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**Molecular phylogenetic analysis of amylase trypsin inhibitors (ATIs) from a selection of  
ancient and modern wheat**

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## 28    **Abstract**

29    Wheat amylase-trypsin inhibitors (ATIs) are a family of wheat proteins, which play an important  
30    role in plant defence against pest attacks. Recently, ATIs have been identified as major stimulators  
31    of human innate immune cells leading to cause Non-coeliac Wheat Sensitivity. Information about  
32    ATI sequence differences among wheat species is scarce, especially considering ancient wheat  
33    genotypes. In this study, ten selected wheat accessions with different ploidy level and year of  
34    release were used for gene sequencing of four representative alpha-amylase/trypsin inhibitor genes  
35    (WMAI, WDAI, WTAI-CM3 and CMx genes). The phylogenetic analysis and the PCA analysis  
36    performed on the deduced amino acid sequences of ATI genes evidenced that the ten wheat  
37    genotypes can be differentiated on the basis of their ploidy level, but not with respect to ancient or  
38    recently developed wheat genotypes. The haplotype analysis based on Nei's genetic distances,  
39    beside confirming these results, also allowed the separation of the hulled from the naked wheat  
40    genotypes. The genetic sequence differences highlighted in this study among the ten genotypes can  
41    be the basis for further studies aimed at identifying proinflammatory sequences in ATI genes. This  
42    is the first study analyzing the ATI genetic sequences of a set of ancient and modern wheat  
43    genotypes with a different ploidy level.

## 44    **1. Introduction**

45    Wheat amylase-trypsin inhibitors (ATIs) represent ~2–4% of total wheat protein and are an  
46    important family of low molecular weight, water-soluble proteins located in the endosperm of the  
47    grain kernel where they play a crucial role in plant defence against pest attacks (Carbonero and  
48    García-Olmedo, 1999). ATIs are able to inhibit enzymes of common parasites such as mealworms  
49    and mealy bugs from digesting wheat starch and protein. Besides having a protective role, they act  
50    as storage proteins (Altenbach et al., 2011).

51    ATIs are classified into four groups: three groups of alpha-amylase inhibitors differentiated  
52    according to their degree of aggregation (monomeric, homodimeric and heterotetrameric forms) and  
53    one group of trypsin inhibitors (Carbonero and García-Olmedo, 1999). The three groups of alpha-  
54    amylase inhibitors include the wheat monomeric amylase inhibitors (WMAI), often called 0.28  
55    proteins; proteins that form the wheat homodimeric amylase inhibitors group (WDAI), sometimes  
56    called 0.19 and 0.53 proteins and the proteins of the wheat heterotetrameric amylase inhibitors  
57    (WTAI). The WMAI inhibitors are encoded by genes on the short arm of chromosome 6 of *T.*  
58    *aestivum*, while the genes for WDAI are located in the short arm of chromosome 3 (Carbonero and  
59    García-Olmedo, 1999), in particular on 3BS and 3DS chromosomes, though not much is known  
60    about homologous common wheat chromosome 3AS (Pandey et al., 2016). The tetrameric  
61    inhibitors are often called CM proteins because of their solubility in chloroform/methanol and in  
62    modern hexaploid wheat five sub-units have been identified: CM1, CM2, CM3, CM16 and CM17.  
63    CM proteins are generally composed by one copy of either CM1 or CM2 sub-unit, encoded by  
64    genes on chromosomes 7D or 7B, plus one copy of either CM16 or CM17 sub-unit, encoded by  
65    genes on chromosomes 4B and 4D, plus two copies of CM3 sub-unit, also encoded on  
66    chromosomes 4B and 4D. WMAI, WDAI and CM proteins have a molecular weight ranging from  
67    13.0 to 13.5 kDa at the subunit level, with the exception of CM3 which shows a mass of about 15.5  
68    kDa (Geisslitz, 2021). All these alpha-amylase inhibitors are active with different specificity against  
69    insect, mite and mammalian alpha-amylases, but are not active against cereal enzymes. The WTAI  
70    inhibitors are also able to inhibit papain, bovine trypsin and subtilisin. The putative wheat trypsin  
71    inhibitors are referred to as CMx proteins, are encoded by genes on the group 4 chromosomes and  
72    are monomeric (Altenbach, 2011; Carbonero and García-Olmedo, 1999).

73    Altenbach et al. (2011) identified 19 different alpha-amylase/protease inhibitors of ~120- to 150-  
74    amino acids expressed in the modern wheat genome. The main ATI species in wheat are 0.19  
75    (WDAI) and CM3 (WTAI).

76 All ATIs are characterized by high homology in their amino acid sequence and by an analogous  
77 compact secondary structure showing five or less (often four) intrachain disulphide bridges and a  
78 main body of the molecule built around four  $\alpha$ -helices arranged in an “up and down” pattern linked  
79 together by loop segments (Carbonero and García-Olmedo, 1999).

80 ATIs are of high interest because they have an important impact on human health. Some ATI  
81 proteins are involved in wheat allergies (Tatham and Shewry, 2008) and, recently, ATIs have been  
82 identified to play a role in the onset of celiac disease (Huebener et al., 2015) and Non-celiac Wheat  
83 Sensitivity (Junker et al., 2012). Several *in vitro* and *in vivo* studies identified ATIs (especially  
84 CM3 and 0.19), but not gluten, as a major nutritional trigger of human and murine innate immunity  
85 on wheat by the activation of the toll-like receptor 4 (TLR4) complex (Geisslitz, 2021; Junker et al.,  
86 2012; Zevallos, 2017).

87 Thus far, there is little information in literature about the diversity of ATIs between different types  
88 of wheat, especially between so-called “ancient” and modern wheat genotypes. Recently, Zevallos  
89 et al. (2017) found that modern hexaploid wheat showed higher ATI inflammatory activity than  
90 some ancient variants like diploid (einkorn) and tetraploid wheat (emmer, KAMUT® khorasan  
91 wheat) or older hexaploid variants like spelt. Another recent paper (Gélinas and Gagnon, 2018)  
92 determined the alpha-amylase inhibitory activity against human  $\alpha$ -amylase in several different  
93 wheat cultivars and showed that the inhibitory potential did not vary with respect to ancient or  
94 recently developed wheat cultivars. To the same conclusion came Call et al. (2020) who analysed  
95 the ATI concentrations and the trypsin inhibitor activity in a set of different *Triticum* species and  
96 EL Hassouni et al. (2021) who determined the content of eight ATI proteins as well as of the total  
97 ATI of 149 European old and modern bread wheat cultivars.

98 However, information about ATI genetic sequences is available only for a few types of wheat,  
99 especially for common wheat (*T. aestivum* L. subsp. *aestivum*) and for wild emmer wheat (*T.*  
100 *turgidum* L. subsp. *dicoccoides*) (Pandey et al., 2016; Wang et al., 2010, 2008, 2007), while ATI  
101 genetic sequences of some other types of wheat like spelt (*T. aestivum* L. subsp. *spelta*), emmer (*T.*  
102 *turgidum* L. subsp. *dicoccum*) and khorasan wheat (*T. turgidum* L. subsp. *uranicum*) have never  
103 been studied. Moreover, gene sequences of some ATIs like WTAI and CMx are very scarce (Liu  
104 and Wang, 2012). Lastly, little is known of the genetic diversity of ATIs between genotypes, in  
105 particular between ancient and modern wheat genotypes (Geisslitz et al., 2021).

106 In the present study, ten wheat accessions were selected to cover different *Triticum* species with a  
107 different genome composition (diploid, tetraploid, hexaploid) and with a different year of release to  
108 determine the genetic sequences of four representative alpha-amylase/trypsin inhibitor genes

109 (WMAI, WDAI, WTAI-CM3 and CMx genes). The genetic sequence differences among the wheat  
110 genotypes were investigated and discussed.

## 111 2. Material and methods

### 112 2.1 Plant material

113 In the present study, ten selected accessions belonging to different *Triticum* species i.e. *T.*  
114 *monococcum* L. subsp. *monococcum* ( $2n = 2x = 14$  chromosomes, AA genome), *T. turgidum* L.  
115 subsp. *dicoccum*, *T. turgidum* L. subsp. *turanicum*, *T. turgidum* L. subsp. *durum* (all  $2n = 4x = 28$   
116 chromosomes, AABB genomes), *T. aestivum* L. subsp. *aestivum*, *T. aestivum* L. subsp. *spelta* (all  
117  $2n = 6x = 42$  chromosomes, AABBDD genomes) were used for the characterization of ATI gene  
118 sequences (Table 1). These genotypes were selected to cover different wheat species with a  
119 different genome composition (diploid, tetraploid, hexaploid) and with a different year of release.  
120 The pedigrees of heritage and modern wheat varieties were retrieved in GRIS – Genetic Resources  
121 Information System for Wheat and Triticale (<http://wheatpedigree.net>) and are shown in Table 1.

### 122 2.2 DNA isolation and PCR amplification

123 Ten seeds for each wheat genotype were grown in sprouters at room temperature in daylight for  
124 seven days. The leaf tissues were sampled at the four-leaf stage from ten different plants per  
125 genotype, immediately frozen in liquid nitrogen and ground in a mortar with a pestle. Total DNA  
126 extraction was performed with the Nucleospin® Plant II kit (Macherey Nagel, Düren, Germany)  
127 following manufacturer's instructions. DNA quality and quantity was measured by  
128 NanoPhotometer® P-Class (Implen GmbH, München, Germany).

129 The primers used to amplify WMAI, WDAI, WTAI-CM3 and CMx genes are listed in Table 2.  
130 Primers F1, F2, F3, F4, R1 and R2 were previously used by Wang et al. (2008), primers WDAIfor  
131 and WDAIrev were previously used by Wang et al. (2007), the other primers were designed *in*  
132 *house* with the software Primer3 (<http://bioinfo.ut.ee/primer3-0.4.0/primer3/>) on the conserved  
133 sequences of the coding regions of genes obtained from the GenBank database.

134 WMAI 0.28 genes were amplified in common wheat samples using F2 and R1 primers and in  
135 einkorn, spelt and Peliss genotypes using Mfor3 and Mrev2. Emmer, turanicum and Alzada samples  
136 were also amplified using the forward primers F1, F3, F4, Mfor1, Mfor2, Mfor4, Mfor5 and the  
137 reverse primers R2, Mrev1, Mrev3, Mrev4, Mrev5, Mrev6. The different combinations of primer  
138 sets used for the amplification of WMAI genes are listed in Supplementary Table 1 with the  
139 corresponding annealing temperatures used and the expected amplicon sizes.

140 WDAI 0.53 and 0.19 genes were amplified using WDAIfor and WDAIrev primers with an  
141 annealing temperature of 58 °C. The expected amplicon size was 446 bp.



WTAI-CM3 genes were amplified using CM3for and CM3rev primers and an annealing temperature of 55 °C. The expected amplicon size was 647 bp. In order to check for the possible presence of WTAI-CM3 gene sequences also in einkorn, the internal forward primers CM3intfor and CM3intfor2 and the internal reverse primers CM3intrev and CM3intrev2 were designed and used in combinations at the annealing temperature of 50 °C.

CMx1/CMx3 and CMx2 genes were amplified using CMxfor and CMxrev primers and an annealing temperature of 60 °C. The expected amplicon size was 498 bp.

All the PCR amplifications were performed with Biometra® ThermoCycler (Biosense srl, Milan, Italy) in 50 µl volume, consisting of 200-400 ng genomic DNA, 1 µM each primer, 1x HotStar HiFidelity PCR Buffer (containing 1.5 mM MgSO<sub>4</sub> and 0.3 mM dNTPs), 2.5 U of HotStar HiFidelity DNA Polymerase (QIAGEN, Hilden, Germany). Due to the lack of amplification of WMAI genes in emmer, turanicum and Alzada genotypes and WTAI-CM3 genes in einkorn, MgSO<sub>4</sub> was also tested at final concentration of 2 and 3 mM to enhance the stability of primer-template complexes in order to further check the presence of these genes in the mentioned genotypes.

The cycling parameters were 95 °C for 5 min to pre-denature, followed by 45 cycles of 94 °C for 15 sec, annealing temperature (specific for each primer set) for 1 min, and 72 °C for 90 sec, and a final extension at 72 °C for 10 min.

The desired DNA fragment was recovered from 1.5 % agarose gel using QIAquick gel extraction kit (QIAGEN, Hilden, Germany), quantified by NanoPhotometer® P-Class (Implen GmbH, München, Germany), ligated to the pDrive Cloning Vector and used to transform *E. coli* competent cells by using QIAGEN® PCR Cloning Plus Kit (QIAGEN, Hilden, Germany). For each amplified gene (WMAI, WDAI, WTAI-CM3, CMx), five positive clones obtained from each wheat genotype were screened and sequenced with the Sanger method using the Brilliant Dye Terminator 1.1 kit (NimaGen BV, The Netherlands) and with ABI3730 automated sequencer. Sequences were obtained using the service provided by BMR genomics (Padova, Italy).

## **2.3 Bioinformatics analysis**

### **2.3.1 Haplotypes identification**

Open reading frames were obtained by using ORF Finder (<http://www.ncbi.nlm.nih.gov/gorf/gorf.html>) and EMBL-EBI server ([https://www.ebi.ac.uk/Tools/st/emboss\\_transeq/](https://www.ebi.ac.uk/Tools/st/emboss_transeq/)). A search for similarity was executed with the BLASTN and BLASTP programs available at the National Centre for Biotechnology Information

(NCBI). Haplotypes identification was performed by MEGA software Version 7.0.26 (Kumar, 2016). Multiple sequence alignments were performed using Clustal Omega program (<http://www.ebi.ac.uk/Tools/msa/clustalo/>), MEGA software Version 7.0.26 and Jalview version 2.11.0 (Waterhouse, 2009). Sequence identity analysis was carried out with SIAS server (<http://imed.med.ucm.es/Tools/sias.html>). Amino acid sequences of reference proteins were retrieved from the Universal Protein Resource database (Uniprot) (<https://www.uniprot.org/>).

The signal peptides and the location of their cleavage sites in proteins were determined with SignalP 5.0 (<http://www.cbs.dtu.dk/services/SignalP/>).

Obtained sequence data have been deposited in EMBL/GenBank Data Libraries under accession nos. MT739524-38 (WMAI genes); MT861996-2029 (WDAI genes); MT887935-58 (WTAI-CM3 genes) and MT897967-88 (CMx genes).

### 2.3.2 Phylogenetic analysis

The phylogenetic relationships between haplotypes were performed using MEGA software Version 7.0.26. The FASTA multiple sequence alignment was used to infer the Neighbor-Joining (NJ) phylogenetic tree. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic tree. The evolutionary distances were computed using the Poisson correction method and are in the units of the number of amino acid substitutions per site. All positions containing gaps and missing data were eliminated. Evolutionary analyses were conducted with 1000 bootstrap replicates (Kumar, 2016).

### 2.3.3 Principal component analysis (PCA)

PCA was used to transform a number of correlated variables into a smaller number of uncorrelated variables called principal components. With this method the original space for variable measurements was projected down onto two low dimensional subspaces. One of these was case-related (ten wheat genotypes), the other was variable-related. The variables were the amino acid substitutions at a specific position. The variable-related subspace was analysed (factor loading) to understand the correlation between the variables and wheat genotypes (principal component).

First, an interim PCA was run with all the variables included and the quality of representation (squared cosine, cos<sup>2</sup>) of each variable was checked. Second, since a high cos<sup>2</sup> indicates a good representation of the variable on the principal component, a definitive PCA was run with those variables with a quality of representation higher than 0.5. The data were not pretreated before submission to PCA since a homogeneous set of variables was used (i.e. amino acid substitutions at

205 a specific position). The PCA analysis was based on correlation matrix and was performed using  
206 STATISTICA Software v. 7.1 (StatSoft, Tulsa, Oklahoma, USA).

#### 207 **2.3.4 Haplotype analysis**

208 The program POPGENE 1.31 was used to carry out the haplotype analysis. The genetic distances  
209 were calculated to compare the ten wheat genotypes (Nei, 1972) based on the cluster analysis of a  
210 0-1 matrix, where the presence or absence of the haplotypes of all the four genes analysed in this  
211 study was recorded as 1 and 0, respectively. The dendrogram was constructed based on the genetic  
212 distances using UPGMA (unweighted pair group with arithmetic average) method modified from  
213 Neighbor procedure of PHYLIP Version 3.5.

### 214 3. Results and discussion

#### 215 3.1 Sequence analysis of monomeric alpha-amylase inhibitors (WMAI)

216 One desirable PCR product of the expected size was obtained in common wheat samples Judee,  
217 Turkey Red, Marquis, Vida (~529 bp using F2 + R1) and in einkorn, spelt and Peliss genotypes  
218 (~502 bp using Mfor3 + Mrev2).

219 Emmer, turanicum and Alzada samples did not produce any PCR product of the expected size with  
220 the two sets of primers cited above and other primers were used (see Supplementary Table 1). The  
221 primers were designed to cover the most part of the WMAI ORF region. Supplementary Figure 1  
222 shows the gene sequence of 0.28 WMAI and the sequences of all the primers used in this study to  
223 amplify the corresponding ORF. All the combinations of primers listed in Supplementary Table 1  
224 were used, but it was not possible to obtain any PCR product corresponding to the ORF of WMAI  
225 genes in the tetraploid emmer, turanicum and Alzada wheat samples.

226 A previous study (Wang et al., 2010) determined the monomeric  $\alpha$ -amylase inhibitor gene  
227 sequences of 14 populations of wild emmer wheat (*T. dicoccoides*), which is the wild progenitor of  
228 modern tetraploid and hexaploid wheat, however, it was not possible to retrieve any monomeric  $\alpha$ -  
229 amylase inhibitor gene sequence from *T. durum*, *T. turanicum* and *T. dicoccum* in the GenBank  
230 non-redundant DNA database (access performed on May 11<sup>st</sup>, 2021).

231 Regarding the available studies determining the amount of monomeric  $\alpha$ -amylase inhibitors in  
232 tetraploid wheat, the results were mixed. Rogniaux et al. (2015) using a targeted MS/MS approach  
233 were able to detect and quantify WMAI protein in *T. aestivum*, *T. durum* and *T. monococcum*, but  
234 not in KAMUT<sup>®</sup> khorasan wheat sample (*T. turgidum* ssp. *turanicum*). Geisslitz et al. (2020) used a  
235 new targeted LC-MS/MS method and found that emmer had similar contents of ATI 0.28 as  
236 common wheat and spelt but obtained mixed results for durum wheat with six out of eight cultivars  
237 showing a content near or even below LOD.

238 The lack of WMAI sequences in *T. durum*, *T. turanicum* and *T. dicoccum* from available databases  
239 together with the amplification of WMAI genes in *T. durum* cv. Peliss, but not in other tetraploid  
240 wheat in this study, and the mixed results regarding the presence of WMAI protein in tetraploid  
241 wheat confirmed that further studies are needed to deepen knowledge about WMAI in tetraploid  
242 wheat.

243 Regarding the other seven wheat genotypes (einkorn, Peliss, spelt, Turkey Red, Judee, Marquis and  
244 Vida) the DNA sequences from five clones per each wheat genotype were determined and used to

deduce the full amino acid sequence of the proteins. As described before (Wang et al., 2008), there was no intron in the monomeric  $\alpha$ -amylase inhibitor sequences. So, it was possible to isolate the complete coding sequences of monomeric  $\alpha$ -amylase inhibitor gene by direct PCR amplification. In this study, no ins/del in the coding region of WMAI were found in any of the wheat samples.

From a total of 35 deduced amino acid sequences, 16 haplotypes were identified (HM1-HM16). Haplotype HM7 was shared by all the hexaploid wheat samples (it occurred in 13 out of 25 hexaploid wheat sequences) and it turned out to be the most abundant WMAI haplotype. No pseudogenes were found (Table 3).

All the putatively functional genes encoded for a 151 amino acid protein (30 amino acid signal peptide and a 121 amino acid mature protein).

In this study, after aligning the 16 monomeric  $\alpha$ -amylase inhibitor haplotypes from 7 wheat genotypes and WMAI 0.28 (GenBank: AJ223492), 18 nsSNPs were identified. The frequency of nsSNPs was 1 out of 25.2 bases.

The alignment of the 16 monomeric  $\alpha$ -amylase inhibitor haplotypes and the position of the nsSNPs were shown in Supplementary Figure 2.

Sequence alignment showed that all the monomeric  $\alpha$ -amylase inhibitors were highly homologous and are part of a monomeric  $\alpha$ -amylase inhibitor family including WMAI 0.28 and suggested that these inhibitors might have derived from a very limited number of ancestral genes (Wang et al., 2008). Haplotype HM7, which is the most abundant WMAI haplotype, had the same sequence as WMAI 0.28; the other haplotypes showed a percentage of identity with WMAI 0.28 ranging from 96.7% to 99.3%.

All the deduced proteins of monomeric  $\alpha$ -amylase inhibitors had 10 Cys residues which form five disulphide bonds. The disulphide bonds are essential for the inhibitory activity (Carbonero and García-Olmedo, 1999) and in fact the Cys residues were at conserved positions.

Monomeric  $\alpha$ -amylase inhibitors are highly active against  $\alpha$ -amylase of *Tenebrio molitor* and show a low inhibitory activity against mammalian and some avian  $\alpha$ -amylases. The 3-D structure of the complex between  $\alpha$ -amylase from *T. molitor* and 0.28  $\alpha$ -amylase inhibitor has been determined (Payan, 2004) and three regions of contact have been identified in the mature protein: the N-terminal segment (residues 1-10); residue 53 (inside the second loop segment) and sequence including residues 103-119, which are part of the fourth loop (shown as residues 31-40, 83 and 133-149 respectively in Supplementary Figure 2). In particular, the last region, which corresponds to the

276 C-terminal segment of 0.28, plays an important role in filling the central substrate-binding subsites  
277 of *T. molitor*  $\alpha$ -amylase and in targeting its catalytic residues. Moreover, this region plays an  
278 important role in the specificity of 0.28 inhibitor. Most of the haplotypes shared the same sequence  
279 as WMAI 0.28 in these regions, with the following exceptions: HM1, HM9, HM10 and HM13.

280 Other two regions critical for the inhibitory activity have been identified in the mature protein: the  
281 N-terminal sequence (positions 1-6) and the sequence after the CRC motif (positions 57-59) (Liu  
282 and Wang, 2012). Only haplotype HM1 (found in einkorn) displayed a change at position 6 (shown  
283 as residue 36 in Supplementary Figure 2).

284 The current literature lacks information on the peptide sequences and epitopes responsible for the  
285 allergies triggered by wheat albumin and globulin proteins (Tatham and Shewry, 2008). Regarding  
286 ATIs, there is only one study which identified a high IgE-binding region at the position 9 to 26 in  
287 the mature WMAI protein (Walsh and Howden, 1989). As shown as residues 39-56 in  
288 Supplementary Figure 2, only two haplotypes had an amino acid change at this level: HM5 and  
289 HM14.

290 Whether the sequence changes found in this study and cited above are able to reduce or even delete  
291 the inhibitory activity and/or the immunogenic properties is not possible to say from these data.

### 292 **3.2 Sequence analysis of dimeric alpha-amylase inhibitors (WDAI)**

293 The primers WDAIfor and WDAIrev were used to amplify the ORF of the dimeric alpha-amylase  
294 genes (WDAI). All the ten wheat genotypes gave the PCR product of the expected size (~509 bp).

295 From a total of 50 deduced amino acid sequences, 35 haplotypes were identified (HD1-HD35),  
296 among which 6 haplotypes represented pseudogenes (indicated with a p after the number), due to  
297 the presence of one or more in-frame stop codons because of a single nucleotide polymorphism  
298 (substitution, insertion or deletion), although we cannot predict from the genomic data whether  
299 these sequences are being expressed (Table 3). All the sequences obtained from the einkorn sample  
300 showed an insertion of C at position 160 (from the start codon ATG) which resulted in a premature  
301 stop codon and the impossibility to synthesize the correct mature protein. The same insertion of C  
302 was observed in the DNA sequences from all the *T. monococcum* accessions and from above all *T.*  
303 *boeoticum* accessions in a previous study (Wang, 2007). The absence of gene sequences coding for  
304 functional WDAI genes in *T. monococcum* is in agreement with functional studies using targeted  
305 MS/MS approach to detect WDAI proteins (Geisslitz et al., 2020, 2018; Rogniaux et al., 2015).

306 All the putatively functional genes were 426 bp long, encoding for a 141 amino acids protein (17  
307 amino acids signal peptide and 124 amino acids mature protein). Haplotype HD18 was found in 6  
308 clones from 5 different wheat genotypes and it was the most abundant.

309 In this study, after aligning the 35 dimeric  $\alpha$ -amylase inhibitor haplotypes from ten wheat  
310 genotypes, 69 nsSNPs were identified (Supplementary Figure 3).

311 The frequency of nsSNPs was 1 out of 6.2 bases. The WDAI gene sequences had more divergence  
312 than the WMAI gene sequences of this study, which is in agreement with previous studies (Liu and  
313 Wang, 2012).

314 All the deduced proteins of WDAI but HD1 and ATI 0.53 had 10 Cys residues which formed five  
315 disulphide bonds and were at conserved positions.

316 Studying the modelled complex of human salivary  $\alpha$ -amylase (HSA) with 0.19 inhibitor, three  
317 spots important for the inhibitory activity of WDAI have been highlighted (Payan, 2004): His47;  
318 Ser49 and the sequence Val104 – Val105 – Asp106 – Ala107. Amino acids changes at the level of  
319 these three spots can result in a different ability to inhibit human  $\alpha$ -amylases. As shown as residues  
320 64, 66 and 121-124 respectively in Supplementary Figure 3, the only haplotypes displaying all the  
321 above-mentioned amino acids residues at the three inhibitor spots were HD5, HD9, HD11, HD15  
322 and HD17. Interestingly, all these haplotypes were found only in hexaploid wheat genotypes, which  
323 showed in fact the highest  $\alpha$ -amylase inhibitory activity *in vitro* (data not published yet).

324 Considering the percentage of identity with the two amino acid sequences of the well-studied  
325 dimeric  $\alpha$ -amylase inhibitors 0.19 and 0.53, only the haplotype HD9 had a 100% of identity with  
326 0.19. However, the level of homology was high with both 0.19 and 0.53 (ranged from 87.1% to  
327 100% for 0.19 and from 84.7% to 99.2% for 0.53), with the exception of the pseudogene HD29p.

### 328 **3.3 Sequence analysis of tetrameric alpha-amylase inhibitors CM3 (WTAI-CM3)**

329 WTAI-CM3 was selected as representative of WTAI genes in this study because of its extended  
330 interaction with  $\alpha$ -amylases compared to the other CM subunits (Capocchi et al., 2013) and because  
331 it was the most abundant expressed WTAI proteins in cv. Butte 86 (Altenbach, 2011).

332 The sequence of CM3 encoding gene revealed that there is only one ORF encoding the entire  
333 protein (Singh et al., 2012).

334 The primers CM3for and CM3rev were designed *in house* and were used to amplify the ORFs of  
335 the tetrameric  $\alpha$ -amylase gene CM3 and all the samples but einkorn gave the PCR product of the

336 expected size (~ 647 bp). Two sets of internal primers were designed *in house* and were used in all  
337 the possible combinations with the einkorn sample, but no band of the expected size was amplified.  
338 Moreover, using the sequence of inhibitor WTAI-CM3 (NCBI: X17574) a blast search against the  
339 GenBank non-redundant DNA database was performed (access performed on May 11<sup>th</sup>, 2021), but  
340 it was not possible to retrieve any WTAI-CM3 sequence from *T. monococcum*.

341 Some studies were previously performed to quantify allergens abundance in different wheat  
342 genotypes and raised doubts on the presence of WTAI-CM3 proteins in diploid wheats. Rogniaux et  
343 al. (2015) detected WTAI-CM3 proteins only at very low levels in two *T. monococcum* (cv.  
344 Engrain, cv. DV92). Geisslitz et al. (2020, 2018) detected WTAI-CM3 in only four out of eight  
345 einkorn cultivars, but in very low concentrations near the LOQ or even below.

346 From a total of 45 deduced amino acid sequences derived from 9 wheat genotypes, 25 haplotypes  
347 were obtained (HC1-HC25), among which one haplotype was identified as a pseudogene (HC20p)  
348 due to the presence of one in-frame stop codon (table 3). As described before (Singh et al, 2012)  
349 there was no intron in the WTAI-CM3 sequences. In this study, no ins/del in the coding region of  
350 WTAI-CM3 was found in any of the wheat samples.

351 All the putatively functional genes encoded for a 168 amino acid protein (25 amino acid signal  
352 peptide and 143 amino acid mature protein).

353 In this study, after aligning the 25 WTAI-CM3 haplotypes from 9 wheat genotypes, 34 nsSNPs  
354 were identified. The frequency of nsSNPs was 1 out of 14.8 bases. Previously, Liu et al. (2012)  
355 stated that WTAI genes (CM2, CM3, CM16) are more conserved than WMAI and WDAI genes  
356 probably because these subunits are combined to form the tetrameric inhibitor and an amino acid  
357 change in one of them can affect the structure and consequently the resulting inhibitory activity.  
358 However, in this study results showed that WTAI-CM3 was more conserved than WDAI genes  
359 where it was found a snSNPs frequency of 1 out of 6.2 bases, but less conserved than WMAI genes  
360 where the snSNP frequency was 1 out of 25.2 bases.

361 The alignment of the 25 WTAI-CM3 haplotypes and the position of the nsSNPs were shown in  
362 Supplementary Figure 4.

363 Haplotype HC6, which was the most abundant WTAI-CM3 haplotype, had the same sequence as  
364 WTAI-CM3 (X17574) studied by García-Maroto et al. (1990), the other haplotypes showed a high  
365 percentage of identity with WTAI-CM3 (X17574) ranging from 92.9% to 99.4%.

366 The most part of the deduced proteins of WTAI-CM3 had 10 Cys residues which form five  
367 disulphide bonds and are at conserved positions. The exceptions were HC3 (Tyr instead of the



368 second Cys of the CRC motif); HC9 (Tyr at position 148); HC11 (Arg at position 29); HC18 (Tyr at  
369 position 52).

370 The sequence DLPGCPRE (amino acid positions 101-108 of the mature protein, shown as residues  
371 126-133 in Supplementary Fig. 4) is the most conserved in CM proteins and may be the  
372 characteristic sequence of alpha amylase/ trypsin - bifunctional inhibitors. Most of the haplotypes  
373 shared this conserved sequence, with the following exceptions: HC14, HC3, HC4, HC7, HC8,  
374 HC21 and HC25 (Supplementary Fig. 4).

375 Capocchi et al. (2013) proposed a structural model for the emmer tetrameric  $\alpha$ -amylase inhibitor  
376 and showed that two sequences Phe30-Lys55 and Lys116-Gln122 of the mature protein (shown as  
377 residues 55-80 and 141-148 respectively in Supplementary Fig. 4) are important for the inhibitory  
378 activity against *T. molitor*  $\alpha$ -amylase. In this study four amino acid changes occurred in some  
379 haplotypes at level of the first sequence: Thr instead of Met at amino acid position 65 occurred in  
380 HC16; Val instead of Ala at amino acid position 68 occurred in HC1; Phe instead of Tyr at amino  
381 acid position 71 and Met instead of Gly at position 74 both occurred in HC3, HC7, HC8, HC21,  
382 HC25 (Supplementary Fig. 4). Interestingly, the haplotypes which had the last two amino acid  
383 changes also had changes at the conserved sequence DLPGCPRE and all these were found only in  
384 hexaploid wheat genotypes.

385 ATI proteins from wheat have been demonstrated to trigger inflammation by eliciting strong innate  
386 immune effects *in vitro* and *in vivo* with the activation of the TLR4–MD2–CD14 complex (Junker  
387 et al., 2012). Cuccioloni et al. (2016) investigated the interaction between WTAI-CM3 and human  
388 TLR4 and predicted the ATI-TLR4 binding interface regions. These sequences are residues 33-44  
389 and 90-100 in the mature protein sequence (shown as residues 58-69 and 115-125 respectively in  
390 Supplementary Fig. 4). The haplotypes which showed one or two amino acid changes within these  
391 regions were: HC1, HC5, HC8, HC15 and HC16 (Supplementary Figure 4). From the existing  
392 evidence it is not possible to say if these changes are able to decrease or delete the ability of CM3-  
393 WTAI to bind human TLR4.

394 Several studies suggested that the apparent absence of alpha-amylase inhibitory activity of *T.*  
395 *monococcum* against mammalian enzymes could be explained by the fact that the corresponding  
396 coding genes might be expressed at very low level or even silenced, perhaps because of gene  
397 mutations that prevent the translation into the mature protein (García-Maroto et al., 1990). This  
398 hypothesis is in agreement with the gene sequencing results of this study. In fact, einkorn did not  
399 show any WTAI-CM3 gene sequence and so it should not be able to produce any tetrameric alpha-  
400 amylase inhibitor proteins. Moreover, all the WDAI gene sequences showed an insertion of C at

position 160 which resulted in a premature stop codon and the impossibility to synthesize the correct WDAI mature protein. Lastly, einkorn showed WMAI sequences, but it is known that WMAI proteins are highly active against alpha-amylase of *T. molitor* and only weakly inhibits the alpha-amylases from human saliva and pancreas (Payan, 2004). Notwithstanding low levels of ATI proteins and a low alpha-amylase inhibitory activity, Call et al. (2020) found that einkorn samples have high trypsin inhibitory activities and they hypothesized that it could be due to the fact that besides ATIs also lipid transfer proteins (LTPs), grain softness proteins and beta-amylases show high affinity to trypsin.

### 3.4 Sequence analysis of trypsin inhibitors CMx

The primers CMxfor and CMxrev were designed *in house* and were used to amplify the ORFs of the CMx genes and all the ten wheat genotypes gave the PCR product of the expected size (~ 498 bp). Since CMxfor was designed 9 nucleotides downstream the A of the start codon, these first 9 nucleotides were taken from the reference sequence (GenBank: X75608) and added to each DNA sequence in order to deduce complete protein sequences for subsequent analyses.

From a total of 50 deduced amino acid sequences, 22 haplotypes were obtained (HX1-HX22), among which two haplotypes were identified as pseudogenes due to the presence of one in-frame stop codon (Table 3).

HX20 occurred in 15 out of 50 CMx sequences and it turned out to be the most abundant CMx haplotype.

The deduced proteins from haplotypes HX1-HX10 were 145 amino acids long and were found only in hexaploid wheat genotypes and the deduced proteins from haplotypes HX13-HX20 were 146 amino acids long. Differently, two nucleotide sequences showed a premature stop codon after nucleotide 363 as observed by Sanchez de la Hoz et al. (1994), which resulted in a deduced protein of 121 amino acids (haplotypes HX11 and HX12). Interestingly, emmer wheat showed only the two haplotypes with the deduced protein of 121 amino acids. The homology between the deduced HX11 and HX12 amino acid sequences and those of the other CMx haplotypes was maintained beyond the premature stop codon, up to the second stop codon which appeared in the same position as the other haplotypes. All the deduced proteins showed a 25 amino acid signal peptide.

In this study, after aligning the 22 CMx haplotypes from 10 wheat genotypes, 39 nsSNPs were identified (Supplementary Figure 5). The frequency of nsSNPs was 1 out of 11.2 bases, similar to what was found for CM3 genes in this study.

The most part of the CMx deduced proteins had 10 Cys residues which form up to five disulphide

bonds and were at conserved positions. The exceptions were HX3, HX4, HX9, HX11 HX12 and HX17. Interestingly, all the haplotypes found in emmer in this study didn't show all the 10 Cys residues.

### 3.5 Phylogenetic analysis

The Neighbor-Joining method was used to calculate the phylogenetic distances and to construct the phylogenetic tree among all the haplotypes for each gene (Figure 1).

For the 16 WMAI haplotypes the optimal tree with the sum of branch length 0.12061355 was shown (Figure 1A). Cluster III, which displayed all the haplotypes of the tetraploid Peliss and the diploid einkorn, diverged from cluster I, cluster II and HM8, the latter displaying only hexaploid wheat genotypes (in particular cluster II and HM8 displayed only haplotypes from spelt). So, the clusters clearly separated the wheat genotypes according to their ploidy.

For the 35 WDAI haplotypes the optimal tree with the sum of branch length 0.96668030 was shown (Figure 1B). HD29p was the most divergent and was highly separated from the other haplotypes. The other pseudogenes from einkorn (HD28p, HD34p, HD35p) diverged together in a separated group within cluster III. Each cluster showed haplotypes from both tetraploid and hexaploid wheat genotypes and from both modern and ancient wheat genotypes, so it was not possible to make any differentiation based on WDAI gene sequences.

For the 25 WTAI-CM3 haplotypes the optimal tree with the sum of branch length 0.20342301 was shown (Figure 1C). Two clusters were identified with cluster II displaying only haplotypes from hexaploid wheat genotypes (both modern and ancient genotypes).

For the 22 CMx haplotypes the optimal tree with the sum of branch length 0.27715519 was shown (Figure 1D). A first subdivision was between haplotypes HX1-HX10 (which were all 145 amino acids long and found only in hexaploid wheat genotypes) and HX11-HX22p. Then a further subdivision distinguished HX9 and HX10 (cluster IV) from the other haplotypes of the first group (cluster V), while in the second group three clusters were identified: cluster I with haplotypes found in both tetraploid and hexaploid genotypes, cluster II showing the two 121 amino acids long CMx proteins, cluster III showing only haplotypes found in einkorn.

According to the phylogenetic analysis of the haplotypes of all the four genes analysed, it was not possible to differentiate the wheat genotypes with respect to ancient or recently developed wheat genotypes.

### 463 3.6 Principal Component Analysis (PCA) of WMAI, WDAI, WTAI-CM3 and CMx sequences

464 A PCA analysis was performed based on deduced amino acid sequences of all the four genes  
465 sequenced in this study (WMAI, WDAI, WTAI-CM3 and CMx).

466 To visualise relationships between the tested variables, principal component analysis was run for  
467 each species in two steps. First, an interim PCA was run including all the variables (160 nsSNPs),  
468 and the quality of representation (Cos2) of each variable was checked. Second, a definitive PCA  
469 was run with those variables with a quality of representation higher than 0.5 (103 nsSNPs).

470 The scatter plot reported the projection of cases (10 wheat genotypes) on the first two components  
471 PC1 and PC2 and explained the 63.6% of total variance with the first PC accounting for 40.9% and  
472 the second PC for 22.7% (Figure 2). Einkorn diverged from all the other genotypes according to the  
473 projection of the first component (PC1 axis). The tetraploid wheat genotypes can be divided from  
474 the hexaploid wheat genotypes according to the second projection (PC2 axis) with the tetraploid  
475 Peliss which was closer to the hexaploid group. This fact can be explained in part by the fact that  
476 Peliss was the only tetraploid wheat which was amplified with primers for WMAI genes. It was not  
477 possible to differentiate the wheat genotypes with respect to ancient or recently developed wheat  
478 genotypes.

### 479 3.7 Haplotype analysis

480 The Nei's genetic distances were calculated for paired comparisons of the ten wheat genotypes  
481 (Supplementary Table 2). The values of genetic distances varied from 0.0085 to 0.0321 with an  
482 average of 0.022. The genetic distance between turanicum and Alzada was the lowest (0.0085)  
483 whereas those between Peliss and both spelt and Vida were the largest.

484 Figure 3 showed the dendrogram based on the Nei's genetic distances. First, einkorn diverged from  
485 the other genotypes and this confirmed the results of the PCA analysis and this is in line with the  
486 literature which affirmed that *T. monococcum* did not contribute to the evolution of both tetraploid  
487 species of *Triticum* and *T. aestivum* through polyploidization (Zhao et al., 2021).

488 Then, the group of the naked tetraploid wheat genotypes diverged from the others (in this group  
489 turanicum and Alzada are closer to each other than Peliss probably because they lacked WMAI  
490 sequences, unlike Peliss). The hulled tetra/hexaploid wheat genotypes diverged from the naked  
491 hexaploid wheat genotypes. In this last group it is interesting to note that the heritage genotypes  
492 (Turkey Red and Marquis) were closer than the modern genotypes (Judee and Vida). It is also  
493 interesting to highlight that emmer was close to spelt and this is in line with recent studies based on

494    HMW glutenin analysis which showed that the European spelt is a result of hybridization between  
495    free-threshing hexaploid wheat and domesticated emmer (Blatter et al., 2004).

#### 496    **4. Conclusions**

497    In this study, the sequences of four representative alpha-amylase/trypsin inhibitor genes have been  
498    determined in ten wheat genotypes with different ploidy level and year of release, and some of  
499    these, for example *T. monococcum* and *T. turgidum* spp. *turanicum*, have never been sequenced for  
500    some or all ATI genes before. So, this study has expanded the available information about ATI  
501    sequences.

502    Considering the deduced amino acid sequences of all the four ATI genes studied, both the  
503    phylogenetic analysis and the PCA analysis evidenced that the ten wheat genotypes can be  
504    differentiated on the basis of their ploidy level. This is in agreement with the fact that ATI genes are  
505    present in different chromosomal sets. Overall, einkorn was the most divergent genotype.

506    Based on both the phylogenetic analysis and the PCA analysis of the haplotypes of the four genes  
507    analysed, it was not possible to differentiate the wheat genotypes with respect to ancient or recently  
508    developed wheat genotypes. Further studies with a higher number of genotypes per each ploidy  
509    level are needed to confirm these results. The haplotype analysis, beside confirming these results,  
510    also allowed the separation of the hulled from the naked wheat genotypes.

511    The literature lacks information on which sequences of ATIs have immunogenic and inflammatory  
512    potential (Geisslitz et al., 2021), so it was not possible to make this kind of consideration on the  
513    basis of the deduced amino acid sequences. However, the genetic sequence differences highlighted  
514    in this study among the ten genotypes can be the basis for further studies aimed at identifying those  
515    sequences in ATI genes able to trigger innate and adaptive immune response. Moreover, the  
516    sequence variants of ATIs can be useful information for future breeding programs aimed at  
517    selecting wheat varieties with potential pest resistance and minimum inflammatory effect on human  
518    health.

519    Finally, it is desirable to measure the inhibitory activities against both alpha-amylase and trypsin  
520    enzymes to find possible differences among the ten wheat genotypes and to eventually find a  
521    correlation with their genetic sequences.

522    To the best of our knowledge, this is the first study analyzing the ATI sequences of a set of ancient  
523    and modern wheat genotypes with a different ploidy level.

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## Tables

**Table 1**

List of wheat genotypes used in this study.

<i>Triticum</i> species	Genotypes	Genome formula	Year of release	Typology	Pedigree	Provenance	Name used in this study
<i>T. monococcum</i> L. subsp. <i>monococcum</i>	Local landrace	AA	/	Hulled, ancient wheat	N.D.	Montana State University (Montana, USA)	Einkorn
<i>T. turgidum</i> L. subsp. <i>dicoccum</i>	Local landrace	AABB	/	Hulled, ancient wheat	N.D.	Montana State University (Montana, USA)	Emmer
<i>T. turgidum</i> L. subsp. <i>turanicum</i> *	QK-77	AABB	/	Naked, ancient wheat	N.D.	Kamut International (Montana, USA)	Turanicum
<i>T. turgidum</i> L. subsp. <i>durum</i>	Peliss	AABB	1900	Naked, heritage durum wheat	LV-Oran	University of Saskatchewan (Canada)	Peliss
<i>T. turgidum</i> L. subsp. <i>durum</i>	Alzada	AABB	2004	Naked, modern durum wheat	Mohawk/Kofa	Montana State University (Montana, USA)	Alzada
<i>T. aestivum</i> L. subsp. <i>spelta</i>	Local landrace	AABBDD	/	Hulled, ancient wheat	N.D.	Montana State University (Montana, USA)	Spelt
<i>T. aestivum</i> L. subsp. <i>aestivum</i>	Turkey Red	AABBDD	1873	Naked, heritage hard red common wheat	(S)Crimean	Hartland Mills (Kansas, USA)	Turkey Red
<i>T. aestivum</i> L. subsp. <i>aestivum</i>	Judee	AABBDD	2011	Naked, modern hard red common wheat	93-X-312-E-14/NuHorizon	Montana State University (Montana, USA)	Judee
<i>T. aestivum</i> L. subsp. <i>aestivum</i>	Marquis	AABBDD	1913	Naked, heritage hard red common wheat	Hard-Red-Calcutta/Red-Fife	Montana State University (Montana, USA)	Marquis
<i>T. aestivum</i> L. subsp. <i>aestivum</i>	Vida	AABBDD	2006	Naked, modern hard red common wheat	Scholar/Reeder	Montana State University (Montana, USA)	Vida

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**Table 2**

List of primers used in this study.

Gene	Name	Sequence
WMAI (NCBI GenBank: AJ223492)	Mfor1	5' – CACCACTTATATCCAAGGACCA – 3'
	Mfor2	5' – AGCTTGCTTTGATTCTGCTGAT – 3'
	Mfor3	5' – ACTAAATTGAAACAATGTGGAT – 3'
	F2	5' – ATGTGGATGAAGACCGKGT – 3'
	F4	5' – ATGCTCGTGGCGACAACAAT – 3'
	F3	5' – ACAACAATGGCGGTTCGAGTA – 3'
	F1	5' – CATAACAGTGGTCCTTGGAGT – 3'
	Mfor4	5' – GCAATGGTGAAGCTCCAGT – 3'
	Mfor5	5' – AGCTGGCCGACATCAACA – 3'
	Mrev1	5' – TCTCTGAGAGGACACATACACCA – 3'
	R1	5' – CACGCACCGCACCATTACTT – 3'
	Mrev2	5' – CACCGCACCAATTAAGAT – 3'
	Mrev3	5' – CGCACCAATTAAGATGCAGA – 3'
	Mrev4	5' – TTAAGATGCAGATTCGCTTGAC – 3'
	R2	5' – GACTAGRYGTCCGKATACGC – 3'
	Mrev5	5' – GATGGGCACCTTGCAGAC – 3'
	Mrev6	5' – GCACCTCCTTCCCCTCAC – 3'
WDAI (NCBI GenBank: AK330823)	WDAIfor	5' – CTATGTATGCTCGTGGCGAC – 3'
	WDAIrev	5' – ACTCATTYGCTTGACTAGGC – 3'
WTAI-CM3 (NCBI GenBank: X17574)	CM3for	5' – CGAACCAGACTTGGCTAGAATA – 3'
	CM3intfor2	5' – ACAACAACTTGTGGCACCT – 3'
	CM3intfor	5' – GCGCTGCGCTACTTCATA – 3'
	CM3rev	5' – ATTCATAGCAGATAGCCCACAC – 3'
	CM3intrev2	5' – TGTGAATGGTCGCCAAGT – 3'
	CM3intrev	5' – GCCGCTCTCACCAACAT – 3'
CMx (NCBI GenBank: X75608.1)	CMxfor	5' – AAGCACCAGCTCATCCTCTC – 3'
	CMxrev	5' – ATACACATATGCGATTTCGTCCA – 3'

**Table 3**

Distribution of WMAI, WDAI, WTAI-CM3 and CMx haplotypes in the ten wheat genotypes. The numbers under parenthesis indicate how many times the haplotype occurred in the 5 sequences analysed per each wheat genotype.

<b>GENOTYPES</b>	<b>WMAI HAPLOTYPES</b>	<b>WDAI HAPLOTYPES</b>	<b>WTAI-CM3 HAPLOTYPES</b>	<b>CM<sub>x</sub> HAPLOTYPES</b>
Einkorn	HM1 (1), HM2 (1), HM3 (3)	HD28p (2), HD29p (1), HD34p (1), HD35p (1)	/	HX13 (3), HX14 (1), HX21p (1)
Emmer	/	HD8 (1), HD18 (1), HD26 (2), HD27 (1)	HC6 (3), HC16 (1), HC17 (1)	HX11 (4), HX12 (1)
Turanicum	/	HD18 (2), HD20 (1), HD21 (1), HD22 (1)	HC6 (1), HC9 (1), HC10 (1), HC11 (1), HC12 (1)	HX17 (1), HX18 (1), HX20 (3)
Peliss	HM3 (3), HM4 (1), HM5 (1)	HD1 (1), HD2 (1), HD4 (1), HD18 (1), HD32 (1)	HC6 (4), HC18 (1)	HX20 (5)
Alzada	/	HD8 (1), HD18 (1), HD23 (1), HD24 (1), HD25 (1)	HC6 (2), HC13 (1), HC14 (1), HC15 (1)	HX19 (1), HX20 (4)
Spelt	HM6 (1), HM7 (1), HM8 (1), HM9 (1), HM10 (1)	HD15 (2), HD31 (1), HD30p (2)	HC6 (2), HC7 (1), HC22 (1), HC23 (1)	HX3 (1), HX8 (1), HX11 (3)
Turkey Red	HM7 (3), HM11 (1), HM12 (1)	HD5 (1), HD6 (1), HD7 (1), HD8 (1), HD9 (1)	HC6 (2), HC19 (1), HC25 (1), HC20p (1)	HX1 (1), HX5 (1), HX8 (2), HX9 (1)
Judee	HM7 (3), HM13 (1), HM14 (1)	HD9 (1), HD10 (1), HD11 (1), HD12 (1), HD33 (1)	HC1 (1), HC2 (1), HC3 (1), HC4 (1), HC5 (1)	HX6 (1), HX15 (1), HX16 (1), HX20 (2)
Marquis	HM7 (4), HM15 (1)	HD13 (1), HD14 (1), HD15 (2), HD16 (1)	HC6 (2), HC7 (1), HC21 (1), HC24 (1)	HX7 (1), HX8 (1), HX10 (1), HX20 (1), HX22p (1)
Vida	HM7 (2), HM15 (2), HM16 (1)	HD3 (1), HD9 (1), HD17 (1), HD18 (1), HD19p (1)	HC6 (2), HC7 (1), HC8 (2)	HX2 (1), HX4 (1), HX8 (3)

### Supplementary Table 1

Combinations of primer sets used to amplify WMAI genes. For each primer set the expected amplicon size (bp) and annealing temperature used (°C) are listed.

Reverse Forward	R1	R2	Mrev2	Mrev3	Mrev5	Mrev6	Mrev4	Mrev1
<b>F1</b>	445 bp 50°C	374 bp 62°C	404 bp 50°C	401 bp 50°C				
<b>F2</b>	529 bp 58°C	458 bp 50°C	488 bp 50°C	485 bp 56°C				
<b>F3</b>	478 bp 50°C	407 bp 56°C	437 bp 52°C	434 bp 55°C				
<b>F4</b>	490 bp 50°C	419 bp 50°C	449 bp 50°C	446 bp 55°C				
<b>Mfor3</b>	543 bp 54°C	472 bp 50°C	502 bp 50°C	502 bp 50°C				
<b>Mfor4</b>				329 bp 54°C	240 bp 54°C	169 bp 54°C		
<b>Mfor5</b>				262 bp 58°C	173 bp 54°C	102 bp 54°C		
<b>Mfor1</b>							550 bp 50°C	
<b>Mfor2</b>								623 bp 50°C

## Supplementary Table 2

Nei's genetic distance matrix of WMAI, WDAI, WTAI-CM3 and CMx haplotypes in the ten wheat genotypes.

Wheat genotype	Einkorn	Emmer	Turanicum	Peliss	Alzada	Spelt	Turkey Red	Judee	Marquis
Emmer	0.0274								
Turanicum	0.0224	0.0206							
Peliss	0.0296	0.0287	0.0167						
Alzada	0.0249	0.0205	<b>0.0085</b>	0.0124					
Spelt	0.0264	0.0133	0.0221	<b>0.0321</b>	0.0229				
Turkey Red	0.0256	0.0220	0.0212	0.0312	0.0212	0.0192			
Judee	0.0247	0.0272	0.0169	0.0286	0.0177	0.0237	0.0166		
Marquis	0.0281	0.0254	0.0212	0.0294	0.0211	0.0174	0.0130	0.0157	
Vida	0.0273	0.0237	0.0212	<b>0.0321</b>	0.0229	0.0201	0.0130	0.0210	0.0148



## Figure captions

### Figure 1.

Phylogenetic analysis of WMAI (A), WDAI (B), CMx (C) and WTAI-CM3 (D) haplotypes. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) was shown next to the branches. In red are indicated in which genotypes the haplotypes occurred. *T. turgidum* L. subsp. *durum* and *T. aestivum* L. subsp. *aestivum* genotypes are indicated with the abbreviations DU and AE respectively.

### Figure 2.

Principal component analysis based on deduced WMAI, WDAI, WTAI-CM3 and CMx amino acid sequences. The scatter plot reports the projection of cases (10 wheat genotypes) on the first two components PC1 and PC2 (accounting for 63.6% of total variability): the diameter of each balloon is proportional to the variability observed for the wheat genotype. Each colour of the balloons corresponds to a different wheat ploidy: green for diploid wheat; red for tetraploid wheat; blue for hexaploid wheat.

### Figure 3.

The dendrogram of the ten wheat genotypes based on Nei's genetic distance matrix.

### Supplementary Figure 1.

The gene sequence of monomeric  $\alpha$ -amylase inhibitor 0.28 (AJ223492) and the primers used to amplify WMAI genes. The sequences of signal peptide and mature protein are highlighted in red and blue respectively. The stop codon is marked with an asterisk.

### Supplementary Figure 2.

Multiple sequence alignment of 16 WMAI haplotypes obtained from seven wheat genotypes and WMAI 0.28 (GenBank: AJ223492). The sequences related to the signal peptide have been included (residues 1 – 30 highlighted with a light blue box). The dots indicate conserved residues and the letters correspond to the substituted amino acid residues for each alignment gap. Blue arrows highlight the positions of the 10 Cys residues. The amino acids at the three inhibitor spots (Payan et al., 2004) are highlighted with yellow boxes. The two regions critical for the inhibitory activity (Garcia-Maroto et al., 1991) are marked with red rectangles.

### Supplementary Figure 3.

Multiple sequence alignment of 35 WDAI haplotypes obtained from ten wheat genotypes with WDAI 0.19 (Uniprot: P01085) and 0.53 (Uniprot: P01084). The sequences related to the signal peptide have been included (residues 1 – 17 highlighted with a light blue box). The dots indicate conserved residues and the letters correspond to the substituted amino acid residues for each alignment gap. The pseudogenes have been included: in-frame stop codons are indicated with asterisks \*; insertions and deletions are indicated with blue and red triangles respectively and the sequences after these changes until the end of the ORF of the corresponding functional gene is shown. Blue arrows highlight the positions of the 10 Cys residues. The amino acids at the three inhibitor spots are highlighted with yellow boxes.

#### **Supplementary Figure 4.**

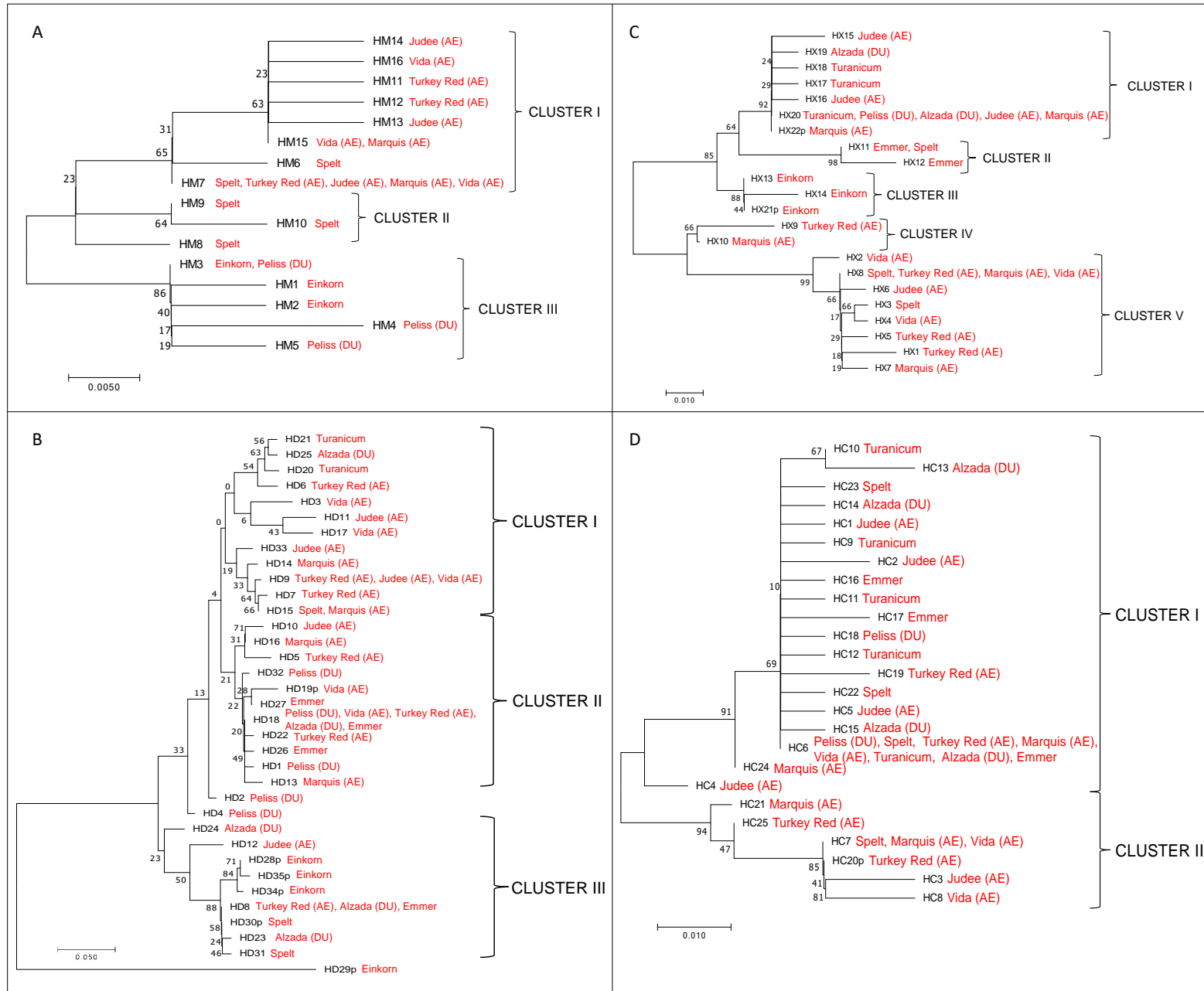
Multiple sequence alignment of 25 WTAI-CM3 haplotypes obtained from nine wheat genotypes and WTAI-CM3 (GenBank: X17574). The sequences related to the signal peptide have been included (residues 1 – 25 highlighted with a light blue box). The dots indicate conserved residues and the letters correspond to the substituted amino acid residues for each alignment gap. The asterisk \* shows in-frame stop codons of pseudogene. Blue arrows highlight the positions of the 10 Cys residues.

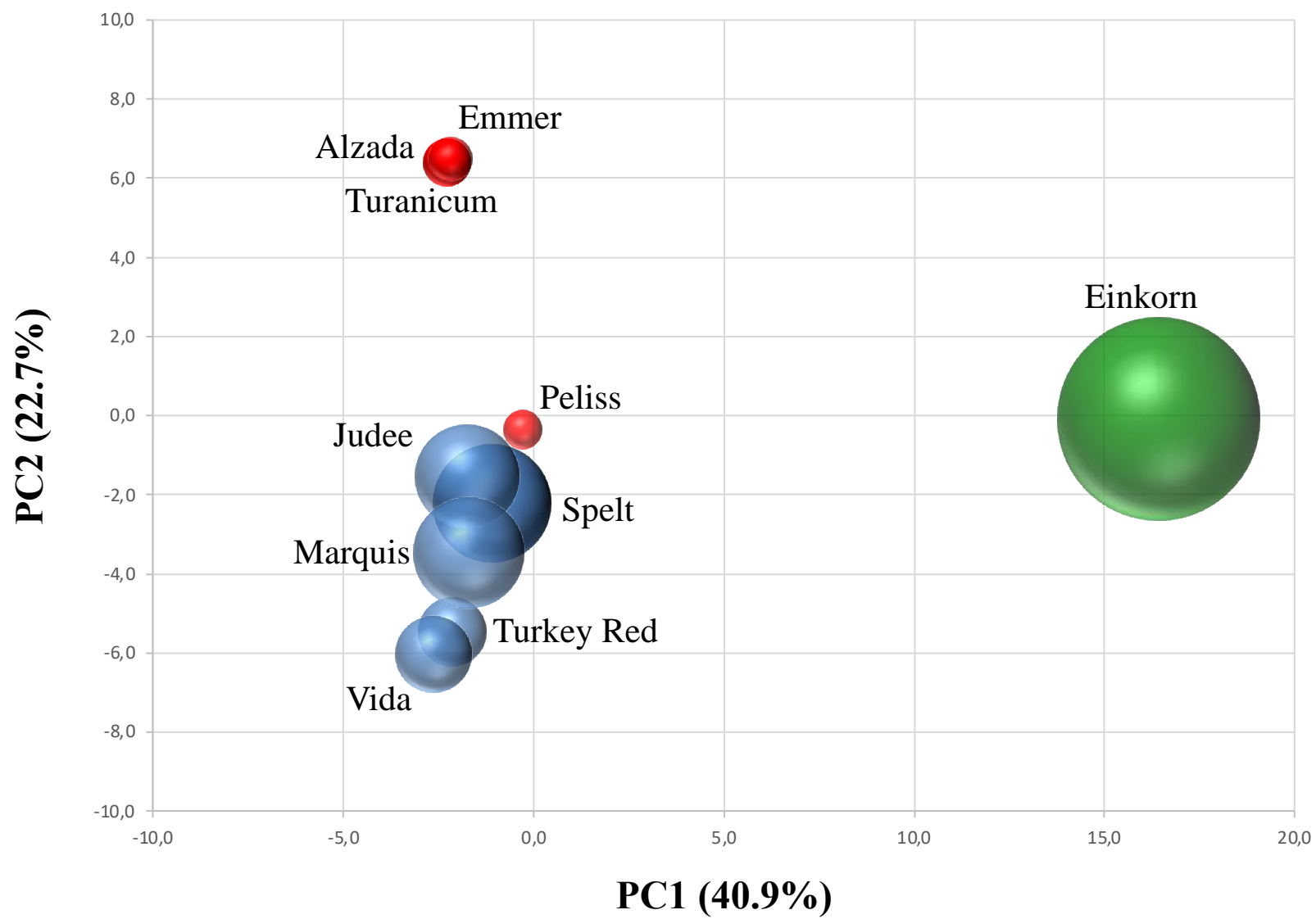
The conserved sequence DLPGCPRE is marked with a red rectangle, the amino acid residues of contact with TMA with yellow boxes and the ATI-TLR4 binding interface regions with blue rectangles.

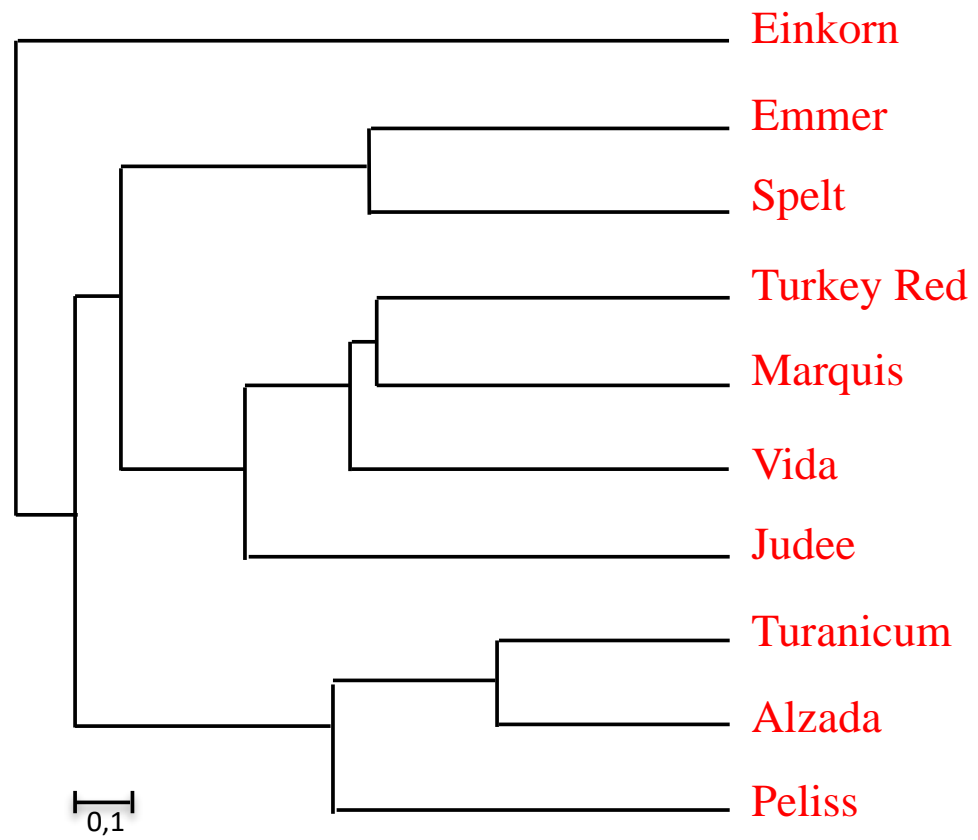
#### **Supplementary Figure 5.**

Multiple sequence alignment of 22 CMx haplotypes obtained from different wheat genotypes with CMx1/CMx3 (deduced from the sequence GenBank: X75608) and with CMx2 (deduced from the sequence GenBank: X75609). The sequences related to the signal peptide have been included (residues 1 – 24 highlighted with a light blue box).

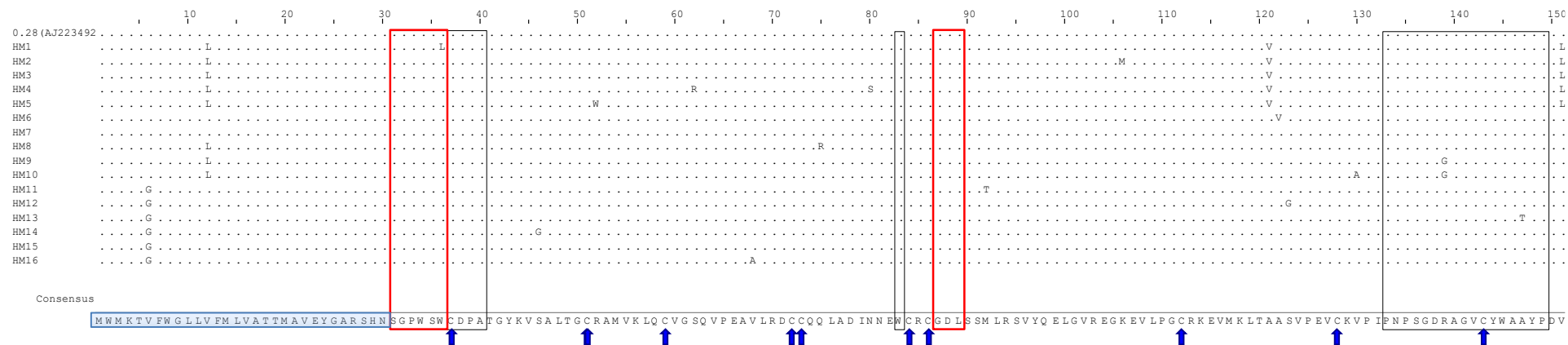
The asterisks \* show in-frame stop codons. The red triangle shows the positions of the deletion found in the nucleotide sequence used to deduce the amino acid sequence. The dots indicate conserved residues and the letters correspond to the substituted amino acid residues for each alignment gap. Blue arrows highlight the positions of the 10 Cys residues.







1 GTCACCACTTATATCCAAGGACCATGAGCGAGCTTGCTTTGATTCTGCTGATTAAGATAG  
 61 CACTAAATTGAAACA **ATGTGGATGAAGACCGTGTCTG**GGGGCTCCTAGTATTCATGCTC  
 M W M K T V F W G L L V F M L  
 121 **GTGGCGACAACAATGGCGGTCGAGTATGGTGCAAGGAGCCATAAC** **AGTGGTCCTGGAGT**  
 V A T T M A V E Y G A R S H N S G P W S  
 181 **TGGTGCATCCGGCGACGGGCTACAAGGTGAGCGCACTACGGGCTGCCGGGCAATGGTG**  
 W C D P A T G Y K V S A L T G C R A M V  
 241 **AAGCTCCAGTGTGTGGGCAGTCAGGTGCCCAGGCTGTCCTAAGAGATTGCTGCCAGCAG**  
 K L Q C V G S Q V P E A V L R D C C Q Q  
 301 **CTGGCCGACATCAACAACGAATGGTGCAGGTGCGGGGACCTCAGCAGCATGTTGCGTAGT**  
 L A D I N N E W C R C G D L S S M L R S  
 361 **GTTTATCAGGAGCTCGGCGTGCCTGAGGGGAAGGAGGTGCTCCAGGTTGCCGGAAGGAG**  
 V Y Q E L G V R E G K E V L P G C R K E  
 421 **GTGATGAAGCTCACGGCGGCGAGCGTGCCTGAGGTCTGCAAGGTGCCATCCCCAACCCG**  
 V M K L T A A S V P E V C K V P I P N P  
 481 **TCGGGAGACAGAGCAGGTGTCTGCTACTGGGCCGCTATCCGGACGCT** **TAG**TCAGCGAA  
 S G D R A G V C Y W A A Y P D V \*  
 541 **TCTGCACTTAATTGGTGCGGTGCGTACGCTCTGATAATAAATTAAGTAATGGTGCGGTG**  
 601 **CGTGCGCACAAAGATAAATAAAATTAAGTAATGGTGTATGTGCTCTCAGAGATCAACTC**  
 661 ATGAATAAATGTGAACTTGTTCTGTTGA



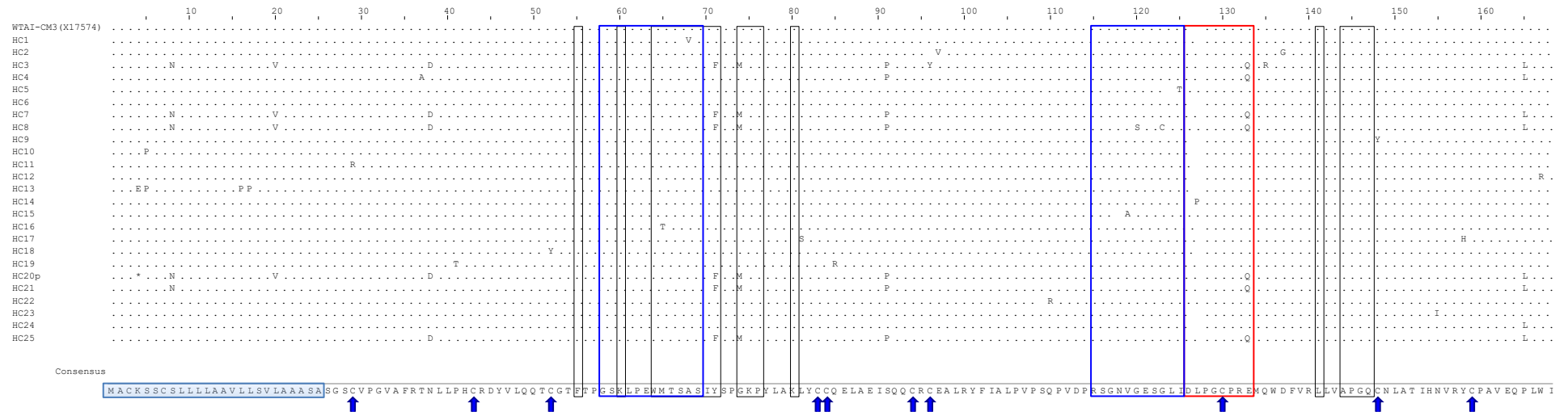
	10	20	30	40	50	60	70	80	90	100	110	120	130	140
HD1	.	S	.	G	.	.	.	.	.	V	S	.	.	.
HD2	.	.	V	.	K	.	.	.	.	.	V	S	.	.
HD3	.	.	.	.	.	.	.	.	.	.	.	E	.	.
HD4	.	.	.	K	.	.	D	.	.	.	.	.	T	.
HD5	.	.	.	.	.	.	.	.	.	.	.	.	K	.
HD6	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD7	.	.	.	.	.	.	V	.	.	.	.	.	.	.
HD8	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD9	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD10	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD11	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD12	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD13	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD14	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD15	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD16	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD17	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD18	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD19p	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD20	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD21	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD22	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD23	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD24	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD25	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD26	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD27	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD28p	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD29p	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD30p	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD31	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD32	.	.	.	S	.	G	.	.	.	.	.	.	.	.
HD33	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD34p	.	.	.	.	.	.	.	.	.	.	.	.	.	.
HD35p	.	.	.	.	.	.	.	.	.	.	.	.	.	.
0.19	.	.	.	.	.	.	.	.	.	.	.	.	.	.
0.53	.	.	.	.	.	.	.	.	.	.	.	.	.	.

Consensus

M L V A T P I A A E Y D A W S Y N S G P W M C Y P G Q A F Q V P A L P G C R P L L K L Q C N G S Q V P E A V L R D C C Q Q L A I S E W C R C G A L Y S M L D S M Y K E H G A Q E G Q A G T G A F P R C R R E V V K L T A A S I T A V C R L P I V V D A S G D G A Y V C K D V A A Y P D A







	10	20	30	40	50	60	70	80	90	100	110	120	130	140
CMX1/CMX3	.	I	V	.	EQ	.	R	.	W	.	R	.	.	.
CMX2	.	I	V	.	EQ	.	R	.	W	.	R	.	D	.
HX1	.	I	V	.	EQ	.	G	.	A	K	.	.	.	.
HX2	.	I	V	.	EQ	.	A	.	K	F	.	H	.	.
HX3	.	I	V	.	EQ	.	A	.	K	F	.	H	.	.
HX4	.	I	V	.	EQ	.	A	.	K	F	.	H	.	.
HX5	.	I	V	.	EQ	.	D	.	H	A	.	.	.	.
HX6	.	I	V	.	EQ	.	A	.	K	F	.	H	.	.
HX7	.	I	P	V	.	EQ	.	A	.	K	F	.	H	.
HX8	.	I	V	.	EQ	.	A	.	K	F	.	H	.	.
HX9	.	.	V	.	G	S	.	R	.	V	.	E	.	.
HX10	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX11	.	.	V	.	G	.	.	R	.	.	.	.	.	.
HX12	.	.	V	.	G	.	.	R	.	.	.	.	.	.
HX13	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX14	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX15	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX16	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX17	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX18	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX19	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX20	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX21p	.	.	.	.	G	.	.	R	.	.	.	.	.	.
HX22p	.	.	.	.	G	.	.	R	.	.	.	.	.	.

Consensus

MAFKHQ LILSTAVLLAALAAASAFDRRCVPGREITYESLNACREYAVRQTCGYLLSAERQKRCCDELSKVPCLRCCEVLRILMDGRVTKEGVVKGSLQEDMSRCKKLTREFIAGIVGREECNLETVFGRYHYCPSEY+GPEVVV

