copy number variation; CDS, coding sequence). **b**, Intact gene number variations. Each dot represents a gene cluster consisting of DW (x axis) and WEW (y axis) genes. Dots on the diagonal represent clusters with identical member numbers from both accessions. Functional predictions for some groups of genes with pronounced differences in member numbers are annotated on the diagram. **c**, Relative distance of DW genes from the centromere separated by gene cluster type. Proportionately more unigenes displaying intact gene number variation than balanced groups are observed towards the ends of the chromosome. HC genes unique to Svevo or Zavitan (not shown) are most highly represented at the ends of the chromosomes, with the median (black line) furthest from the centromere. Shape width represents the relative gene frequency.

about 10,000 years ago¹. Although the first evidence of DW dates to 6,500–7,500 years ago, DW became established as a prominent crop only 1,500–2,000 years ago². Thus, the human-driven tetraploid wheat evolution process is the result of domestication (WEW to DEW), continued evolution under domestication (DEW to durum wheat landraces, DWL) and breeding improvement from DWL to modern durum wheat cultivars (DWC).

Wild relatives of modern crop plants can serve as sources of valuable genetic diversity for various traits (for example, disease resistance^{3,4} and nutritional quality⁵). Comprehensive comparative genomic analyses between cultivated crops and wild progenitors is a key strategy to detect novel beneficial alleles and structural variations that could constrain breeding efforts, as well as to understand the broader genetic consequences of evolution and selection history^{6,7}.

Here we report the fully assembled genome of the modern DW cultivar (cv.) Svevo and provide a genome-wide account of modifications imposed by thousands of years of empirical selection and breeding. This was achieved by comparing the Svevo genome with the assembled genome of WEW accession Zavitan⁸ and through a survey of the genetic diversity and selection signatures in a Global Tetraploid Wheat Collection consisting of 1,856 accessions. A region bearing a signature of historic selection

co-locates with *Cdu-B1*, a quantitative trait locus (QTL) spanning 0.7 cM on chromosome 5B⁹ known to control cadmium (Cd) accumulation in the grain. Identification of the gene(s) responsible for *Cdu-B1* has been hampered by the large and repetitive nature of the DW genome and the low recombination rate in the region of interest. The efficient, genome-enabled dissection of the *Cdu-B1* locus reported here demonstrates the value of the Svevo genome assembly for wheat improvement.

Results

The durum wheat reference genome. The Svevo genome sequence was assembled de novo using protocols previously described⁸ and its main features are illustrated in Fig. 1. After sequencing (Supplementary Table 1) and assembly, the scaffolds (length of the shortest contig needed to cover 50% of the genome (N50) = 6.0 megabases (Mb); Supplementary Table 2) were ordered and oriented using the Svevo×Zavitan genetic map as previously described¹⁰. Thereafter, chromosome conformation capture sequencing (Hi-C)¹¹ resulted in a set of pseudomolecules (9.96 Gb; Supplementary Table 3) corresponding to the 14 chromosomes of DW and one group of unassigned scaffolds (499 Mb). The pseudomolecules encompass 95.3% of the assembled sequences and have 90% of the scaffolds oriented. Alignment of the DW genome with

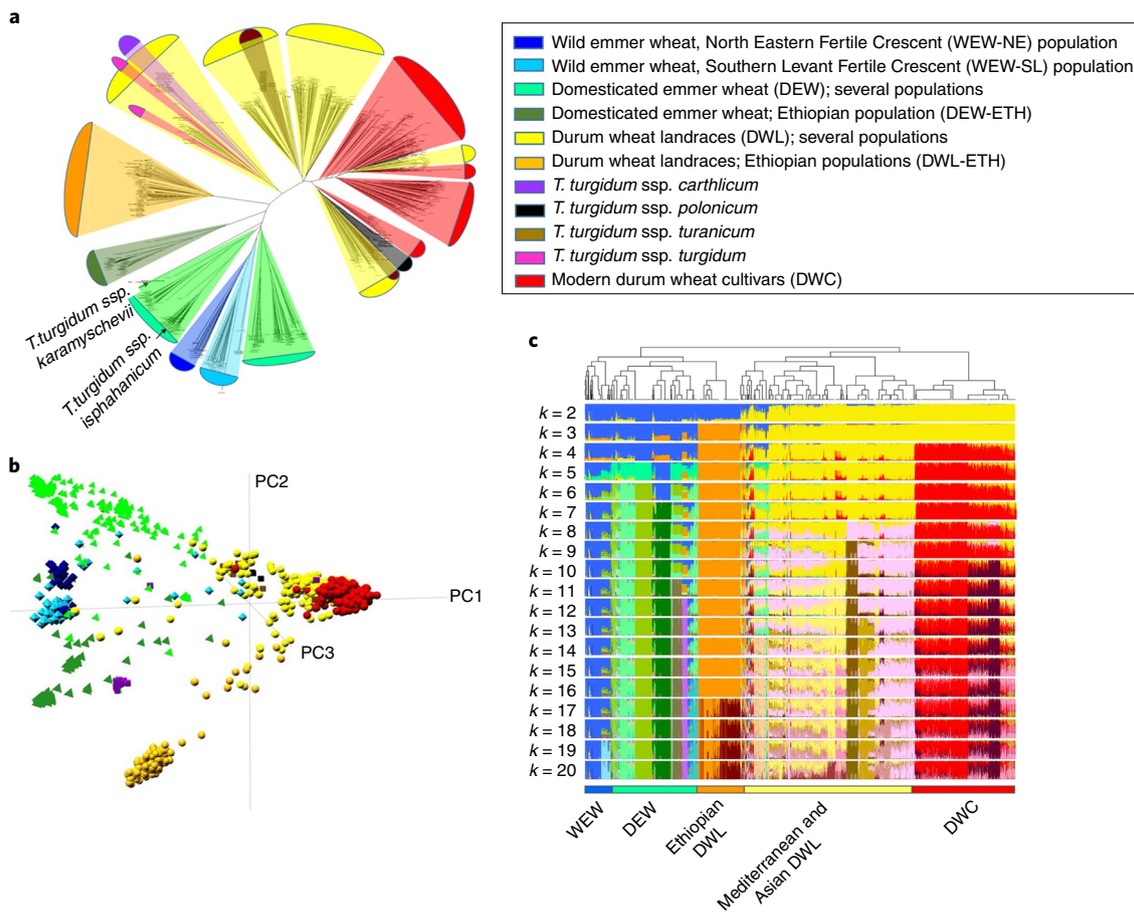


Fig. 3 | Tetraploid germplasm structure and phylogenetic relationships. **a**, Neighbor joining tree from Nei's genetic distances among the 1,856 accessions of the Global Tetraploid Wheat Collection. Genetic distances were computed from a set of 5,775 whole-genome linkage disequilibrium-pruned ($r^2 = 0.5$) SNPs. Correspondence between branches and main tetraploid wheat taxa/populations on the basis of ADMIXTURE and other population structure analyses are indicated by color code with the exception of *T. turgidum* ssp. *karamyschevii* and ssp. *isphanicum* that are indicated directly on the graph. Significances were estimated through 1,000 bootstrap resampling. **b**, Principal component analysis plot of the Global Tetraploid Wheat Collection on the basis of genome-wide pairwise distances calculated on the basis of linkage disequilibrium-pruned SNPs. **c**, ADMIXTURE analyses of the Global Tetraploid Wheat Collection with k (number of populations assumed for the analysis) from 2 to 20.

high-density SNP genetic maps¹² showed highly recombinogenic distal chromosome regions exhibiting an almost linear relationship between genetic and physical distance (Supplementary Fig. 1). These regions account for about 22% of the genome with an average recombination rate of 1.8 Mb cM^{-1} (Supplementary Table 4). In contrast, large pericentromeric regions are nearly devoid of recombination and represent about 44% of the genome, with a mean recombination rate of 107 Mb cM^{-1} . Annotation of the Svevo genome led to the identification of 66,559 high confidence (HC) genes, 90.5% of which exhibited detectable evidence of expression in at least one of the 21 RNA-seq datasets listed in Supplementary Table 5. A detailed description of the DW genome is presented in the Supplementary Note (Sections 1.1 and 2.1). Projection onto the DW genome of 2,191 previously reported QTLs resulted in a full meta-QTL analysis (Supplementary Table 6b and Supplementary Dataset 1), revealing a QTL density distribution that closely mirrors the gene density distribution (Supplementary Table 7 and Fig. 1d,e).

Comparison between Svevo and Zavitan genomes. To gain insights into short-term evolutionary changes, we compared the genome divergence between the modern DW cultivar Svevo and the WEW accession Zavitan⁸. The comparison revealed strong overall synteny (Fig. 1c) with high similarity in total HC gene number

(DW 66,559; WEW 67,182; Supplementary Table 8), chromosome structure and transposable element composition (Supplementary Table 9). We identified syntenic LTR-retrotransposon insertions (Supplementary Fig. 2) not yet subjected to the rapid transposable element turnover of the intergenic space^{1,13} because of the relatively short separation time between Svevo and Zavitan. To monitor structural variations in the HC gene set, including minor changes that might have been generated during the short evolutionary timespan, a graph-based sequence clustering of all Svevo and Zavitan HC genes (in total 133,741) was undertaken. Stringent clustering (alignment e value $< 10^{-10}$, overlap $> 75\%$ and identity $> 75\%$) grouped only highly similar gene models in the same cluster. This approach produced 36,434 unigene groups, 79% (28,794) of which were clusters with at least two members, while 21% (7,640) contained only singletons (Supplementary Table 10). The main scenarios for conserved and variable genes are summarized in Fig. 2a. The most frequent cluster configuration is made of two homoeologous gene copies per genome (one per A and B sub-genomes), which occurs in 35% of all unigene groups. Altogether, the unigene groups with balanced copy numbers for Svevo and Zavitan represent up to 60% of all unigenes and involve 63% of all Svevo genes. The remaining 40% of unigenes (14,660) display asymmetric numbers of intact, full-length genes between Svevo

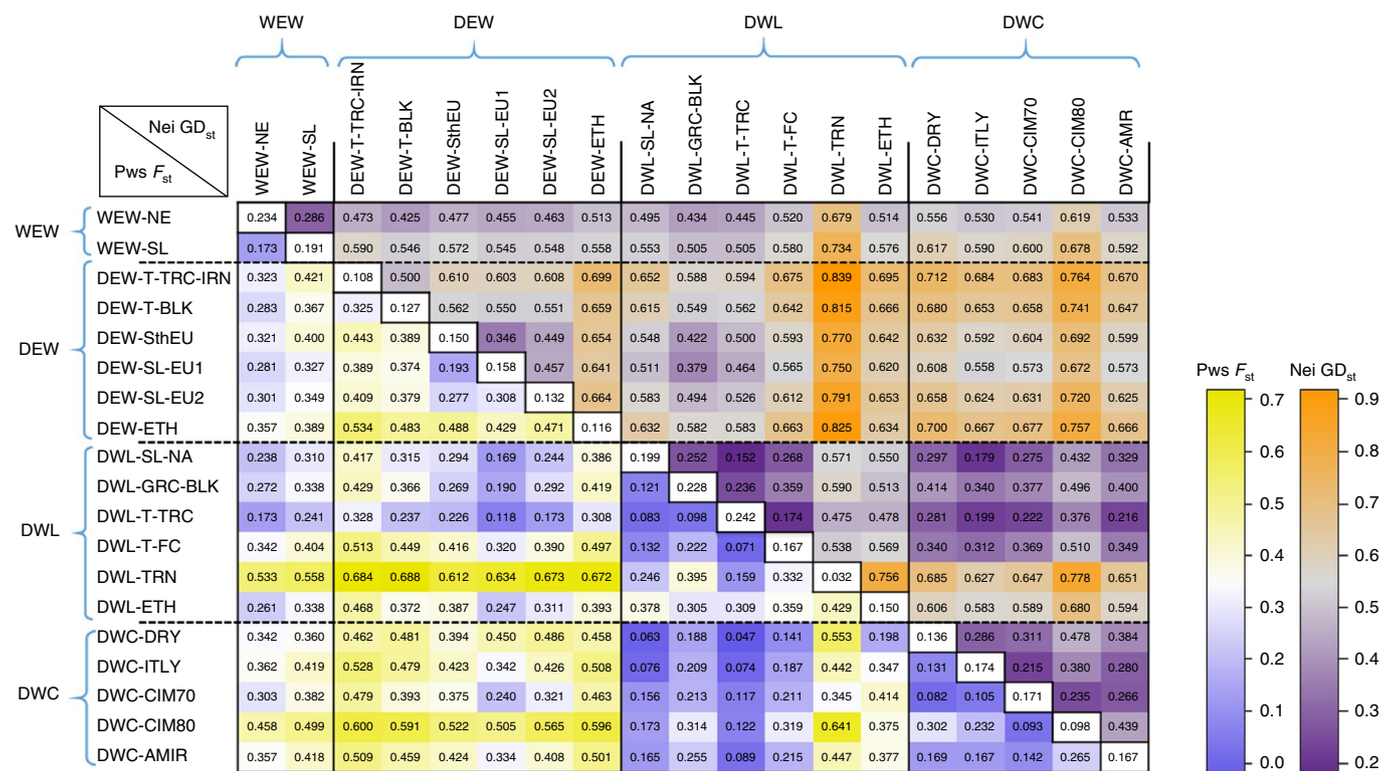


Fig. 4 | Summary of Nei's genetic distances GD_{st} (above diagonal) and pairwise F_{st} (below diagonal) between main tetraploid wheat populations. Diagonal numbers represent within-population genetic diversity (expected heterozygosity) values. Only low-admixture accessions were used (Q-membership higher than 0.5 for WEW, DEW; Q-membership higher than 0.4 for DWL, DWC). Statistics were estimated with 5,775 linkage disequilibrium-pruned ($r^2=0.5$) SNPs. WEW-NE, WEW from the North Eastern Fertile Crescent, Turkey, Iran and Iraq; WEW-SL, WEW from Southern Levant including Lebanon, Syria, Israel and Jordan; DEW-T-TRC-IRN, DEW from Turkey to Transcaucasia and Iran; DEW-T-BLK, DEW from Turkey to the Balkans; DEW-SthEU, DEW spread in Southern Mediterranean areas; DEW-SL-EU1, DEW from Southern Levant Fertile Crescent to Europe (population 1); DEW-SL-EU2, DEW from Southern Levant Fertile Crescent to Europe (population 2); DEW-ETH, DEW from Oman, India and Ethiopia; DWL-SL-NA, DWL from Southern Levant Fertile Crescent to North Africa and Iberia; DWL-GRC-BLK, DWL from Greece to Balkans; DWL-T-TRC, DWL from Turkey to Transcaucasia; DWL-T-FC, DWL diffused in Turkey to the whole Fertile Crescent; DWL-TRN, *T. turanicum*; DWL-ETH, DWL from Ethiopia; DWC-DRY, DWC from Italian and ICARDA breeding programs adapted to dryland areas; DWC-ITLY, DWC from Italy; DWC-CIM70, DWC from the wide adaptation, temperate-adapted photoperiod insensitive CIMMYT and ICARDA germplasm bred in the 1970s; DWC-CIM80, DWC from the high-yielding CIMMYT germplasm bred in the 1980s; DWC-AMR, DWC from the photoperiod-sensitive North American and French germplasm.

and Zavitan (we named this intact gene number variation). Since the unigene classification is on the basis of HC genes, any mutation leading to a frameshift and/or premature stop codon that rules out a gene from the HC class in Svevo or in Zavitan, results in an asymmetric unigene distribution. For at least two-thirds of the unigenes displaying variation of intact gene number, counterparts for the missing copies can still be found in LC or pseudogene class. The complete gene loss caused by large structural variations was responsible for asymmetric gene distribution in only one-third of the cases. Among the unbalanced gene clusters, there are 6,120 mixed clusters with copies from both genomes, subdivided into more Svevo members or more Zavitan members, as well as 4,313 and 4,227 lineage-specific unigene groups (mostly singleton genes) in Svevo and Zavitan, respectively, which have no close homoeolog in the HC gene set of the other accession. A detailed example of the type of variation leading to intact gene number variation is given for the lineage-specific unigenes (Supplementary Table 10a). In Svevo, this class includes 4,811 genes that represent 7.2% of all HC genes, a value similar to the 5% found after the comparison between two cultivars in a recent pangenome study of hexaploid wheat¹⁴. When the Svevo-specific genes were mapped onto the Zavitan genome (Fig. 2a and Supplementary Table 10b), 1,493 genes (31%) were not found on the Zavitan sequence, 1,225 (26%) correspond to shorter counterparts of annotated Zavitan

HC genes and 1,095 (23%) were annotated as low confidence (LC) genes or pseudogenes. The remaining 965 (20%, that is, 1.4% of all Svevo HC genes) map to unannotated regions and are candidates for genes missed in the automated annotation.

Loss and gain events can occur from ancestral four-member unigene clusters with one gene per A and B subgenome of both Svevo and Zavitan. One loss would result in a three-member unigene cluster, with one subgenome location missing. A total of 1,121 clusters with one lost Svevo member were found. The reverse situation with one lost member from Zavitan was found 852 times. The presumed HC gene losses are located predominantly in the more distal chromosomal regions (Supplementary Fig. 3a). A gain event would result in clusters with at least five members with one subgenome carrying two members. This condition was found 472 times with Svevo gains and 503 times with Zavitan gains. Most of the gains are located on the same chromosome indicating tandem gene duplication as the prevailing mechanism (Supplementary Fig. 3b).

Functional categories associated with relevant copy number differences between the two accessions are highlighted in Fig. 2b. A statistical analysis for gene ontology over-representation revealed that specific functions can be classified as: differentially enriched in Svevo, differentially enriched in Zavitan, balanced and Svevo-specific unigene groups (Supplementary Fig. 3c). The balanced

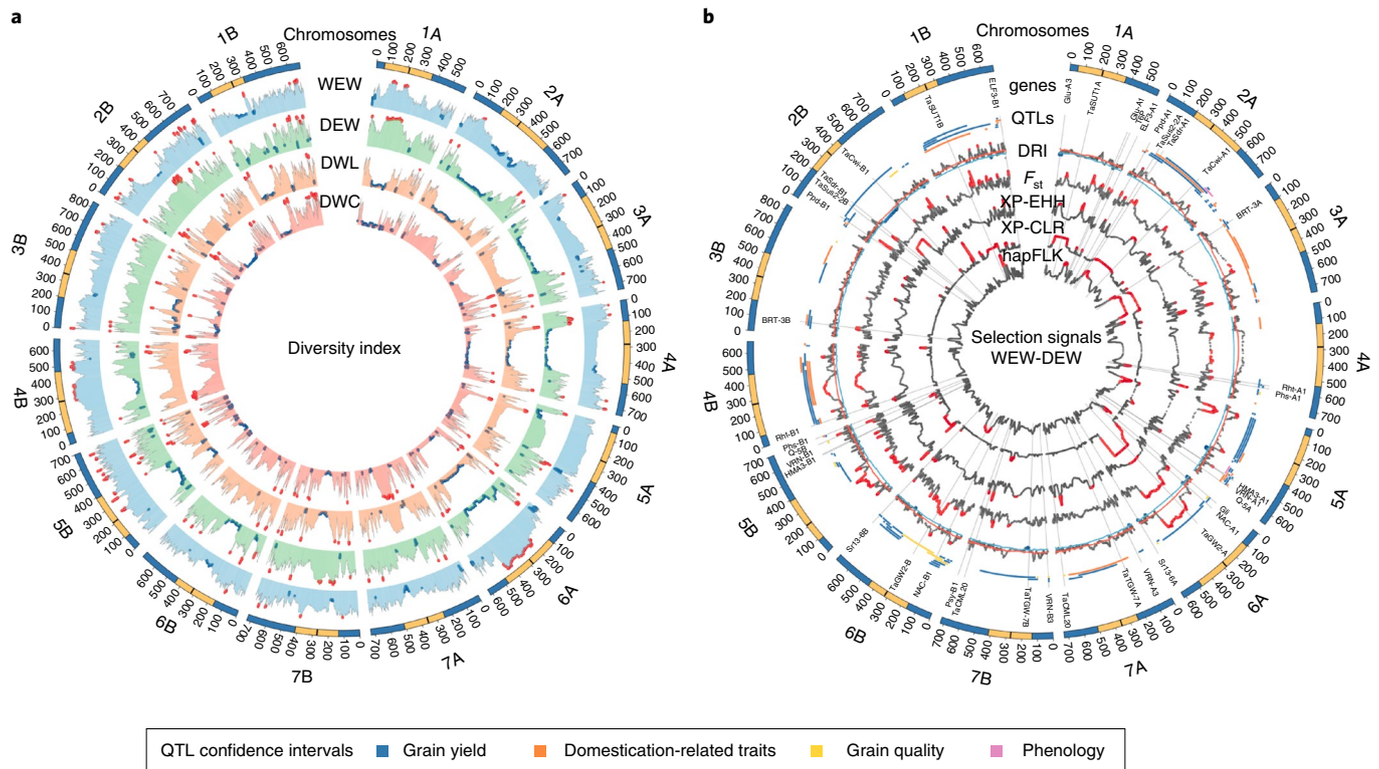


Fig. 5 | Genome-wide analysis of SNP diversity in the Global Tetraploid Wheat Collection and cross-population selection signatures from wild to domesticated emmer transition (WEW to DEW) on the basis of 17,340 informative SNPs. a, SNP-based diversity index (DI) for the main germplasm groups identified in the Global Tetraploid Wheat Collection: WEW, DEW, DWL and DWC. DI is reported as a centered 25 SNP-based average sliding window (single SNP step). Top and bottom 2.5% DI quantile distributions are highlighted as red- and blue-filled dots, respectively. **b**, Cross-population selection index metrics for the comparison between WEW and DEW. Selection metrics are provided for: diversity reduction index (DRI), divergence index (F_{st}), cross-population extended haplotype homozygosity (XP-EHH), multilocus test for allele frequency differentiation (XP-CLR) and haplotype-based differentiation test (hapFLK). For DRI, top and bottom 2.5% DI quantile distributions are highlighted as red- and blue-filled dots, respectively, while for the other selection metrics top 5% quantile distributions are highlighted as red-filled dots. The physical location of genes (Supplementary Table 12) and QTL confidence intervals relevant to domestication and breeding is reported.

gene groups are enriched for a multitude of regulatory functions and indicate a stronger sensitivity to gene dosage effects for the main regulatory networks. Unbalanced unigene groups with more Svevo members and Svevo unique groups are enriched for functions involving protein phosphorylation, for example kinases, which are known to trigger signal transduction cascades in response to environmental cues. The distal highly recombinogenic regions of chromosomes are enriched in unigenes displaying variation of intact gene number (Fig. 2c), contain most of the known QTLs (Fig. 1e) and the HC genes display a reduced expression breadth (that is, average expression value across all tissue/treatment conditions, Fig. 1i). This indicates the presence of an increased number of condition-specific genes. The less dynamic interstitial regions contain a greater number of balanced gene families (Fig. 2c). Here the genes are expressed in nearly all conditions, indicative of an enrichment in housekeeping genes that is consistent with reports from the barley¹¹ and bread wheat¹⁵ genomes. The positive correlation between recombination rate and DNA variants supports previous evidence that higher recombination rates and illegitimate recombination are drivers for tandem duplications¹⁶.

The balanced copy number groups contain much longer genes (median 1,152 base pairs (bp)) than the groups displaying variation of intact gene number (median 879 bp; Fig. 2a). The highest median gene lengths are found in groups with two copies in each genome (1,242 bp), whereas the lowest are among the unique genes (Svevo, 735 bp; Zavitan, 768 bp) and, surprisingly, in groups with one copy in each genome (756 bp). Such a pronounced shift towards shorter

genes indicates an ongoing gene decay by frameshifts and mutations leading to premature stop codons. Collectively, the relatively high number of genes undergoing degeneration could be a consequence of more freedom for gene loss facilitated by the functional redundancy of the tetraploid genome state.

Germplasm structure and phylogenetic relationships. The wheat iSelect 90K SNP Infinium assay¹⁷ was used to genotype the Global Tetraploid Wheat Collection consisting of 1,856 accessions representing the four main germplasm groups involved in tetraploid wheat domestication history and breeding: WEW, DEW, DWL and DWC (Supplementary Table 11). A set of 17,340 SNPs (non-redundant, genetically and physically mapped, sub-genome-specific Mendelian loci) was used for analysis of genetic diversity, population structure and identification of the selection signatures. Four non-hierarchical clustering analyses (DAPC, sNMF, ADMIXTURE, fineSTRUCTURE^{18–21}), principal component analysis, and pairwise dissimilarity analysis on the basis of neighbor joining generated global and highly concordant pictures of the genetic relationships among taxa and populations (Fig. 3 and Supplementary Datasets 2 and 3). WEW, DEW, DWL and DWC clearly separated in the neighbor joining tree (Fig. 3a), a result indicative of strong demographic and founder effects and little evidences for polyphyletic origin. Principal component analysis (Fig. 3b) illustrates the broad genetic diversity of DEW, while DWC showed a comparatively limited genetic diversity and a close relationship to a specific DWL population. DEW and

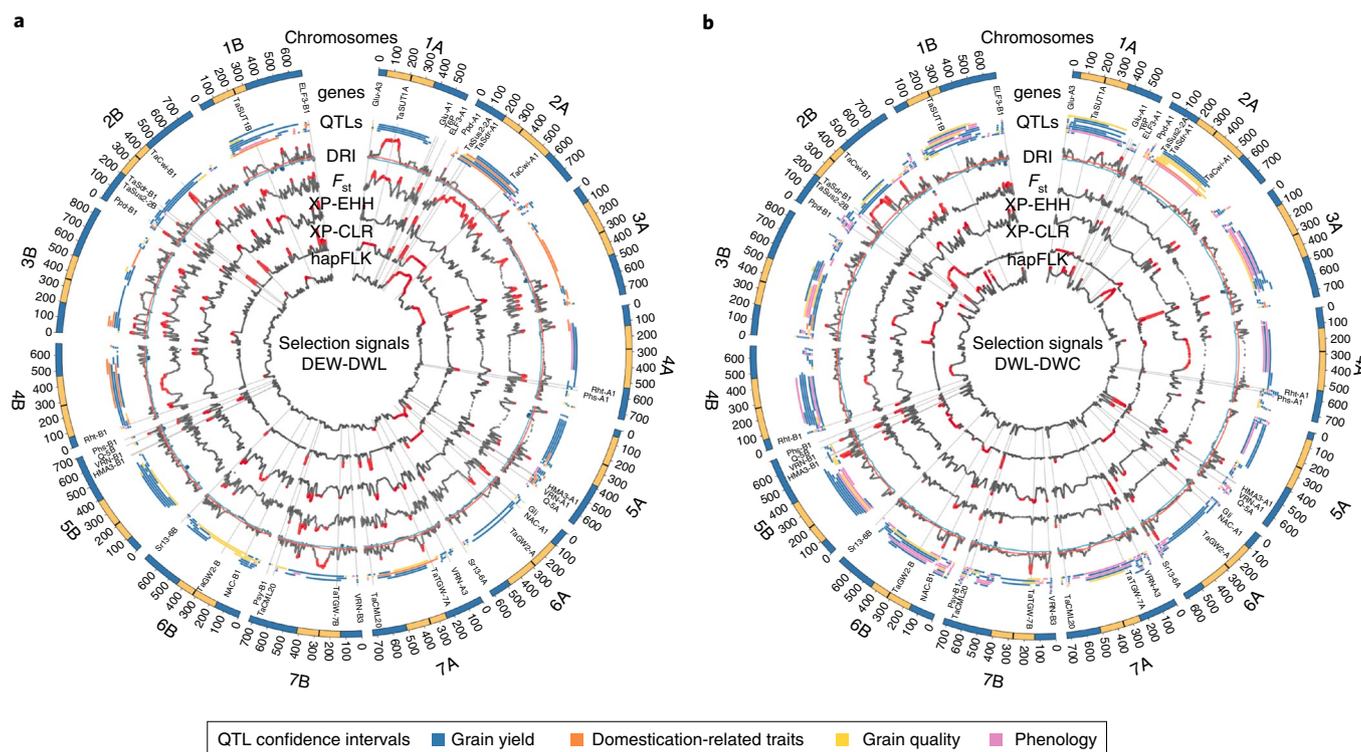


Fig. 6 | Analysis of diversity and selection signatures in tetraploid wheat. Genome-wide cross-population selection signatures in DEW to DWL and DWL to DWC on the basis of 17,340 informative SNPs. **a**, Cross-population selection index metrics for the DEW to DWL. **b**, Cross-population selection index metrics for the DWL to DWC. For both panels, selection metrics are provided for: DRI, F_{st} , XP-EHH, XP-CLR and hapFLK. For DRI, top and bottom 2.5% DI quantile distributions are highlighted as red- and blue-filled dots, respectively, while for the other selection metrics top 5% quantile distributions are highlighted as red-filled dots. The physical location of genes (Supplementary Table 12) and QTL confidence intervals relevant to domestication and breeding is reported.

DWL from Ethiopia, *T. turgidum* ssp. *turanicum* and *T. turgidum* ssp. *carthlicum* were genetically isolated.

ADMIXTURE analysis (Fig. 3c) showed that WEW and DEW have highly structured genetic diversity even at high k values, while DWL showed a high rate of admixture already at low k values. WEW germplasm was divided into two main populations from North Eastern Fertile Crescent and Southern Levant (WEW-NE and WEW-SL, respectively). WEW-NE was further divided into several populations from Turkey, Iran and Iraq, while WEW-SL included distinct populations from Israel (3), Jordan, Syria and Lebanon (Supplementary Fig. 4). DEW and DWL germplasm was characterized by a similar though independent radial dispersal pattern: Northern-to-Southern Fertile Crescent and from Fertile Crescent to Mediterranean basin (Western), Greece to Balkans (Western), Iran to Transcaucasia (Eastern) and Oman to India and Ethiopia. The germplasm belonging to DEW and DWL was subdivided in six main populations each, while all DWC clustered to a further distinct group that represents a wide branch of the durum North African and Turkey to Transcaucasian landrace populations (Supplementary Fig. 4).

After removing the accessions with a high level of admixture, the genetic relationships among the main tetraploid germplasm groups were further investigated using hierarchical analysis of variance (ANOVA), by computing the pairwise divergence index (or fixation index) F_{st} and Nei's genetic distances (Fig. 4), and by generating population-based whole-genome phylogenetic trees (Supplementary Fig. 5). The results confirm the radial dispersal patterns already reported and indicate the WEW-NE from Turkey as the most probable ancestor of all DEW populations (F_{st} and genetic distance values consistently lower for all WEW-DEW pairs). Two DEW populations

from Southern Levant Fertile Crescent (Fig. 4) showed the closest relationship to all DWL populations (except *T. turgidum* ssp. *turanicum*), while the DWC germplasm was mostly related to the two DWL populations from North Africa and Transcaucasia (Fig. 4). The Ethiopian and *T. turgidum* ssp. *turanicum* populations were the most differentiated among the DWL germplasm and their contribution to the modern durum varieties was minimal.

Diversity reduction and signature of selection. The pattern of diversity for each germplasm group was assessed through a SNP-based gene diversity index²² (Fig. 5a). WEW showed the highest average diversity with only two pericentromeric regions (chromosomes 2A and 4A) with a lower than average diversity. Thus, WEW provides a valuable reference for assessing the reduction of diversity associated with domestication and breeding in tetraploid wheat. Compared to WEW, each of the subsequently domesticated/improved germplasm group showed several strong diversity depletions that arose independently and were progressively consolidated through domestication and breeding. With few exceptions, the diversity depletions that occurred in the early transition (WEW to DEW, Fig. 5b or DEW to DWL, Fig. 6a) are confirmed or even reinforced in the subsequent ones (Fig. 6a,b). Consequently, the genome of DWC is characterized by numerous regions showing near-fixation of allelic diversity (Fig. 6b). We applied five different metrics to detect selection signatures: diversity reduction index (DRI²³), single site divergence index (F_{st} ; ref. ²⁴), haplotype-based frequency differentiation index (hapFLK²⁵), cross-population extended haplotype homozygosity (XP-EHH²⁶), and spatial pattern of site frequency spectrum (XP-CLR²⁷). Genomic regions supported by one or more indexes were considered as putative signatures of selection.

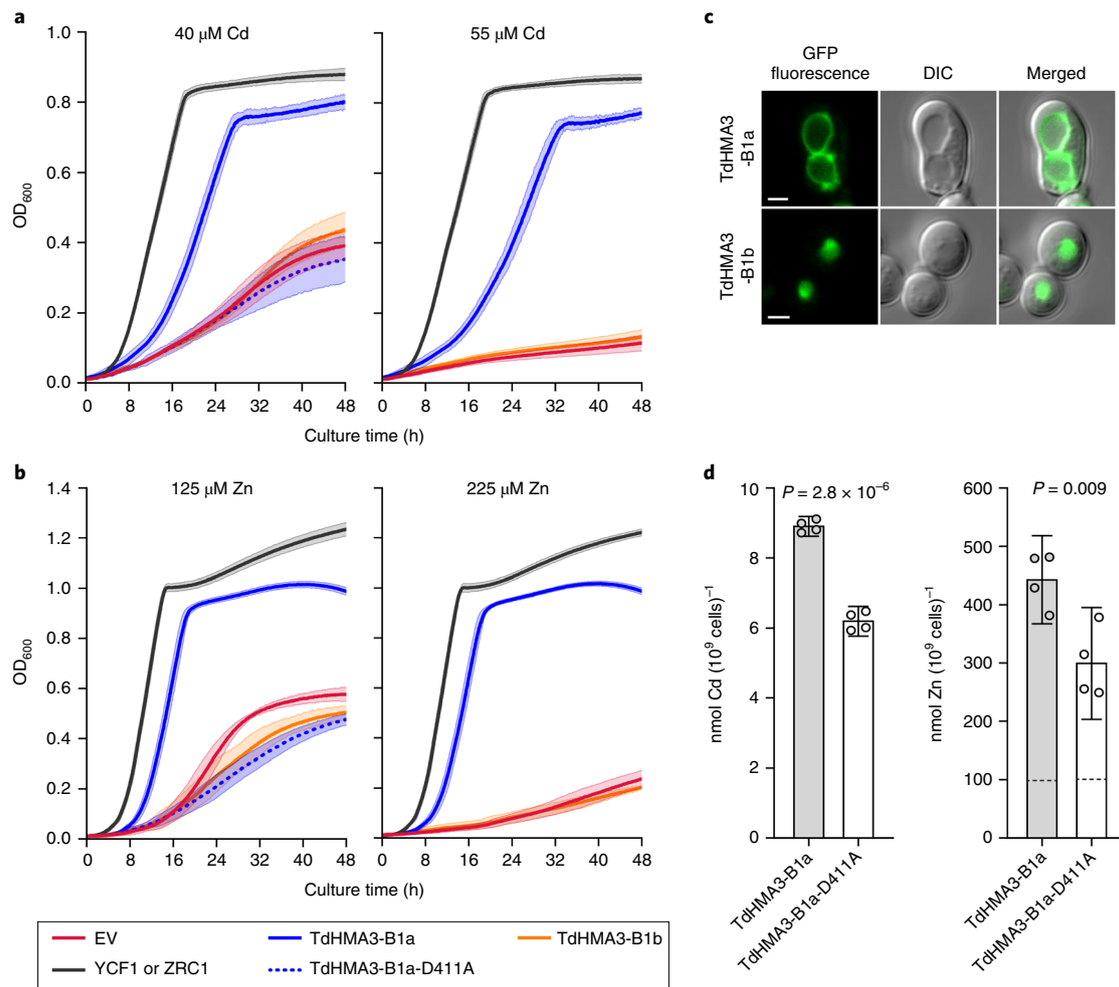


Fig. 7 | *TdHMA3-B1a* complements Cd- and Zn-sensitive mutant phenotypes of yeast. **a**, Growth (OD₆₀₀) of Cd-sensitive *ycf1* yeast expressing empty vector (EV, p413TEF), *YCF1*, *TdHMA3-B1a*, *TdHMA3-B1a-D411A* and *TdHMA3-B1b* in the presence of 40 and 55 μM Cd. Plotted growth curves are means of three (D411A), five (*YCF1*, *B1a*, *B1b*), or six (EV) experiments \pm 95% confidence intervals shown as shaded backgrounds. **b**, Growth (OD₆₀₀) of Zn-sensitive *zrc1cot1* yeast expressing empty vector (EV, p413TEF), *ZRC1*, *TdHMA3-B1a*, *TdHMA3-B1a-D411A* and *TdHMA3-B1b* in the presence of 125 and 225 μM Zn. Plotted growth curves are means of three (*B1b*) or five (EV, *ZRC1*, *B1a*, D411A) experiments \pm 95% confidence intervals shown as shaded backgrounds. **c**, Localization of *TdHMA3-B1a*-GFP and *TdHMA3-B1b*-GFP expressed in *ycf1* (differential interference contrast (DIC) and merged images provide spatial references). Scale bars, 2 μm . **d**, Cd accumulation in *ycf1* and Zn accumulation in *zrc1cot1* expressing *TdHMA3-B1a* and transport activity knockout, *TdHMA3-B1a-D411A*, after exposure to 5 μM Cd or 50 μM Zn for 4 h. Data are shown as means \pm 95% confidence intervals for $n = 4$ independent cultures (circles). Experiments repeated with similar results. P values were calculated by two-tailed, unpaired t -tests (d.f. = 6). The dashed lines indicate the Zn concentrations of the inoculating yeast cultures.

Frequently, two or more indexes occurred in overlapping regions, hereafter referred as selection clusters. In total, 104 pericentromeric (average size 107.7 Mb) and 350 non-pericentromeric (average size 11.4 Mb) clusters were identified in one, two or three transitions.

When 41 loci known to be under selection during emmer domestication and durum wheat evolution or breeding (Supplementary Table 12) were projected on the genome, many of them overlapped with selection clusters (Figs. 5 and 6; Supplementary Dataset 4). Most of the strongest pericentromeric diversity depletions (DRI > 4) occurred during emmer domestication (chromosomes 2A, 4A, 4B, 5A, 5B, 6A and 6B). Furthermore, one of the two brittle rachis regions marking the early domestication process (*BRT-3B*⁸) showed a localized sharp reduction in diversity confirmed by F_{st} and XP-CLR indexes. The same region, then, underwent an extreme diversity reduction in the DEW-to-DWL transition (DRI 3.4). Additional 14 pericentromeric and 90 non-pericentromeric (DRI > 2) diversity depletions, including one harboring the major tough glume QTL

governing threshability (Tg-2B²⁸), occurred during the DEW-to-DWL transition. Finally, several reductions in diversity (75 with DRI > 2) were specifically associated with breeding of modern durum cultivars, including some associated with disease resistance (for example, *Sr13*; ref. ²⁹ and *Lr14*; ref. ³⁰) and grain yellow pigment content loci (for example, *Psy-B1*; ref. ³¹). A detailed description of the selections signatures is presented in Supplementary Note.

Variation for cadmium grain content in tetraploid wheat. *Cdu-B1* is a QTL located on the long arm of chromosome 5B, which accounts for >80% of the phenotypic variation in cadmium (Cd) concentration in grain^{9,32,33}. The *Cdu-B1* region corresponds to a physical interval of 4.27 Mb. A detailed comparison of the Zavitan and Svevo (low and high Cd, respectively) genomes, coupled with exome sequencing, revealed a segment of increased nucleotide variation in this refined region (Supplementary Fig. 6). Furthermore, the region contains 192 gene models, 48 of which have informative

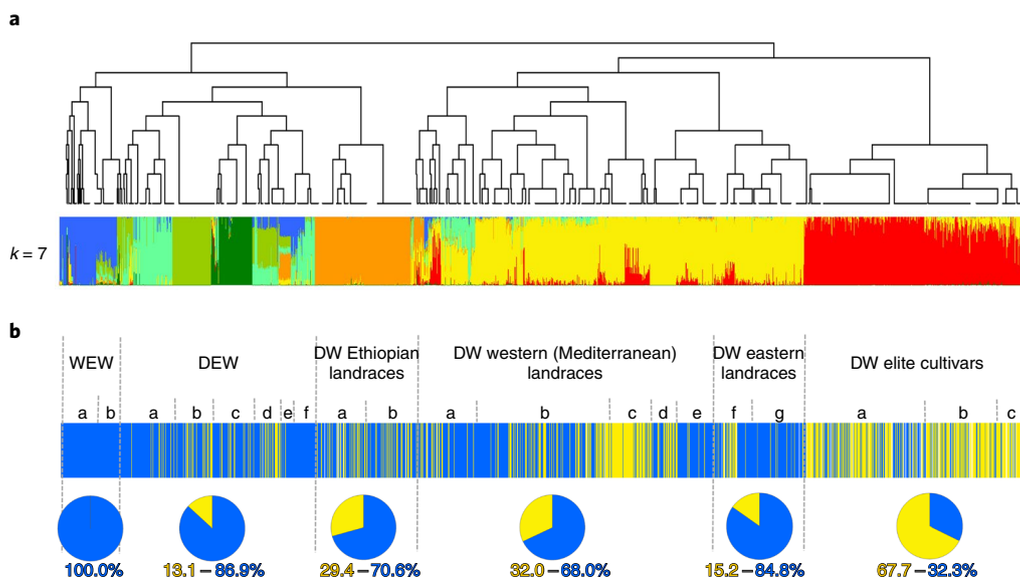


Fig. 8 | Population structure and *TdHMA3-B1a/b* allelic distribution in the Global Tetraploid Wheat Collection (1,856 accessions; Supplementary Table 11). **a**, FineSTRUCTURE clustering and bar plots of individual ADMIXTURE membership coefficients at critical $k=7$ for the tetraploid diversity panel. **b**, Bar plot of individual *TdHMA3-B1a/b* allelic score in WEW, DEW, DWL and DWC. Pie charts of *TdHMA3-B1a/b* relative ratio in main tetraploid germplasm groups. Blue bars, accessions carrying *TdHMA3-B1a*; yellow bars, accessions carrying *TdHMA3-B1b*.

functional annotations (Supplementary Table 13). One of these genes, *TRITD5Bv1G197370*, encodes a P_{1B} -type heavy-metal ATPase 3 (HMA3) that is orthologous to rice HMA3 (*OsHMA3*) (Supplementary Table 13 and Supplementary Fig. 7). *OsHMA3* is a tonoplast-localized transporter that transports Cd into vacuoles, thereby limiting its translocation from roots to shoots and grain^{34–36}. Closer inspection of *TRITD5Bv1G197370* (*TdHMA3-B1* herein) revealed a 17-bp duplication in the first exon that creates two alternative alleles, namely *TdHMA3-B1a* in Zavitan (functional; low grain Cd) and *TdHMA3-B1b* in Svevo (non-functional; high grain Cd; Supplementary Fig. 7). The alternative alleles are clearly identified by *Xusw59*, a diagnostic polymerase chain reaction (PCR) marker that amplifies the region including the 17-bp duplication (Supplementary Fig. 7) and perfectly discriminated low- and high-Cd accumulators in a global collection of DW lines (Supplementary Fig. 8). Heterologous expression of *TdHMA3-B1a/b* alleles in Cd- and zinc (Zn)-sensitive yeast confirmed that *TdHMA3-B1a* is a tonoplast-localized Cd and Zn transporter (Fig. 7 and Supplementary Figs. 9–12). Furthermore, disruption of P-ATPase ion transport activity (that is, *TdHMA3-B1a-D411A* knockout) reduced both *TdHMA3-B1a*-mediated yeast complementation (Fig. 7a,b and Supplementary Fig. 11) and cellular Cd and Zn accumulation (Fig. 7d and Supplementary Fig. 13). These findings demonstrate that *TdHMA3-B1a* transports Cd and Zn into vacuoles. Functional analysis of the *TdHMA3-B1* homoeolog, *TdHMA3-A1* (Supplementary Figs. 7b,c and 9) and the longest alternative open reading frame for allele *TdHMA3-B1b*, *TdHMA3-B1b-ORF2* (Supplementary Fig. 7b), are provided in the Supplementary Note. Consistent with the predicted function of *TdHMA3-B1*, the primary in planta effect of the non-functional allele, *TdHMA3-B1b*, is the reduced Cd retention in roots and a two- to threefold increase in Cd transport to shoots and grain (Supplementary Figs. 14 and 15 and Supplementary Table 14)³⁷. Although other genes in the *Cdu-B1* physical interval could potentially contribute to *Cdu-B1*, only *TdHMA3-B1* is functionally consistent with the *Cdu-B1* phenotype and is supported by these results as a candidate for the Cd phenotypic differences in DW.

The distribution of the *TdHMA3-B1a* and *B1b* alleles in the Global Tetraploid Wheat Collection revealed a clear association

with domestication status and geographical provenance (Fig. 8, Supplementary Fig. 16, Supplementary Tables 15 and 16 and Supplementary Dataset 2). The functional *TdHMA3-B1a* allele was genetically fixed in all 115 WEW accessions, while the *TdHMA3-B1b* allele showed a trend of increasing frequency in DEW (13%), DWL (26%) and DWC (68%). Among DEW accessions, the non-functional, high-Cd-accumulating *TdHMA3-B1b* allele was enriched in the subgroup from Turkey^{38,39} (38%), indicating Turkey to be the region of origin and location from which the allele spread. Among DWL accessions, *TdHMA3-B1b* showed a maximum of 41% occurrence in North African landraces (Fig. 8b). The steady increase in the frequency of the *TdHMA3-B1b* allele from DEW to DWC indicates a process of systematic selection or divergence during durum wheat evolution and breeding as demonstrated by the local F_{st} data (Supplementary Dataset 5). This result might indicate either a positive effect of the high-Cd allele per se or the presence of a selective sweep for another gene in linkage disequilibrium with *TdHMA3-B1b*. To assess the presence of genes which could have been targeted by selection in this locus, we identified the boundaries of high linkage disequilibrium with *TdHMA3-B1* (squared correlation coefficient $r^2 \geq 0.5$) in both DWL and DWC. The region contains 219 HC and LC genes in both Svevo and Zavitan as well as 73 and 58 genes present only in WEW and DW, respectively (Supplementary Dataset 6). Although the vernalization responsive gene *VRN-B1* (ref. 40) lies in this region (6.94 Mb distal from *TdHMA3-B1*), inspection of allelic diversity revealed *VRN-B1* as a low-variant gene, with the ancestral wild-type *vrn-B1* dominating across the entire Global Tetraploid Wheat Collection, thus excluding *VRN-B1* as a contributor to a possible selective sweep around *Cdu-B1*.

Discussion

The genome assembly of the modern DW cv. Svevo, with a quality level consistent with those recently obtained for other species^{8,11,15}, represents an essential tool to study durum wheat domestication, evolution and breeding as well as to gain new insights into gene function and the genome-wide organization of QTLs for relevant agronomic traits. This study presents an inclusive analysis of a large

panel of tetraploid wheat representing all known taxa and provides a global picture of genetic relationship and population structure. The process leading to modern durum wheat was revealed by the four main germplasm groups of the Global Tetraploid Wheat Collection. The combination of genetic diversity and selection signature analysis revealed a dynamic description of the modifications imposed on the genome by domestication and breeding. The strongest reductions in diversity occurred in well-defined pericentromeric regions during the domestication of WEW. Then, the reduction of diversity continued more moderately, but spread over the genome, during the evolution of DWL and, more recently, as a consequence of the breeding activity^{23,41}. Multiple divergence and haplotype metrics identified several regions coincident with known domestication loci, as well as others that might indicate new putative loci under domestication or selection.

Identification of *TdHMA3-B1* as the gene most likely responsible for phenotypic variation in grain Cd accumulation, a result supported by genetic and functional evidence, and the recovery of the *TdHMA3-B1a* allele for low Cd accumulation, provides an example of the relevance of the genomic tools presented here. The increase in frequency of *TdHMA3-B1b* during DW breeding could be due to the presence of a selective sweep for another gene in the linkage disequilibrium region, although no evidence has been found and further studies are required to support this hypothesis. Alternatively, the non-functional *TdHMA3-B1b* allele could exert some beneficial effects on plant fitness. Zinc assimilation by plants results in the co-transportation of Cd, and like other P_{1B,2}-type ATPase transporters¹², *TdHMA3-B1* can transport both metals. Although *Cdu-B1* has no effect on agronomic performance under Zn-sufficient conditions, the low-Cd line from a pair of *Cdu-B1* near-isogenic lines showed reduced biomass compared to the high-Cd line when grown under Zn-deficiency⁴³. Therefore, *TdHMA3-B1b* could provide a growth benefit in Zn-deficient soils, such as those that widely occur in wheat-growing regions of Turkey⁴⁴ where *TdHMA3-B1b* originated. A reduction in root vacuolar sequestration of Cd and Zn in high-Cd genotypes (non-functional *TdHMA3-B1b* allele) under Zn-limiting conditions may increase the pool of Zn available for transport to the shoot, thereby sustaining shoot growth.

Access to the fully annotated genome sequence in combination with the wealth of genotypic, genetic mapping¹² and gene expression data provides great potential for future innovation for the wheat scientific community and the breeding sector. Gene discovery, QTL cloning and the precision of genomics-assisted breeding to enhance grain quality and quantity of pasta wheat will benefit from the resources presented here. Furthermore, the durum sequence provides a fundamental tool to more effectively bridge and harness the allelic diversity present in wheat ancestors most of which remains largely untapped.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, statements of data availability and associated accession codes are available at <https://doi.org/10.1038/s41588-019-0381-3>.

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Author contributions

L.C., C.J.P., K.F.X.M., A.C. and RT conceived the study. A.S., G.S., P.D.V., N.P., L.M., H.B., A.D., A.C., C.J.P. and L.C. planned, and organized the sequencing steps. S.C.M., A.H., M.Mas. and N.S. carried out chromosome conformation capture and curated the final genome assembly. S.O.T., H.G., M.S., D.O., T.L., V.P., P.B., P.F., P.C., M.L., B.L., A.S., S.C., M.Mac., S.B., F.D., C.M., C.C., E.M., D.M., E.M., D.N., A.G., H.B. and A.M.M. contributed to genome annotations. A.M., M.G., M.Mo. and L.M. curated the genome visualization. S.O.T., H.G., M.S., D.O., T.L., V.P., R.A., J.D., A.D. and K.F.X.M. conducted the comparison between Svevo and Zavitan genomes. A.T.O.M. and I.H. carried out the identification of loss of functional mutations. M.Mac., R.K.P., D.O., E.F., S.C., S.S., F.D., H.O., B.K., D.M., R.J., E.M., S.C., J.D.F., M.P., A.M.M., S.S.X., A.D. and M.J.H. planned and carried out the analysis of genetic diversity and the identification of selective sweeps. N.S.H., K.W., J.E., R.P.M.L., J.M.C., A.G.S., S.C.K., K.Y.H.L., G.J.T., R.K., S.W. and C.J.P. planned and carried out the isolation and functional analysis of TdHMA3-B1. L.C., M.Mac., A.M.M., S.W., K.F.X.M. and C.J.P. organized and managed the contributions of others to this publication and produced the first draft of it. L.C. served as Senior Project Coordinator. All authors discussed the results and commented on the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Methods

DW genomic sequencing and assembly. *DNA extraction and sequencing.* DW genomic DNA was isolated from fresh leaf tissue (~10 g) of DW cv. Svevo using a phenol/chloroform large-scale nucleus extraction protocol⁴⁵. Five size-selected genomic DNA libraries ranging from 450 bp to 10 kb were sequenced to generate a total of 3.35 terabases (Tb) of data (equivalent to 279× coverage, on the basis of an estimated genome size of 12 Gb) using standard protocols. The libraries were constructed at the University of Illinois Roy J. Carver Biotechnology Center and sequenced at University of Salerno and Genomix4life by using an Illumina HiSeq2500 platform and HiSeqX instrument, according to the manufacturer's instructions (Supplementary Table 1).

Scaffold assembly. The DW genome was assembled from the 3.35 Tb of data using the proprietary software package DenovoMAGIC2 (NRGene) as previously described⁸. PCR duplicates, adapters and linkers were removed, and paired-end reads found to contain probable sequencing errors (that is, sub-sequences of ≥23 bp not found in at least one other independent read) were culled. From the paired-end libraries, only pairs with at least 10 bp sequence overlap were merged to create stitched reads. The stitched reads were then used to build an initial De Bruijn graph of contigs (*k*-mer, 191 bp). By exploring graph structure, the software identifies non-repetitive contigs and uses stitched reads information to resolve repeats and extend non-repetitive sequences of the contigs, where possible (Supplementary Table 2). Scaffolding was completed using a directed graph containing contigs as nodes and edges were on the basis of the paired-end and mate-pair links as vertices. Erroneous connections were identified and filtered out to generate unconnected sub-graphs that were ordered into scaffolds. Paired-end reads were used to find reliable paths in the graph for additional repeat resolving. This was accomplished by searching the graph for a unique path of contigs connecting pairs of reads mapping to two different non-repetitive contigs. The scaffolds were then further ordered and linked using the mate-pair libraries, estimating gaps between the contigs according to the distance of mate-pair links. Linking scaffolds with mate-pair reads required confirmation of at least three filtered mate-pairs or at least one filtered mate-pair with supporting confirmation from two or more filter failed mate-pairs where the Nextera adapter was not found. Scaffolds shorter than 380 bp were masked and links between non-repetitive contigs mapping to the same scaffolds were merged, generating a directed scaffold graph. The scaffolding procedure identified the non-branched components in the scaffolds graph, filtered out the rare connections between them and generated topological sorting based ordering of the initial scaffolds into the final scaffolds (Supplementary Table 2).

Chromosome conformation capture sequencing (Hi-C) and pseudomolecule construction. Scaffolds were assembled into pseudomolecules using data from two chromosomes conformation capture sequencing (Hi-C) libraries generated using the TCC protocol⁴⁶ as previously described¹¹. The libraries were sequenced using an Illumina HiSeq2500 instrument following the manufacturer's instructions (each TCC library on two lanes, paired-end, 2×100). In silico HindIII digestion of the Svevo sequence assembly by NRGene and assignment of the pre-processed reads to the restriction fragments was done following the methods described⁴⁷.

To identify possible misjoins in the Svevo NRGene assembly, we aligned the IWGSC chromosome survey sequencing hexaploid wheat contigs³⁸ to the assembly and lifted the POPSEQ map positions and flow-sorted chromosome assignment from the chromosome survey sequencing contigs to the durum scaffolds as described⁸. A total of 18 putative chimeric scaffolds (an example is reported in Supplementary Fig. 17) were detected and split. All the identified misjoins had also evidence from flow-sorting sequencing data and Hi-C links from different chromosomes. The improved durum wheat assembly (10.45 Gb, scaffold

N50 of 5,972,063 bp; Supplementary Table 2) was used to construct the final pseudomolecules (Supplementary Table 3).

The construction of a chromosome-scale Hi-C map of the NRGene v.2 assembly comprised two steps: (1) chromosome assignment and genetic anchoring of the Svevo scaffolds, (2) ordering and orienting scaffolds by Hi-C link information. Scaffolds were assigned to chromosomes using POPSEQ and flow-sorting data as previously described¹⁰. We used the same methods for Hi-C map construction as for the barley and WEW genomes^{8,47}. We considered only intrachromosomal Hi-C links between pairs of scaffolds less than 20 cM apart. In the case of chromosome 3B, we used a threshold of 5 cM to avoid erroneous joins. Moreover, we did not consider scaffolds with fewer than 100 HindIII restriction sites for Hi-C mapping. Scaffolds were oriented as previously described⁴⁷ using a bin size of 1 Mb. After the manual removal of four outlier scaffolds on chromosome 5A, the Hi-C map was highly collinear to the Svevo×Zavitan genetic map (Hi-C, Supplementary Fig. 18).

The FASTA sequence assemblies representing the 14 durum wheat chromosomes (that is, pseudomolecules, 2,938 scaffolds and 9.96 Gb in total) were constructed on the basis of the final Hi-C map. One hundred N characters were inserted as gap sequence between adjacent scaffolds. In addition, 126,526 scaffolds not assigned to chromosomes were placed in a sequence named 'chrUn' (499 Mb). The inclusion of gap sequences increased the total size of the assembly from 10.45 Gb to 10.46 Gb.

Additional methods. Additional methods are detailed in the Supplementary Note.

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

Data availability

A dedicated DW genome browser with all genome information presented in this manuscript (sequences, annotations, markers, passport data, etc.) is available at <http://www.interomics.eu/durum-wheat-genome>. Svevo genome and gene model were submitted to EBI-ENA under the study PRJEB22687. Svevo genome, gene model, SNP and QTL data and passport information of the Global Tetraploid Wheat Collection are available at GrainGenes (https://wheat.pw.usda.gov/GG3/browse_Durum_Svevo). RNA-seq and microRNA (miRNA) datasets can be downloaded at the SRA database under accessions SRP149116 (study: PRJNA473404). Durum wheat cv. Svevo GFF3 and VCF annotation files can be downloaded at figshare, <http://doi.org/10.6084/m9.figshare.6984035>. The WEW acc. Zavitan gene model version 2 is available at <https://doi.org/10.5447/ipk/2019/0>. The accessions used in this study are available on request and on the basis of a Material Transfer Agreement from the corresponding author (L.C.).

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Software and code

Policy information about [availability of computer code](#)

Data collection

This work has taken advantage from the novel genome sequence of durum wheat cv. Svevo generated by the authors and of existing dataset downloaded from public databases as detailed in the Supplementary note (section 1: Additional Materials and Methods). Swiss-Prot (version 02-15-10), Arabidopsis Araport 11 (version 201606), a TrEMBL (version 02-15-10) and UniProt databases were the main sources.

Data analysis

This work employed a vast number of software and data analyses, each software is cited in the Supplementary note (section 1: Additional Materials and Methods) along with the corresponding reference or website information. In summary (numbers refer to the references listed in the Supplementary note file):

1- Genome assembly was carried out using the proprietary software package DenovoMAGIC2TM (NRGene, Nes Ziona, Israel) as described⁸.

2- Genome annotations was supported with the following tools:

- HISAT2 (version 2.0.4)⁹ to align multiple sets of RNA-seq data to the assemblies;
- Stringtie (version 1.2.3)⁹ to assemble mapped reads into transcript sequences for each dataset separately;
- GMAP11 (version 06/30/2016) to align all sequences to the assemblies;
- Cuffcompare from Cufflinks software suite¹² for transcript predictions;
- Transdecoder package (version 3.0.0) to extract the longest open reading frames for each transcript sequence and to translate them into predicted protein sequences;
- AHRD tool (Automated Assignment of Human Readable Descriptions, <https://github.com/groupschoof/AHRD>, version 3.3.3) to annotate gene functions;
- BUSCO (Benchmarking Universal Single-Copy Orthologs) tool (version 2, Embryophyta odb9)¹⁴ to determine the abundance of strongly conserved genes in the sets of all annotated genes;
- Tallymer¹⁵ to calculate the Basic k-mer defined repetitivity;
- Vmatch (www.vmatch.de) for transposons detection and classification by a homology search against the REdat_9.7_Triticeae and the

PGSB transposon library¹⁶;

- LTRharvest¹⁷ to identify full-length LTR retrotransposons (fl-LTRs);
- NLR-annotator version 0.7 pipeline (<https://github.com/steuernb/NLR-Annotator26>) to annotate the loci associated with Nucleotide-Binding Leucine-Rich Repeat domains (NLRs);
- CpGCluster²⁷ to detect CpG islands;
- Find Individual Motif Occurrences (FIMO)²⁸ to annotate Transcription factors binding sites (TFBS);
- Blast2GO PRO to assign GO terms;
- Bioconductor package GOSTATS²⁹ (version 2.42.0) for GO enrichment call;
- CD-HIT^{33,34} (version 4.6.5) and Tandem Repeats Finder³⁶ to identify domains with high copy number;
- OrthoMCL (version 2.0, www.orthomcl.org) to search for gene families.

3- Genetic maps were produced following a pipeline including: scripts (https://github.com/plantinformatics/Durum_iSelect_90kSNP_GenotypeCalling) for genotype calling in unrelated samples, sample cluster assignment, confidence score estimates, and final genotype call from Illumina raw data project files; ii) quality check and filtering of genotype calls; iii) marker grouping and ordering in MST-map⁴² (<http://www.mstmap.org/>), MetaQTL analysis and projection of QTLs to DW assembly was calculated using the BiomeRCator version 4.2 software⁴⁶

4- For the analysis of genetic diversity the raw data from Illumina genotyping were jointly analyzed for cluster assignment and genotype calling using a custom script for genotype calling in unrelated samples (https://github.com/plantinformatics/Durum_iSelect_90kSNP_GenotypeCalling). Pairwise LD values were estimated by means of the `snpGdsLDMat` function in the R package `SNPRelate`⁵³,

The NJ phylogenetic tree was obtained by calculating the pairwise genetic distances, performing 1,000 bootstrap resampling, and obtaining the tree in R, using the `dist.gene`, `boot.phylo`, `write.tree` and `write.nexus` functions (`poppr`, `pegas`, `ape`, `adegenet`, `ade4` libraries). PCA was performed using `EIGENSTRAT`⁵⁶. Phylogenetic networks were computed using `SplitsTree4` version 4.14.665 while Fst statistics at each locus was computed by R package `pegas`⁷⁴. `fastPHASE` v1.4.876 and R package `imputeq`⁷⁷ were used to reconstruct the haplotypes from SNP data.

5- For construction of the refined interval for Cdu-B1 markers from the array were mapped to the genome of Svevo by GMAP¹¹. Markers uniquely mapping to chromosome 5B were used for Single Marker Regression analysis using Windows QTL Cartographer (<https://brwebportal.cos.ncsu.edu/qtlcart/index.php>). Sequence reads from exome capture experiment were processed by Trimmomatic version 0.32.81 and aligned to the genome of Svevo using Novoalign version 3.02.05 (www.novocraft.com/products/novoalign). Variants were called using the SamTools version 1.2.139.

6- For comparative sequence analysis of HMA gene, protein sequences or translated CDS sequences for P1B-ATPases (HMAs) from *Arabidopsis*, *Brachypodium distachyon*, and rice were compiled. DW HMA genes were identified by TBLASTN of the Svevo genome using *Brachypodium* and rice HMA proteins as queries (E-value < 10⁻³), and the DW HMA gene models were predicted with Fgenesh+⁸⁴ using relevant wheat or barley HMAs as homologs. The sequences and locus identifiers of the proteins included in the phylogenetic analysis are shown in Supplementary Data Set 12. Sequences were aligned with MAFFT L-INS-i (version 7.311, <https://mafft.cbrc.jp/alignment/server/>) using the default settings⁸⁵. Gaps and poorly aligned regions were removed from the multiple sequence alignment (MSA) by Gblocks version 0.91b86 using less stringent selection criteria⁸⁷; http://molevol.cmima.csic.es/castresana/Gblocks_server.html). The trimmed MSA consisted of 570 positions (31% of the untrimmed MSA), including 58 invariant sites. Phylogenetic tree was reconstructed using the maximum-likelihood method with PhyML version 3.188

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A dedicated durum wheat genome browser with all genome information presented in this manuscript (sequences, annotations, markers, passport data, etc.) is available at <http://www.interomics.eu/durum-wheat-genome>. Svevo genome and gene model were submitted to EBI-ENA under the study PRJEB22687. Svevo genome, gene model, SNP and QTL data and passport information of the GTC are available at GrainGenes (https://wheat.pw.usda.gov/GG3/jbrowse_Durum_Svevo). RNA-Seq and miRNA datasets can be downloaded at the SRA database under accessions SRP149116 (study: PRJNA473404). Durum wheat cv. Svevo GFF3 and VCF annotation files can be downloaded at: <https://figshare.com/>, accession: 10.6084/m9.figshare.6984035. The wild emmer wheat acc. Zavitan gene model version 2 is available at: <https://doi.org/10.5447/ipk/2019/0>. The accessions used in this study are available on request and based on a Material Transfer Agreement from the corresponding author (LC).

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Sample size

The article refers to 3 main experiments.

The first deals with the genome sequence assembly of a modern cultivar of durum wheat (Svevo), representing the first high quality cultivar released in Europe, for which a high resolution genetic map (Svevo x Zavitan) was already available. The Svevo genome was sequenced at 279X coverage (= 3.35 Tb of NGS data) using a combination of five size-selected genomic DNA libraries of insert size ranging from 450 bp (paired-end) up to 10 kb (mate pairs), in balanced relative amount. This sequencing data, analysed with the software package DenovoMAGIC2TM (available to NRGene, Nes Ziona, Israel), ensured to assemble 10.45 Gb of unique Svevo sequences, structured into contigs of N50-length = 56.20 Kb, scaffolds of N50-length = 5.972 Mb, and 14 Hi-C ordered pseudomolecules including gaps for 149.190 Mb only (1.42%). We provide a Svevo "golden-standard" assembly of quality comparable to that reported for the tetraploid wild emmer (Avni et al. 2017), Ae. tauschii (Luo et al. 2017), and Chinese Spring hexaploid wheat (IWGSC, 2018). The empirical and statistical experimental procedure that led to the choice of the sequencing depth is reported in Avni et al. (2017). The final assembly provides 66.559 high-confidence and 303,404 low-confidence genes. The completeness of gene content was then validated based on gene representativeness from independent public databases (98.1% of BUSCO genes, 97.7% of a dataset of experimentally determined genes, 92.3% of Triticeae reference proteins). It also provides 51,077 high-quality resolved full-length LTR retrotransposons.

The second experiment describes the diversity in tetraploid wheats using 1,854 accessions, this is the largest diversity panel reported so far for tetraploid wheat. It represents all known AABB-genome subspecies and samples all regions where tetraploid wheat (both wild and cultivated) is spread. In particular, the collection samples wild emmer wheat (WEW), domesticated emmer wheat (DEW), durum-related subspecies, durum wheat landraces (DWL) and durum wheat cultivars (DWC) in balanced counts. The collection has been assembled from world-wide researchers long time-involved in diversity and genomics of tetraploid wheat, and supplemented by additional 490 accessions specifically sampled from domestication areas. The whole-genome genetic diversity survey of the four main domestication-related sets (WEW, DEW, DWL, DWC) of tetraploid wheat was carried out based on the well-established and highly informative iSelect 90K wheat SNP assay. After filtering for uniqueness and information content, it provided a total of 17,416 transcript-associated SNPs (1.66/Mb or 16.6/10Mb). Based on the linkage disequilibrium decay rate in the four germplasm sets considered, the number of SNPs was considered adequate to assess the relative genetic diversity losses/gains among domestication-related sets.

The third experiment describes the cloning of a locus controlling Cd accumulation. This work was carried out using a large panel of genetic materials including 3 segregating populations: the high-density Svevo x Zavitan reference mapping population, a large F2 population of 5,081 individuals suitable for fine-mapping at 0.1 cM scale and a panel of cultivars and breeding lines. Further, a survey of the causal polymorphism was extended to the whole diversity panel (1,854 worldwide accessions).

Data exclusions

No specific data were excluded.

Replication

The analysis and assembly of genomic Illumina NGS data involved initial several standard quality-control and filtering steps as described in Supplemental Materials & Methods.

At the core of the analysis, the software package DenovoMAGIC2TM (available to NRGene, Nes Ziona, Israel) uses very stringent parameters and thresholds for all contig assembling and scaffolding steps as described in Supplemental Materials & Methods. In details, elongated or stitched reads were produced upon an overlap threshold of 10 bp minimum. Initial contig assembly was carried out based on De Bruijn graphs at very high kmer = 191 bp. The assembler has special features for the identification of unique non-repetitive sequences, for resolving repeats and, in the scaffolding step, to identify erroneous connections. Stringent parameters were also used for scaffold ordering and linking based on the mate-pairs data.

Both high-density genetic map and chromosome conformation capture sequencing (Hi-C) were used for scaffold validation, correction and re-ordering of final pseudomolecules. Only intra-chromosomal Hi-C links below certain distance-thresholds were used. Available IWGSC contigs and POPSEQ for hexaploid wheat were used as further validation/correction steps.

The annotation pipeline is robust as it has been developed and tested on the most advanced Triticeae genome assemblies, including barley, Aegilops tauschii and hexaploid wheat, combining in-silico bioinformatics and gene expression data. Several quality-thresholds were applied to assemble RNA-seq reads into transcripts. Finally the whole-assembly validation was carried out by considering its gene representativeness when tested against three independent and validated gene/protein sets (BUSCO, lab-validated gene set, Triticeae protein set).

Micro-RNAs, long non coding RNAs, transposons and LTR retrotransposons, prolamin and NB-LRR gene families were annotated based on homology searches against Triticeae and more-comprehensive databases. Functional non tandem duplicated gene clusters were found based on GO-enrichment specific tests.

Genetic mapping was based on 17 interconnected maps, mostly based on the same iSelect 90K wheat SNP Illumina array platform, for which a common pipeline for genotype call and genetic map was adopted, starting from the original Illumina iSelect raw data files. The pipeline included several quality-check and filtering steps. Based on the available reference sequences of SNP and other markers, the genetic maps were anchored to the Svevo genome assembly using the most complete, gap-free high-density Svevo x Zavitan unique reference binned map. This genetic-to-physically anchored map was also used in the process of QTL projection on the final Svevo assembly.

The highly-robust and widely used iSelect 90K wheat SNP Infinium array-based genotyping platform was used for the diversity survey. The genotyping projects were all carried out in high-standard Illumina-certified genotyping labs. The raw iSelect Illumina SNP genotyping data obtained for the wide tetraploid wheat diversity panel from various genotyping projects were processed with the same pipeline described above for the genetic mapping data. Only unique Mendelian-segregating loci, whose genetic map location was coincident with the corresponding Svevo physical address, were used for further analysis.

In all the Illumina SNP genotyping projects including both genetic mapping and diversity accession panels, common reference genotypes were used as internal quality-standard checks. For this purpose, the well-known "Cappelli", "Langdon" and "Svevo" reference DNA, together with those from several parents of mapping populations were used.

After passport analysis and clustering of genetic profiles, the 2,558 tetraploid diversity panel accessions that were initially assembled from various genotyping projects were reduced to 1,854 non-redundant tetraploid accessions. Passport info were mostly useful to predict cases of duplicated samples or high genetic relationships, thus representing an additional quality check control.

The population genetic structure of the tetraploid diversity panel was assessed with two independent well-known softwares (ADMIXTURE and fineSTRUCTURE), based on replicated analysis runs as recommended by software analysis instruction manuals. The two softwares gave bi-directional comparable, highly-complementary results.

The whole-genome genetic diversity survey based on the four main domestication-related sets of tetraploid wheat was carried out using a

SNP dataset filtered for uniqueness of map location (dingle-locus Mendelian loci), absence of sigletons and double-singletons, and, in order to limit the interference effects caused by ascertainment bias particularly relevant for wild emmer and domesticated emmer accessions, the SNPs were further selected for overall null allele frequency < 0.25 (failure rate).

All the fine-mapping and positional cloning steps of the Cadmium transporter locus, including several genotyping and phenotyping steps, were carried out using standard replications. Same applies to the candidate gene expression analysis, including both in-planta and in yeast expression and complementation experiments. Importantly, near-isogenic line stocks, collections of diverse germplasms of adequate counts, and yeast mutant stocks were used as to most adequate materials to guarantee highly-repeatable and reliable results.

In conclusion, the experimental design, methods of replication and validation, and statistical methods used meet the standards outlined by Nature Genetics. All attempts at replication were successful. Several experiments required multiple replications and in each case the number of replications is described.

Randomization

When generating the Illumina NGS Svevo genome sequencing data, library construction was carried out assuring the high quality and integrity of the original genomic DNA used. DNA random fragmentation (shearing/sonication) and tagmentation for Nextera libraries were carried out by following carefully optimized Illumina protocols that ensured the random representativeness of genomic sequences for all paired-end and mate-pairs libraries. Concerning the distribution of sequencing depth among the different types of libraries, 123X coverage was dedicated to PE libraries of 450bp-insert size (half of total coverage), this was required to obtain highly-accurate contig assemblies. Coverage dedicated to each of the 750bp-PE and to the three Nextera Mate Pairs libraries (3kb, 5kb, 10kb) was equally partitioned (38-41X each) in order to ensure that balanced sequence information of different insert size was conveyed to the scaffold assembler.

During the selection of the accessions of the tetraploid diversity panel, great attention was given to sample accessions from each of the four main wheat germplasm groups (WEW, DEW, DWL, DWC) to ensure an accurate germplasm representativeness. Within each group, care in uniform sub-regional sampling and further germplasm bank's accession availability inspection was considered to represent all main sub-areas related to diffusion and domestication. Within sub-areas, random sampling in balanced numbers was carried out, after passport inspections of seedbank available accessions, in order to limit as much as possible sampling of duplicated or highly related accessions. Prior to the final diversity analysis, great care was taken in the joint inspection of passport and molecular information available in order to filter out clearly duplicated, highly similar and redundant accessions. Germplasm structure was assessed in great details, based on two independent dedicated software.

As to the whole-genome scan for differential gene diversity among the four main germplasm subgroups, given the predominantly descriptive objective of this analysis, namely describing the diversity present among germplasm collections, selective sweep tests and corrections for population structure were not applied, except for the initial withdrawal of the Ethiopian DEW and DWL accessions, two groups clearly distinct from the main bulks of European, Mediterranean and Central Asian germplasm. A similar approach was applied for the Cadmium-transporter allele survey distribution (carried out for all 1,854 accessions available).

As to the fine-mapping and positional cloning steps of the Cadmium transporter locus, including several genotyping and phenotyping steps, standard randomization best practices were used across all experiments.

Blinding

Not applicable. This research did not include experiments with observer biases.

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