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Cloning of the wheat Yr15 resistance gene sheds light on the plant tandem kinase-pseudokinase family

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Yellow rust, caused by *Puccinia striiformis* f. sp. *tritici* (*Pst*), is a devastating fungal disease threatening much of global wheat production. Race-specific resistance (*R*)-genes are used to control rust diseases, but the rapid emergence of virulent *Pst* races has prompted the search for a more durable resistance. Here, we report the cloning of *Yr15*, a broad-spectrum *R*-gene derived from wild emmer wheat, which encodes a putative kinase-pseudokinase protein, designated as wheat tandem kinase 1, comprising a unique *R*-gene structure in wheat. The existence of a similar gene architecture in 92 putative proteins across the plant kingdom, including the barley RPG1 and a candidate for *Ug8*, suggests that they are members of a distinct family of plant proteins, termed here tandem kinase-pseudokinases (TKPs). The presence of kinase-pseudokinase structure in both plant TKPs and the animal Janus kinases sheds light on the molecular evolution of immune responses across these two kingdoms.

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heat was domesticated in the Fertile Crescent over 10,000 years ago¹. Today, wheat is cultivated more extensively than any other food crop, covering more than 244 million ha and yielding more than 750 million tons annually (Food and Agriculture Organization Corporate Statistical Database (FAOSTAT)). It is a major component in human diet, providing starch, proteins, vitamins, dietary fiber, and phytochemicals².

Wheat crops are subjected to numerous types of biotic and abiotic stresses, with fungal diseases comprising one of the most serious threats to wheat production. Yellow (stripe) rust disease of wheat is caused by the fungus *Puccinia striiformis* f. sp. *tritici* (*Pst*). Due to the rapid evolution of the pathogen and development of new virulent races, severe yield losses occurred during recent decades in Europe and worldwide³. Given that 88% of wheat production is susceptible to yellow rust, more than five million tons of wheat harvest, valued at around one billion USD, are estimated to be lost annually due to this pathogen⁴. These epidemics demonstrate that disease management relying on single race-specific resistance genes has become ineffective. As an alternative, broad-spectrum resistance genes are much sought after by breeders, because such genes generally can provide robust protection against diseases, in particular against yellow rust.

The plant surveillance system can detect pathogen-secreted molecules and activate powerful defense responses, which often trigger local programmed cell death (PCD) at the site of attempted colonization, known as a hypersensitive response (HR)⁵. In cereals, three different resistance mechanisms have been identified: (i) surface-localized receptors that recognize microbial-associated or pathogen-associated molecular patterns (PAMPs) and activate PAMP-triggered immunity⁶; (ii) race-specific resistance conferred by intracellular immune receptors that recognize pathogen effector proteins and activate effector-triggered immunity⁶; (iii) broad-spectrum quantitative (partial) disease resistance via various molecular pathways⁷.

Although hundreds of disease resistance (R)-gene loci have been genetically mapped to wheat chromosomes, only a small number of yellow rust resistance genes (Yr) have been isolated so far (Yr10, Yr18/Lr34/Sr57, Yr36, Yr46/Lr67/Sr55, Yr5/YrSP, and Yr7)^{8,9}. Most of the cloned R-genes in wheat encode intracellular nucleotide-binding leucine-rich-repeat receptors (NLRs), which recognize pathogen-secreted effectors delivered into the host cytoplasm⁷. Nevertheless, among the cloned Yr genes only three are NLRs (Yr10, Yr5/YrSP, and Yr7), while the others represent different protein families, which act through unique mechanisms and result in broad-spectrum resistance to yellow rust $(Yr36)^{10}$, and even in multi-pathogen (partial) resistance against the three wheat rusts and powdery mildew (Yr18/Lr34/Sr57/Pm38, Yr46/ Lr67/Sr55/Pm39)⁷. Yr36 encodes a wheat kinase start 1 (WKS1) protein having a non-RD kinase domain fused to a START lipidbinding domain¹⁰. WKS1 contributes to cell death, restricting pathogen growth and sporulation. However, this response takes several days longer than typical HR responses, thereby resulting in partial resistance¹¹. Yr46/Lr67 is a hexose transporter involved in sugar uptake¹². Changes in two critical amino acids in the resistant allele of Yr46/Lr67 probably alter the hexose transport in infected leaves and may explain the reduced growth of multiple biotrophic pathogen species¹². Yr18/Lr34 encodes an ATPbinding cassette (ABC) transporter of a yet unknown substrate¹³.

The genetic bottlenecks associated with wheat polyploidization and domestication, as well as selections in agroecosystems, led to a decrease in wheat genetic diversity and an increase in its vulnerability to biotic and abiotic stresses. One of the ways to improve wheat tolerance to these stresses is to recruit the adaptive potential of the wild wheat germplasm. Wild emmer wheat (WEW; *Triticum turgidum* ssp. *dicoccoides*), the tetraploid

progenitor of hexaploid common wheat (*Triticum aestivum*), has valuable adaptive diversity to various diseases, including yellow rust^{1,14}

Yr15 is a yellow rust resistance gene discovered in the 1980s in WEW accession G25 (G25)¹⁵. Yr15 confers broad-spectrum resistance against a worldwide collection of more than 3000 genetically diverse Pst isolates, including modern races, such as 'Warrior' (race DK09/11), which is currently threatening wheat production^{8,16–18}. Although some Pst races were reported to be virulent on Yr15 in 2000s¹⁹, since 2003 no virulence on Yr15 was reported among new incoming Pst isolates (Global Rust Reference Center (GRRC)).

Previously, we had mapped *Yr15* on chromosome arm 1BS distal to the nucleolar organizer locus *Nor-B1* using restriction fragment length polymorphism and simple sequence repeat markers^{20,21}. The development of near isogenic lines (NILs) for *Yr15* in the common wheat Avocet background⁸ and the identification of DNA markers linked to this gene²¹ have accelerated the introgression of *Yr15* into a variety of pre-breeding and commercial materials, including cultivars, such as Clearwhite 515, Expresso, Patwin 515, and Seahawk, that are widely grown in western United States^{8,21}. In light of the above, *Yr15* will find use in wheat breeding in many other wheat-growing regions⁸.

Here, we describe the identification of the Yr15 gene sequence by map-based cloning and demonstrate its function in conferring resistance to yellow rust in durum and common wheats, by both ethyl methanesulfonate (EMS) mutagenesis of resistant Yr15 introgression lines (ILs) and transgenic complementation of susceptible varieties. We show that Yr15 encodes a protein composed of putative kinase and pseudokinase domains in tandem, which we name here as WTK1 (wheat tandem kinase 1). Orthologs, paralogs, and homologs of WTK1 were found in other cereal species. Furthermore, we have discovered similar tandem kinase-pseudokinase (TKP) structures in a wide range of plant taxa and suggest that they belong to the same protein family, designated here as TKP family.

Results

Variation in resistance response of Yr15 ILs. A set of Yr15 ILs inoculated with Pst isolate #5006 showed a range of resistance responses, probably due to differences in their genetic backgrounds (Supplementary Fig. 1a). Infection types (ITs) for these ILs ranged from 0 (no visible response, e.g., B70 (Yr15)) to 2 (large chlorotic blotches, e.g., Sel46 (Yr15)), with clear HR visible to the naked eye in most of the Yr15 ILs (Supplementary Fig. 1a).

Previous studies showed that although Yr15 conferred broadspectrum resistance against a worldwide collection of Pst isolates $^{16-18}$, a few isolates that appeared in Europe more than a decade ago (e.g., isolate DK92/02) showed virulence on Yr15 IL Avocet $+ Yr15^{19}$. Therefore, we have challenged a set of Yr15, Yr5, and Yr5/Yr15 ILs with Pst races known to be virulent on Yr15 (DK92/02) 19 or Yr5 (AU85569) 22 . Indeed, DK92/02 was virulent on Avocet + Yr15 (IT = 5-7), and AU85569 was virulent on Avocet + Yr5 (IT = 7), while both of these races were virulent (IT = 8-9) on Avocet S NIL (Supplementary Fig. 1b). However, pyramiding Yr15 with Yr5 in four different backgrounds (YecoraRojoYr5Yr15, PatwinYr5Yr15, SummitYr5Yr15, and DirkwinYr5Yr15) showed full protection against both virulent isolates (Supplementary Fig. 1b).

Mechanism of resistance associated with HR. Microscopic observations of the interactions between plant host cells and the invading pathogen showed the formation of fungal structures within leaf tissues, including substomatal vesicles, primary infection hyphae, haustoria mother cells (HMCs), and haustoria

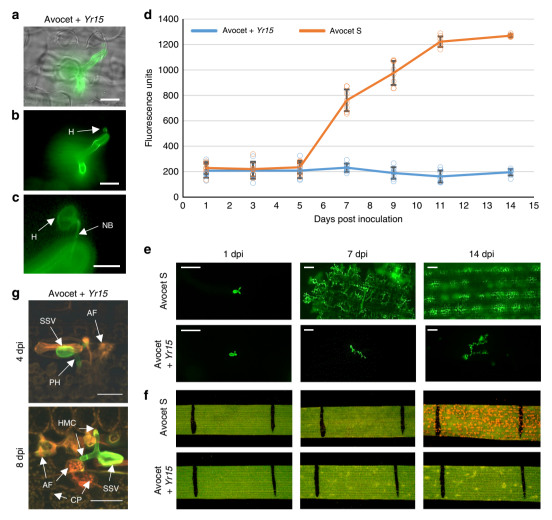


Fig. 1 Fungal development in *Pst*-infected Avocet S and Avocet + *Yr15* NILs during 14 days post inoculation (dpi). **a-c** *Pst* haustorium (H) within invaded plant host cell observed in Avocet + *Yr15* at 3 dpi under bright field (**a**) and fluorescence (**b**), connected to HMC via neckband (NB; **c**). Bars = $20 \, \mu m$ (**a**, **b**), $5 \, \mu m$ (**c**). **d** Comparative amounts of fungal biomass (chitin) within leaf tissues of NILs during 1-14 dpi. Error bars denote standard deviation (s.d.) based on eight biological replicates. **e** Fluorescence micrographs of fungal colonies and feeding structures at 1, 7, and 14 dpi in susceptible and resistant NILs. Bars = $100 \, \mu m$. **f** Leaf segments at 1, 7, and 14 dpi in susceptible and resistant NILs. The black lines delineate a 1 cm segment in the middle of the second leaf of the same plant. **g** Fungal structures and host responses in Avocet + *Yr15* at 4 and 8 dpi. The fungal colony (in green) consist of a substomatal vesicle (SSV) and primary infection hyphae (PH) with haustorial mother cells (HMC). Indicative signs of host cell autofluorescence (AF) are visible already at 4 dpi, while at 8 dpi the infected host cells show bright orange AF and appear to be collapsed and distorted; a couple of them contain aggregated chloroplasts (CP) with bright fluorescence as an indicator of host HR. Bars = $40 \, \mu m$

already at 1 day post inoculation (dpi) in both the susceptible common wheat genotype Avocet S and its resistant NIL Avocet + Yr15 (Supplementary Fig. 2a). Moreover, high-resolution closeup images presented in Fig. 1a-c clearly show HMCs, very thin penetrating hyphae (neckband) and haustoria feeding structures inside invaded mesophyll cells of Avocet + Yr15 at 3 dpi. Substantially larger fungal colonies and much more abundant pathogen feeding structures were observed in Avocet S compared with Avocet + Yr15 NIL, 3-14 dpi (Fig. 1d, e and Supplementary Fig. 2a). Relative quantification of fungal development and colonization of Avocet S (Fig. 1d) fitted the standard logistic population growth model well and displayed highly significant increase in biomass ($p < 10^{-15}$, likelihood ratio test), which is typical for the growth of microorganisms under favorable conditions²³. By contrast, colonization of Avocet + Yr15 was negligible and did not change significantly 1 to 14 dpi (p = 0.21, likelihood ratio test) (Fig. 1d). At 7 dpi fungal biomass residing within infected leaf tissues began to show significant differences between NILs (p < 0.001, t test, n = 16) (Fig. 1d), when

macroscopic evidence of HR became to be visible on Avocet + *Yr15* leaves, but not on Avocet S (Fig. 1f and Supplementary Fig. 2b). The differences in fungal growth between the NILs can be attributed to the strong HR response in Avocet + *Yr15*, which restricts the fungal growth at very early stages of infection. The host cells that are interacting with fungal feeding structures show clear indications of HR, as evident from the collapse and distortion of these cells, strong autofluorescence as well as the aggregation of the chloroplasts within the infected cells (Fig. 1g and Supplementary Fig. 3a–d). These results suggest that HR plays a central role in restricting the development of fungal feeding structures.

Map-based cloning of *Yr15*. A recombinant inbred mapping population segregating for *Yr15* was generated by crossing the susceptible recurrent durum line D447 with the resistant durum *Yr15* ILs (B9, B10), into which *Yr15* had been introgressed from $G25^{20}$. We screened 8573 F_2 plants with DNA markers to develop a high-resolution map of the *Yr15* gene region. First, we exploited

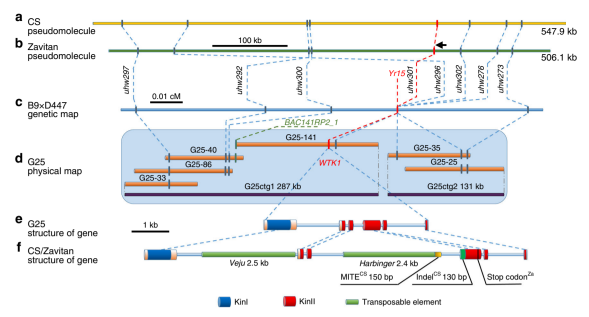


Fig. 2 Map-based cloning of *Yr15.* **a, b** Genomic sequence of CS (**a**) and Zavitan (**b**) 1BS pseudomolecules harboring the *WTK1* region. **c** Genetic map of the 1BS region carrying *Yr15*. Marker *uhw301* was developed from the *WTK1* sequence. **d** G25 BAC-based physical map covering the *Yr15* region. Assembly of BAC sequences yielded contigs G25ctg1 (287 kb) and G25ctg2 (131 kb). Marker *BAC141RP2_1* was used to connect G25-141 and G25-40 BAC clones. **e, f** Structure of *WTK1* alleles from G25 (**e**) and from CS and Zavitan (**f**). Features that are unique to CS are marked with superscript CS, while those that are present only in Zavitan are marked with superscript Za

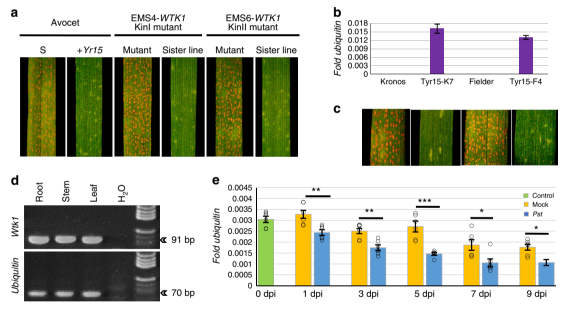


Fig. 3 Validation of *WTK1* as *Yr15*. **a** Loss of resistance to *Pst* in *WTK1* kinase-like domain mutants. EMS4 and EMS6 carry mutations in Kinl and Kinll, respectively. **b-c** *Wtk1* IF1 transcription levels (**b**) and *Pst* resistance phenotypes (**c**) in independent transgenic events Tyr15-K2 and Tyr15-F1. Error bars denoting standard error of mean (s.e.m.) are based on three technical replicates. **d** *Wtk1* expression survey in root, stem, and leaf of B9 by RT-PCR. **e** Transcript levels of the *Wtk1* IF1 in mock-inoculated or *Pst*-inoculated B9 plants. Error bars denoting s.e.m. are based on six biological replicates. Asterisks indicate the level of significance by *t* test, n = 12: p < 0.05 (*), p < 0.01 (**), p < 0.001 (***)

collinearity between the Yr15 region and the syntenic regions in model organisms $Brachypodium\ distachyon$, $Oryza\ sativa\ (rice)$, and $Sorghum\ bicolor\ (Supplementary\ Fig. 4a, b and Supplementary\ Tables 1–4). The nearest collinearity-derived markers, <math>uhw264$ and uhw258, defined Yr15 within an interval of 0.3 cM. The target interval containing Yr15 was then saturated with 24 markers developed from either bacterial artificial chromosome (BAC) sequences positioned on the physical map of chromosome arm 1BS of T. aestivum cv. Chinese Spring $(CS)^{24}$

(Supplementary Table 5), the genome assembly of WEW accession Zavitan (Zavitan)²⁵ (Supplementary Fig. 4c–e and Supplementary Table 6), or G25 BAC sequences. Using these markers, we mapped *Yr15* between markers *uhw300* and *uhw273* (0.013 cM) (Fig. 2a–c).

A pooled BAC library, constructed from Yr15 donor G25, was screened with the closely linked or co-segregating markers uhw297, uhw292, uhw296, and uhw273 (Fig. 2c), which were derived from the CS contig corresponding to the Yr15 region.

Sequencing of the six BACs isolated with these markers yielded two non-overlapping contigs (G25ctg1, 286,738 bp and G25ctg2, 131,485 bp; Fig. 2d). Sequence analysis of these contigs revealed the presence of three putative candidate genes. One of them contained predicted protein domains that previously have been associated with plant responses to pathogens and therefore was selected for validation. This gene contains two distinct kinase-like domains arranged in tandem and is designated here as *WTK1* (Fig. 2e, f). Alignment of the *WTK1* full-length complementary DNA (cDNA) against the *WTK1* genomic sequence (4657 bp) indicated that the gene contains six exons coding for a 665 amino acid protein.

Validation of the candidate gene for Yr15. To validate the function of WTK1, we used EMS to mutagenize a set of Yr15 ILs and identified 2 out of 2112 tetraploid and 8 out of 1002 hexaploid M2 families that segregated for resistance to Pst (Supplementary Table 7). Sequencing of WTK1 in the 10 susceptible M₂ plants confirmed the presence of independent missense mutations in each mutant. Five EMS mutants contained amino acid changes in the WTK1 kinase-like domain I (KinI) and five in kinase-like domain II (KinII) (Supplementary Table 7). Examples of these mutants (EMS4, having a mutation in KinI, and EMS6, with a mutation in KinII) are presented in Fig. 3a. In the F₂ progenies derived from crosses between the resistant wild-type Avocet + Yr15 and two of the susceptible mutants (EMS4 and EMS6), homozygosity for the non-functional WTK1 mutations, designated as *wtk1*, co-segregated with susceptibility to *Pst* (Supplementary Table 8). These 10 independent mutations demonstrate that both KinI and KinII domains of WTK1 are necessary for the resistance conferred by Yr15.

To determine whether WTK1 is sufficient to confer resistance to Pst, we transformed the susceptible varieties T. aestivum cv. Fielder and Triticum durum cv. Kronos with a 9.1 kb genomic fragment harboring the complete WTK1 coding region and its regulatory elements. Expression of the WTK1 transgene was detected in 17 out of 22 T_0 transgenic plants and their progenies. In subsequent generations when challenged with Pst (race PST-130 and isolate #5006), the WTK1 transgene co-segregated with the yellow rust resistance phenotype (Fig. 3b, c and Supplementary Fig. 5a–g), thereby validating that WTK1 is indeed Yr15.

Expression of Yr15 alternative transcript variants. Three alternative transcript variants—isoform 1 (IF1), isoform 2 (IF2), and isoform 3 (IF3)—were revealed by sequencing of 48 WTK1 cDNAs (Supplementary Fig. 6a). IF1, which encodes the complete WTK1 protein, was found to be the most common transcript (85.7% of tested WTK1 cDNAs), while the other isoforms are much less abundant (14.3% together) and encode truncated proteins. WTK1 is expressed in roots, stems, and leaves (Fig. 3d). Transcript levels of the three WTK1 isoforms were significantly lower in leaves of IL B9 inoculated with Pst, at all dpi (except 0 dpi), relative to mock-inoculated plants (Fig. 3e and Supplementary Fig. 6b). In addition, a significant reduction in expression was observed over time (1-9 dpi) for both Pst-inoculated (p $< 8.0 \times 10^{-8}$, F test, df₁,df₂ = 1,30) and mock-inoculated (p < 2.4×10^{-5} , F test, df₁,df₂ = 1,30) plants for the WTK1 IF1 (Fig. 3e) and Supplementary Fig. 6b). These findings suggest that expression of *WTK1* is down-regulated by the presence of the pathogen.

Changes in reading frames of non-functional alleles of Yr15. A search for WTK1 sequences on chromosome arm 1BS of the CS^{26} and $Zavitan^{25}$ whole-genome assemblies revealed only non-functional wtk1 alleles in both of these susceptible lines (Supplementary Fig. 1a). These alleles differ from the functional Wtk1

allele of WEW G25 by the presence of indels (e.g., insertions of transposable elements (TEs) such as *Veju*, *Harbinger*, or MITEs) that have changed the reading frame of exon 4 and generated premature stop codons in both CS and Zavitan, relative to G25 (Fig. 2e, f). In addition, orthologs of *Yr15* were found also on chromosome arms 1AS and 1DS. The predicted protein encoded by the ortholog on 1A of CS shares 83% similarity with WTK1 on 1BS of G25, but is 21 amino acids longer due to intron retentions. In contrast, the predicted proteins encoded by the 1A ortholog in Zavitan and the 1D ortholog in CS are shorter than the G25 1BS WTK1 due to frameshifts resulting in stop codons; their sequence similarities to G25 1BS WTK1 are relatively low (39.9 and 52.7%, respectively).

Phylogeny of orthologs, paralogs, and homologs of WTK1. In addition to the orthologs of WTK1 that were found on all homoeologous group 1 chromosomes of WEW (AABB) and common (AABBDD) wheats, we found WTK1 orthologs also in diploid wheat relatives Triticum urartu (AA), Aegilops speltoides (SS), and Aegilops tauschii (DD), representing the ancestral A, B, and D genomes, respectively, and on chromosome 1R of rye (Secale cereale, RR). A phylogenetic analysis clustered all the predicted proteins into a single branch, consisting of sub-clusters corresponding to the 1A, 1B, and 1D copies (Supplementary Fig. 7). No WTK1 ortholog was detected on chromosome 1H of barley (Hordeum vulgare, HH). Paralogs of WTK1 were found on the homeologous group 6 chromosomes of tetraploid and hexaploid wheat, their diploid relatives T. urartu and A. tauschii, and on chromosome 6H of barley. All group 6 copies clustered together phylogenetically. A homologous copy from chromosome Bd2 of B. distachyon showed an intermediate position between the two clusters, while the closest predicted protein from O. sativa was resolved as an outgroup. The presence of WTK1 paralogs on both group 1 and group 6 chromosomes of almost all diploid wheat species, as well as the relatively low nucleotide sequence identity between group 1 and group 6 paralogs of WTK1 (~80% in KinI and ~30% in KinII), suggests an old duplication event.

Distribution of *Wtk1* allele in wild and cultivated wheats. We have evaluated 390 *Triticum* accessions for the presence of *Wtk1* by gene-specific markers (Supplementary Tables 9–12). *Wtk1* was detected in 18% of the accessions of the southern (Israel, Lebanon, Jordan, and Syria) WEW populations but was not detected in the northern (Turkey and Iran) ones (Supplementary Table 10). We screened 93 modern durum and common wheat accessions and found that the functional *Wtk1* allele was present only in recently developed *Yr15* ILs (Supplementary Table 12).

WTK1 TKP architecture. Both WTK1 domains belong to a protein kinase-like superfamily with a conserved catalytic domain, but contain neither membrane-targeting motifs nor known receptor sequences. The lack of transmembrane domains in WTK1 is consistent with its cytoplasmic localization (with some partition into the nucleus) in barley protoplasts transformed with a WTK1-green fluorescent protein (GFP) fusion (Supplementary Fig. 8a-i). The presence of key conserved residues^{27,28} in KinI and the lack of most of these residues in KinII implies that WTK1 possesses a putative kinase-pseudokinase structure (Supplementary Fig. 9). Moreover, KinI belongs to serine/threonine non-RD kinases, shown previously to be associated with immune receptors²⁹. WKS1 is another Pst resistance gene from WEW that harbors a non-RD kinase domain. However, it is attached to a lipid-binding START domain¹⁰ and does not exhibit high similarity to either domains of WTK1 (Supplementary Fig. 10). Only two plant resistance genes have a tandem

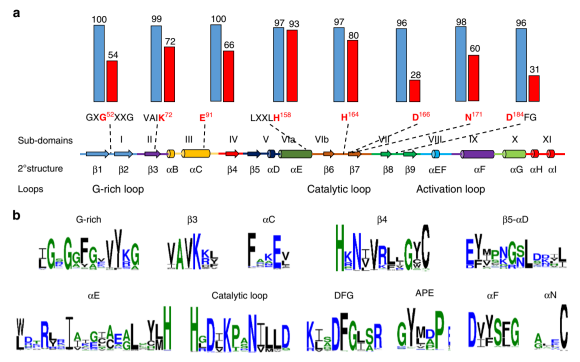


Fig. 4 Predicted structures of putative kinase and pseudokinase domains of TKP proteins based on 92 proteins from ten plant species. **a** Conservation of key motifs, residues, and secondary structure between putative kinase and pseudokinase domains. The secondary structure of a canonical kinase is shown with the standard annotations of its subdomains²⁷. Eight key residues are marked in red, with numbers that are based on their positions in α cAMP-dependent protein kinase catalytic subunit (cAPK)²⁷. Dashed lines indicate positions within secondary structure elements. The histograms above the motifs represent the degree of conservation (% of identical to conserved residuals) for putative kinase (blue columns) and putative pseudokinase (red) domains. **b** Sequence logos representing the conservation of key motifs and neighboring sequences in putative kinase domains. The size of the letters corresponds to their information content

kinase structure similar to WTK1: barley RPG1, conferring resistance to stem rust³⁰, and $MLOC_38442.1$, which was proposed as a candidate for the barley true loose smut resistance gene $Un8^{31}$. Nevertheless, phylogenetic analysis shows that they are quite distant from WTK1 (Supplementary Fig. 10, 11a).

The TKP protein family is found across the plant kingdom.

The occurrence of functional resistance genes with a tandem kinase structure (WTK1 and RPG130), as well as of many WTK1 orthologs and paralogs in wheat and its near relatives, motivated us to search for similar protein architectures across the plant kingdom. Altogether, we found 92 predicted proteins that are composed of putative kinase and pseudokinase domains in tandem; like WTK1, none had additional conserved domains. Most of the putative kinase domains share key conserved residues²⁸ (Fig. 4a, b and Supplementary Table 13), while the putative pseudokinase domains are generally highly divergent in these positions, suggesting that they probably have no or impaired kinase activity³² (Fig. 4a and Supplementary Table 14). The least conserved residues in the putative pseudokinase domains were found in the catalytic loop (D166 and D184) and the glycine-rich phosphate-binding loop (G⁵²)³³ (Fig. 4a). Analyses of motifs shared between putative kinase domains of TKP family members revealed additional conserved residues in sequences neighboring the core motifs (Fig. 4b). These results indicate that the putative kinase domains in this protein family may share a common

The phylogenetic analysis showed that all 184 putative kinase and pseudokinase domains of the 92 predicted proteins could be sorted into 11 major clades and two singletons (Supplementary Fig. 11a). Homology-based comparisons of the individual kinase domains of the 92 TKPs with the annotated *Arabidopsis* kinome

families³⁴ revealed clear relationships between the 11 clades and specific Arabidopsis families or subfamilies (Supplementary Fig. 11a). Members of clades 1-10 and cluster 11.2 were found to correspond to five families of plant receptor-like kinases (RLKs): concanavalin A-like lectin protein kinases (L-LPK), leucine-rich-repeat receptor kinases (LRR), receptor-like cytoplasmic kinases (RLCK), other kinases with no published family (RK), and cell-wall-associated kinases (WAKs). Members of cluster 11.1 were not classified as RLKs. Instead, they were found to be associated with the following families of soluble kinases: cATP-dependent, cGTP-dependent, and phospholipid-dependent kinases, cyclin-dependent kinases, raf-like MAPK (mitogenactivated protein kinase) kinase kinase (MAP3K-raf), MAPK kinase kinase (MAP3K), and SNF1-related kinase 3. All 92 TKPs can be divided into two distinct groups based on the topology of the phylogenetic tree: (i) two kinase domains of the same gene that are clustered together in the same branch most likely resulted from a duplication event; (ii) two kinase domains belonging to the same gene but are positioned on separate branches represent a gene fusion event (Supplementary Fig. 11a). While four clades (2, 5, 7, and 11) included proteins that, by this criterion, resulted from either gene duplications or fusions, three clades (6, 9, and 10) included proteins that originated only from duplication events, and four clades (1, 3, 4, and 8) included proteins that resulted only from fusion events. A total of 52 TKPs were derived from duplication (occurring in kinase domains associated with 10 different families or subfamilies) and the other 40 proteins resulted from gene fusions (represented by 10 combinations of domains belonging to different families or subfamilies; Supplementary Fig. 11b). The most common domains present in TKPs were similar to WAK kinases (63 out of 184) and LRR 8B (cysteine-rich kinases; 49 out of 184). WTK1 appears to be

derived from a fusion of WAK and RLCK_8 kinase domains. Interestingly, most fusion events were found in monocot TKPs. Notably, clade 6 includes proteins of distant species and was presumably originated from a single ancient duplication event of a kinase in the L-LPK family before the divergence of the monocots and eudicots.

Discussion

Although new *R*-genes are frequently discovered from various sources, the rapid evolution of *Pst* populations overcome most of the commonly used *Yr* genes, rendering most of the world wheat production susceptible to devastating epidemics⁴. The combination of marker-assisted selection and classical approaches can speed up the progressive incorporation of individual new *R*-genes into elite wheat varieties. However, it seems that the appearance of new virulent *Pst* races in the fields is faster than the incorporation of novel *R*-genes into elite wheat varieties.

Yr15 and Yr5 showed broad-spectrum resistance to Pst races from around the globe⁸ and are among the most promising genes for protecting wheat yields. Nevertheless, virulent races were detected against each one of them, separately, in Australia for $Yr5^{22}$ (isolate AU85569) and in Europe for $Yr15^{19}$ (e.g., isolate DK92/02). Therefore, we were interested to know whether the pyramiding of Yr5 and Yr15 into one cultivar could provide a more durable resistance. Our results indeed show that Yr5 and Yr15 complement each other and provide full resistance to the virulent races. Pyramiding resistance genes by classical breeding methods may take many years. However, approaches based on genetic transformation of individual cloned R-genes or of Rcassettes that include several R-genes with complementary resistances to various races, preferably via different mechanisms, can be considerably faster and can also overcome the problem of individual genes segregating in backcross populations. Yr5, Yr15, Yr18, Yr36, and Yr46 are obvious candidates to be included in such a cassette, since they represent different gene architectures (NLR, kinase-pseudokinase, ABC transporter, Kinase-START, and hexose transporter, respectively⁷) and most likely confer resistance via different molecular mechanisms.

The comparison of fungal development in susceptible vs. resistant NILs presented here provides indications for the possible mechanism of resistance conferred by Yr15. The macroscopic and microscopic observations, together with the quantification of fungal biomass in the resistant genotype that carries Yr15, show clear development of HR after fungal penetration, establishment of fungal feeding structures within the plant cells and restriction of fungal growth. The microscopic observations showed that a strong PCD response took place as part of the resistance mechanism, resulting in localized cell death that arrested further colonization of host cells by Pst^{5,6}. We showed that the size of chlorotic spots as a macroscopic indication of HR varied between the different genetic backgrounds of the Yr15 ILs. The HR/PCD response can be activated via different cellular pathways;⁵ further experiments are needed to identify the proteins interacting with WTK1 and the exact pathway that activates HR, in order to provide a better understanding of the mechanism of action and the host-parasite co-evolutionary processes behind it.

Two out of the six Yr genes cloned in wheat are derived from WEW (Yr36 (WKS1) and Yr15 (Wtk1)) and provide broadspectrum resistance against Pst. Both are present only in the southern distribution range of WEW populations in the Fertile Crescent (Israel, Lebanon, Jordan, and Syria) and were not detected in the northern populations (Turkey and Iran). Since wheat was probably domesticated from wild emmer in southeast Turkey¹, both Yr15 and Yr36 have, until recently, not found their way into the genepool of cultivated wheat germplasm. While the

WKS1 sequence is completely absent from all three genomes of hexaploid wheat 10, in the current study we have shown that nonfunctional orthologs and paralogs of WTK1 are present on all chromosomes of groups 1 and 6 of the tetraploid (AABB) and hexaploid (AABBDD) wheat genomes. Despite the relatively low similarity of orthologous and paralogous copies of WTK1 proteins in wheat that are associated with the susceptibility phenotype (Zavitan and CS), they still may carry resistance to some pathogens in other genetic backgrounds (such as diploid species) or may have other functional alleles in natural WEW populations. Furthermore, the presence of orthologs, paralogs, and homologs of WTK1 in all ancestral diploid genomes of wheat, and even in the basal Pooid Brachypodium, indicates that WAK-RLCK kinase-pseudokinase fusion represents an old event that occurred prior to the divergence of the wheat and Brachypodium lineages, about 35-40 million years ago³⁵. In contrast, the kinase-START protein architecture of WKS1 has not been found in other species outside of the Triticeae tribe and is likely a recent fusion of existing kinase and START domains¹⁰. For current breeding purposes, both Yr15 and Yr36 constitute untapped resources to improve resistance to Pst in domesticated wheat, since they are absent in all tested modern durum and common wheat varieties, except for recent introgressions.

The kinase protein superfamily represents 1-2% of the functional genes across plant and animal genomes³⁶, as reflected by the wide involvement of kinases in almost all cellular processes. The RLK clade is relatively expanded in flowering plants³⁶ and plays an important role in plant immune responses³⁷. The current phylogenetic analysis of the TKP protein family revealed that 175 out of 184 kinase domains were associated with RLKs, suggesting that TKPs are involved in plant defense mechanisms. Moreover, broad-spectrum resistance to biotrophic fungal pathogens (e.g., rusts and smut) was already demonstrated for WTK1 and RPG1³⁰ and proposed for Un8³¹, raising the question of whether the widespread TKPs serve generally as a family of resistance proteins. Most RLKs in Arabidopsis share multidomain architecture with receptor, transmembrane, and kinase domains³⁸, but 24% of them contain no extracellular or transmembrane domains, which are likewise lacking from TKP family members. Moreover, both RPG130 and WTK1 are localized mainly in the cytoplasm.

The evolution of multi-domain proteins, such as TKPs, presumes monophyletic or polyphyletic origins, respectively, from single or multiple independent evolutionary events³⁹ that can be explained by single or multiple birth models for protein families⁴⁰. Our phylogenetic analysis indicates that TKP family members originated from either gene duplication or gene fusion, suggesting a polyphyletic origin of the TKPs. Moreover, even divergent members of the TKP family, such as RPG130 and WTK1, which evolved, respectively, by duplication and fusion, share not only similar gene architecture but also analogous function (e.g., resistance to pathogens). Most of the TKP proteins (52 out of the considered 92) were formed by tandem duplication, earlier shown to be more frequent than fusion during the evolution of multi-domain proteins³⁹. In addition, we have seen indications on lineage-specific origin of some TKP members, that is, domains related to WAK family that were found only in monocots.

After gene duplication or fusion, one of the copies can gain a new function, a process known as neofunctionalization⁴¹. Interestingly, all members of the TKP family discussed here possess putative kinase-pseudokinase domain architecture and probably fit this neofunctionalization model. This structure may be related to the function of TKPs since some pseudokinases were shown to play an important role as baits for pathogen effectors⁴². Moreover, 20% of the RLK family members are pseudokinases, a higher

percentage than the average in the *Arabidopsis* kinome (13%)³⁶. The WTK1 pseudokinase belongs to the RCLK family. Members of this family were shown to serve as targets for inhibition by pathogen effectors (e.g., BIK1 in *Arabidopsis*)⁴³. Moreover, the duplication event itself was suggested as a source for the decoy model⁴⁴. Thus, the decoy role can be proposed as one of the potential mechanisms of function of the TKP family members in immune response. However, further studies are required in phylogenetically representative plant species in order to elucidate the mechanism of resistance conferred by this unique protein family.

Furthermore, members of the animal Janus kinase (JAK) family contain a similar structure of kinase and pseudokinase domains, a unique feature of JAKs compared to other animal protein kinases⁴⁵. The participation of members of both JAK⁴⁶ and TKP (e.g., WTK1) families in innate immune responses, particularly in PCD, calls attention to not only structural but also possibly functional similarities between these proteins, suggesting convergent molecular evolution of proteins involved in immunity in both the plant and the animal kingdoms.

Methods

Mapping populations and yellow rust assays. The yellow rust resistance gene Yr15 was introgressed from the resistant WEW accession G25 (G25) into the susceptible durum wheat (T. turgidum ssp. durum) accession D447 (LD393/ 2^* Langdon ND58-322) to develop the resistant BC₃F₉ (B9) or BC₃F₁₀ (B10) ILs²¹. A large mapping population, consisting of 8573 F₂ plants, was developed by crossing D447 with these ILs (B9 or B10), which carry Yr15 within a segment of chromosome arm 1BS from G25.

Phenotyping of recombinant, mutagenized, and transgenic lines under growth chamber conditions was carried out with Israeli Pst isolate #5006 (race 38E134) and US Pst race PST-130 using a standard protocol²¹. Plants were inoculated either at the two-leaf to four-leaf stage ("seedling inoculation") or at the stem elongation stage ("adult-plant inoculation"). The yellow rust response variation was evaluated 14 to 18 dpi using a 0 to 9 scale of IT⁴⁷.

Yr15 and Yr5 ILs were phenotyped for their response to Pst isolates DK92/02 and AU85569 from the isolate collection of the GRRC (Aarhus University, Flakkebjerg, Slagelse, Denmark). Plants were point inoculated with urediniospores suspended in engineered fluid (NovecTM 7100; 3M, Maplewood, MN, USA)⁴⁸. Using a 0 to 9 scale⁴⁷, the yellow rust responses were scored 18 dpi; the phenotypes were recorded using a Canon EOS 7D digital SLR camera equipped with a Canon macro EF 100mmf/2.8L IS USM lens (Canon, Tokyo, Japan).

Microscopy of *Pst*-wheat interactions within infected leaves. Fluorescence microscopy of *P. striiformis* structures was performed using wheat germ agglutinin (WGA; a lectin that binds specifically to β (1>4)-N-acetyl-D-glucosamine, i.e., chitin) conjugated with a fluorescent dye 49 . Leaf segments (second leaf, 10 cm long) from the NILs, Avocet S, and Avocet + *Yr15*, each of which had been inoculated with urediniospores of *Pst* isolate #5006, were sampled from 1 until 11 dpi (10:00 to 10:30 a.m., every 2 days), when visible sporulation developed on the susceptible plants. Fluorescence microscopy was carried out on an inverted fluorescence microscope, Leica DMi8 (Leica Microsystems, Wetzlar, Germany), fitted with a filter cube for the FITC excitation range (Ex: 460–500; Dc: 505; Em: 512–542), and a FLUO regime to observe the WGA-stained fungal structures.

Quantification of fungal biomass within infected leaves. After inoculation with Pst isolate #5006, leaf segments (middle, second leaf) of the resistant Avocet + Yr15 and susceptible Avocet S NILs were cut every 2 days (1–14 dpi) from different plants at each time point, with eight biological replicates collected for each line at each time point, and used for quantification of fungal biomass by chitin measurement 50 . Three technical replicates were made for each tissue sample. Fluorescence was measured on a SpectraMax M2e Microplate Reader (Molecular Devices, Sunnyvale, CA, USA), using 485-nm excitation and 535-nm emission wavelengths, a 1.0 s measurement time, and a cross pattern of well-scanning, yielding an average measurement per well. Statistical analyses of parameters associated with the accumulation of fungal biomass were determined by the maximum likelihood estimation method (SPSS) following the logistic population growth model. The null hypothesis (no growth) was tested by the likelihood ratio test

Confocal microscopy. The second leaf of 16-day-old plants of the susceptible Avocet S and resistant Avocet + Yr15 NILs were point inoculated⁴⁸ with fresh spores of the Pst isolate UK75/30. One leaf segment per plant was sampled at 4 and 8 dpi. Leaf samples were fixed in ethanol:chloroform (3:1, v/v) + 0.15% (w/v)

trichloroacetic acid solution for at least 24 h. Segments were then washed twice in 50% ethanol (10 min) followed by clearing in 0.05 M NaOH (30 min). After washing with deionized water (DI) and 0.1 M Tris-HCl buffer (pH 5.8) (30 min), samples were stained for 10 min in Tris-HCl buffer containing 0.1% Uvitex 2B (Polysciences Inc.) (w/v). Specimens were washed four times in DI followed by one wash in 25% glycerol and were left overnight in DI to remove access stain. Specimens were stored in 50% glycerol until further use.

For the microscopic investigations, leaf segments were mounted on glass slides in 75% glycerol. Fungal colonies and host cell autofluorescence were analyzed using an Olympus FV1200 confocal laser scanning microscope. Fungal structures and leaf tissues were excited with 405 and 515 nm lasers, and the emitted light was detected in two channels with filter settings 486–520 and 554–654 nm, respectively. Z-stacks were collected with 1 µm separation. The images from the two detection channels were merged and 3D projections were performed with IMARIS® bitplane 6.2. The resulting 2D images were adjusted for brightness and contrast. This included small adjustments in individual color channels to obtain an equal representation of the background signal from the healthy plant tissue on all images. All adjustments were applied equally across a whole image using the free software paint.net (https://www.getpaint.net/).

Development of a high-density genetic map. We assigned *Yr15* to deletion bin Sat0.31 of chromosome arm 1BS by developing and mapping dominant marker *uhw250*, as well as cleaved amplified polymorphic sequences⁵¹ markers *uhw252* and *uhw254*, based on expressed sequence tags assigned to Sat0.31⁵² (Supplementary Table 2). Using the GenomeZipper approach^{24,51}, which infers collinearity with orthologous regions of *B. distachyon*, *O. sativa*, and *S. bicolor* (Supplementary Table 3), eight additional markers were developed and the *Yr15*-containing region was reduced to a 0.3 cM interval between flanking markers *uhw264* and *uhw259* (Supplementary Fig. 4a).

Single-nucleotide polymorphism (SNP) markers RAC875_c826_839 and BS00022902_51 derived from the 15K wheat SNP array (Trait Genetics GmbH, Gatersleben, Germany) were converted to Kompetitive Allele-Specific PCR (KASP) markers using PolyMarker (http://polymarker.tgac.ac.uk/) (Supplementary Table 4).

The screening of F_2 plants from the D447 × B9 and D447 × B10 crosses for recombination events in the region carrying the Yr15 gene was carried out with the following marker sets (Supplementary Fig. 4b): (i) wmc406 and gwm273 (6); (ii) $RAC875_c826_839$ and $BS00022902_51$; (iii) uhw264 and uhw259. A total of 94 homozygous recombinant inbred lines (RILs), corresponding to 13 independent recombination events, were detected and used for the high-density map and for chromosome walking. These RILs were evaluated for resistance to Pst isolate #5006.

Physical maps of the Yr15 region on chromosome arm 1BS. Markers uhw264 and uhw259, flanking Yr15, were used to screen a gridded CS BAC library⁵³. Three positive BAC clones (TaaCsp364O11, TaaCsp1023G2, and TaaCsp1158K20) were detected with marker uhw264 and two (TaaCsp729H14 and TaaCsp814G12) with uhw259 (Supplementary Fig. 4b-d). The BAC-end sequence (BES) of clone TaaCsp364O11 was used to develop marker uhw267. Screening of the CS library with uhw267 yielded clone TaaCsp691F7. The BES of clone TaaCsp729H14 was used to develop marker uhw268, which was mapped distal to uhw259 (Supplementary Fig. 4c). These efforts generated two BAC contigs, flanking Yr15, with a gap between them, which was bridged using a 1BS physical map constructed from a CS 1BS-specific BAC library²⁴. The contigs of the CS 1BS physical map that cover the region corresponding to Yr15 were assembled as follows²⁴. A wheat BAC library composed of 55,296 BAC clones was constructed from CS 1BS. In total, 49,412 high-quality fingerprints, representing 14.4 1BS equivalents, were assembled into 57 long scaffolds covering 83% of chromosome 1BS using the LTC software²⁴. The LTC network representation of significant clone overlaps of the scaffold covering the Yr15 region is presented in Supplementary Fig. 4e. Screening of the 57 three-dimensionally arrayed BAC pools that comprise the minimal tiling path (6447 BACs selected from 49,412 fingerprinted clones) with markers uhw264, uhw267, uhw268, and uhw259 identified a 1.3 Mb contig spanning the complete Yr15 region (ctg49; Supplementary Fig. 4e). The 21 BAC clones (Supplementary Fig. 4d) that bridged the gap between markers uhw267 and uhw268 were identified and sequenced. Molecular markers developed from the sequenced clones of the 1BS contigs yielded genetic markers uhw297 and uhw292, which are distal to Yr15, uhw296, and uhw276, which co-segregate with Yr15, and uhw273, uhw275, uhw291, uhw274, uhw282, uhw284, and uhw277, as well as physical markers uhw280 and uhw279, which are proximal to Yr15 (Supplementary Table 5 and Supplementary Fig. 4c). The names and order of the 21 BAC clones are presented in Supplementary Fig. 4d.

The Yr15 donor line, G25, was used for construction of a pooled BAC library⁵⁴. High molecular weight genomic DNA was partially digested with *Hind*III to obtain fragments with sizes in the range of 100–250 kb, which were ligated into a pINDIGO vector (Caltech, Pasadena, CA, USA) and transformed into *Escherichia coli* cells⁵⁴. After growing individual *E. coli* colonies on agar plates, 443,880 transformed colonies were collected into 150 pools having an average of 2959 colonies per pool. The genome coverage of the G25 BAC library was calculated as 4.5× (average clone size, 120 kb). The initial screening of the library with marker *uhw280* yielded BAC clone G25-64, which was sequenced and used to develop the

proximal markers uhw288, uhw289, uhw287, uhw286, uhw286, and uhw281 (Supplementary Table 5). Further screening of the G25 BAC library with the closest distal markers uhw297 and uhw292, the co-segregating marker uhw296, and the proximal marker uhw273 yielded six BAC clones (G25-33, G25-86, G25-40, G25-141, G25-35, and G25-25; Fig. 2). DNA samples of these BACs were extracted using a Qiagen Plasmid Midi Kit or Qiagen Large-Construct Kit (Qiagen, Hilden, Germany). Contiguous sequences were generated on a Pacbio RS II (Pacific Biosciences, Menlo Park, CA, USA) at the Institute of Biotechnology, University of Helsinki (Helsinki, Finland). BAC clones were sequenced and assembled (HGAP3 implemented in SMRT portal 2.3) separately. The GAP4 program (Staden package) was used to edit and join the assembled BAC contigs into two contigs (G25ctg1, 286,738 bp; and G25ctg2, 131,485 bp) spanning the Yr15 gene region (Fig. 2). One Yr15 distal marker (uhw300) and two co-segregating markers (uhw302 and uhw301) were developed by comparing of G25 sequences to the 1BS pseudomolecule of Zavitan²⁵ (Supplementary Table 6).

G25 contig annotation and identification of candidate genes. Repetitive elements were masked using the Triticeae Repeat Sequence Database (TREP; http://botserv2.uzh.ch/kelldata/trep-db/index.html). Non-repetitive sequences were analyzed for genes by BLASTN searches against the GenBank (https://blast.ncbi.nlm.nih.gov/Blast.cgi) and the TIGR Wheat Genome (http://tigrblast.tigr.org/euk-blast/index.cgi?project=tae1) databases, using Genscan (http://genes.mit.edu/GENSCAN.html) and FGENESH (http://www.softberry.com/berry.phtml). A BLASTN search of the sequences of the G25ctg1 and G25ctg2 contigs against the high confidence (HC) gene models of the 1BS pseudomolecule of Zavitan²⁵ was used to reveal the presence of putative HC candidate genes.

Isolation and sequencing of the full-length WTK1 cDNA. Total RNA was extracted with the TRIzol reagent (Invitrogen, ThermoFisher Scientific, Waltham, MA, USA). Poly(A)+ RNA was purified from the total RNA with a Qligotex mRNA Midi Kit (Qiagen). First-strand cDNA was synthesized with Superscript II (Invitrogen) using primer E1820 (Supplementary Table 9). Nested PCR was carried out first with E1820 and the Yr15 5'-untranslated region (UTR) primer Y15F0 and then with E2146 (matching part of E1820; Supplementary Table 9) and the 5'-UTR primer Y15F2. The PCR products were purified, cloned, and sequenced.

To determine the start site of the *WTK1* transcript, we used 5' rapid amplification of cDNA ends (RACE)⁵⁵. In brief, the poly (A)+ RNA was treated with tobacco acid pyrophosphatase (Epicentre, Madison, WI, USA), and then purified with an RNeasy Plant Mini Kit (Qiagen). An adapter (RNAoligo, Supplementary Table 9) was ligated to the 5' end of the RNA and reverse transcription was carried out as above, primed with E1820. Amplification of the 5' end was carried out by two rounds of PCR reactions, using the primer 5' RACE (Supplementary Table 9) and sequentially two gene-specific primers, Y15R2 and Y15R1 (Supplementary Table 9).

Candidate gene validation by screening EMS-mutagenized lines. Seeds of the Yr15 durum wheat (B9) and common wheat (Avocet + Yr15, Excalibur + Yr15, and Suncea + Yr15) ILs were treated with 0.4–0.75% EMS solution for 16–18 h at room temperature ⁵⁶. Prior to sowing, the seeds were washed three times with 10% sodium thiosulfate and then twice in water (30 min each time), covered with Whatman paper, and air-dried at 4 °C.

EMS-treated M₁ plants were generated at the University of Haifa (Haifa, Israel) and at the University of Sydney (Cobbitty, Australia). M2 families (10-20 seeds per family) were artificially infected with Pst under field conditions in Israel (Pst isolate #5006), or under greenhouse conditions in Australia (Pst isolate 110 E143A+). All M₃ seedlings obtained from susceptible M₂ plants were inoculated in a growth chamber with Pst isolate #5006 to confirm the homozygosity of the recessive mutations, and then screened for mutations within the coding sequence of the Yr15 candidate gene WTK1. PCR products were amplified from genomic DNA with gene-specific primers (Supplementary Table 9), sequenced, and then compared for nucleotide variations by multiple sequence alignment. The full-length cDNA of WTK1 was amplified from the susceptible M2 plants and sequenced using two cDNA-specific primer pairs (WTK1_L2F and WTK_RE6, Supplementary Table 9), in order to confirm the detected point mutations at the messenger RNA (mRNA) level. The WTK1 mutations were ranked with SIFT (http://sift.jcvi.org/) to predict the effects of non-synonymous mutations on protein function (Supplementary Table 7).

We crossed two of the EMS-mutagenized lines of common wheat (EMS4 and EMS6, Supplementary Table 7) with the resistant wild-type parental line Avocet + Yr15 to produce segregating F_2 families. The response of each F_2 population to yellow rust was assessed at the seedling stage; goodness of fit for the observed and expected ratios in that population was evaluated with a χ^2 test (Supplementary Table 8). Sequence analysis of PCR products amplified from 40 F_2 plants was carried out as described above to confirm co-segregation of homozygosity for the mutations in WTK1 with the loss of resistance to Pst.

Candidate gene validation by transgenic complementation. Agrobacterium tumefaciens-mediated transformation of susceptible durum and common wheat varieties, respectively Kronos and Fielder, served to further verify WTK1 function

by complementation. Phusion* High-Fidelity DNA Polymerase (New England BioLabs, Ipswich, MA, USA) was used to amplify the *WTK1* genomic region from G25 BAC clone G25-141 (Fig. 2). Restriction sites *Xho1* and *AvrII* were added, respectively, to primers *Yr15F1/R1* and *Yr15F2/R2* (Supplementary Table 9) to enable cloning into the pLC41Hm transformation vector. Two overlapping PCR products were digested by the restriction enzyme pairs *Xho1-Msc1* and *Msc1-AvrII*, purified from agarose gel bands, and cloned into *Xho1-SpeI* linearized pLC41Hm. A 9116-bp genomic fragment that contains the full-length *Wtk1* coding region, as well as 3428 bp upstream of the start codon and 1031 bp downstream of the stop codon, was cloned for transformation. Sanger sequencing confirmed the accuracy of the construct. *Agrobacterium tumefaciens*-mediated transformation⁵⁷ of Kronos and Fielder with this construct was carried out at the University of California Plant Transformation Facility (Davis, CA, USA) (http://ucdptf.ucdavis.edu/).

In total, 15 independent T_0 plants in Kronos (Tyr15-K1 to Tyr15-K15) and 7 T_0 plants in Fielder (Tyr15-F1 to Tyr15-F7) were obtained. Three primer pairs (HpyFI/RI, Yr15TestFI/RI, and $Y15KI_F2/Yr15P2$; Supplementary Table 9) served to validate the presence of WTKI in the transgenic plants. In addition, we extracted mRNA from all T_0 plants and estimated the transcript levels of the three Wtk1 isoforms (isoform 1, IF1; isoform 2, IF2; isoform 3, IF3) by quantitative real-time PCR (qRT-PCR) using Ubiquitin as the endogenous control.

We germinated $10\text{--}25~\text{T}_1$ seeds from each transgenic event that expressed WTK1 and inoculated the plants with Pst race PST-130, which is virulent on Kronos and Fielder. All 12 positive T_1 families in Kronos and the 5 positive T_1 families in Fielder showed resistance to this Pst race. We genotyped all tested T_1 plants and confirmed co-segregation of the resistance with the presence of the transgene. Furthermore, 5 to 10 independent plants each of the T_2 families from transgenic events Tyr15-F1, Tyr15-F4, Tyr15-F5, Tyr15-F6, Tyr15-K7, Tyr15-K8, Tyr15-K10, Tyr15-K12, and Tyr15-K15 were also tested with Pst isolate #5006, which is likewise virulent on Kronos and Fielder.

Gene expression analysis by quantitative RT-PCR. Expression analysis was conducted with total RNA isolated from various plant tissues (leaves, roots, and stems) using an RNeasy Plant Mini Kit (Qiagen). First-strand cDNA was generated using the qScript TM Flex cDNA Synthesis Kit (Quanta Biosciences, Beverly, MA, USA). The qRT-PCR was performed on a StepOne Plus Real-Time PCR System (Applied Biosystems, Foster City, CA, USA) using the following program: 95 °C for 20 s; 40 cycles of 95 °C for 3 s and 60 °C for 30 s. The qRT-PCR reaction mixture contained the following components in a total volume of 10 µl: 5 µl Fast SYBR green master mix (Applied Biosystems); 2.5 µl diluted cDNA; 300 nM of each primer. The efficiency of each pair of primers was calculated using four serial fivefold dilutions (1:1, 1:5, 1:25, and 1:125) in triplicates. Amplification efficiencies of all primers were higher than 95%. Transcript levels are expressed as linearized fold-Ubiquitin levels calculated by the formula $2^{(Ubiquitin CT-Target CT)} \pm standard$ error of the mean (SEM).

Assessment of alternative splicing of WTK1 mRNA. Alternative splicing variants (IF1, IF2, IF3; Supplementary Fig. 6a) were revealed by sequencing of 48 B9 cDNA clones, which were amplified with a poly(T) primer and WTK1-specific primers, or with WTK1-specific primers alone. To determine the relative transcript levels of these three splicing variants, we designed PCR primers for each WTK1 isoform (Supplementary Table 9). Leaves of B9 were collected after the following treatments: (i) before inoculation (0 h), as the control; (ii) after inoculation with spores of Pst isolate #5006 suspended in Soltrol® 170 light oil; (iii) after spraying with Soltrol® 170 light oil lacking Pst spores, as a mock control. Samples of treatments (ii) and (iii) were collected at 1, 3, 5, 7, and 9 dpi, with six biological replicates for each treatment at each time point. All data were subjected to statistical analysis using the general linear model in SPSS.

Structure of wtk1 from CS and Zavitan. A search for *WTK1* sequences in the chromosome arm 1BS of CS²⁶ and in the Zavitan²⁵ whole-genome assemblies revealed the presence of non-functional alleles in both of these susceptible lines. The presence of three TEs: (i) a non-autonomous TRIM LTR retrotransposon of the Veju family (RLX_Taes_Veju) in intron 1 of CS and Zavitan); (ii) a DNA transposon of the DTH_Harbinger family (trep3042) in intron 3 of CS and Zavitan; (iii) a non-autonomous DNA transposon of the DTM_MITE type (trep1674-1) in intron 3 of CS, was identified by a search in two TE databases—mipsREdat_9.3p_Poaceae_TEs and trep-db_complete_Rel-16.

Evolutionary history of cereal WTK1 orthologs and paralogs. WTK1 protein sequences, used for phylogenetic analyses, were obtained from the genome assemblies of WEW (Zavitan pseudomolecules)²⁵, common wheat (CS pseudomolecules)²⁶, barley *H. vulgare* (Morex pseudomolecules) (http://webblast.ipk-gatersleben.de/barley_ibsc/), *A. speltoides* (accession #29; https://wheat-urgi.versailles.inra.fr/Seq-Repository/Assemblies), *A. tauschii* (http://plants.ensembl.org/Aegilops_tauschii/), *T. urartu* (NCBI number PRJNA182347), and rye *Secale cereale* (http://webblast.ipk-gatersleben.de/ryeselect/).

The public resource eggNOG version 4.5 (http://eggnogdb.embl.de) was searched for WTK1 Orthologous Group proteins at various taxonomic levels. The resulting set of WTK1 orthologs, designated as ENOG4115QHQ, included 11

protein sequences from nine different species. Two of these proteins were used for further phylogenetic analysis (*B. distachyon* BRADI2G38370.1 and *O. sativa* LOC_Os01g20880.1).

The analysis included 21 protein sequences. All sequence positions that contained gaps, as compared with WTK1-1B from G25, were eliminated from the analysis. Therefore, the analysis was conducted using 263 amino acid residues out of the 665 of the full-length WTK1 predicted from the DNA sequence of clone G25-141. The evolutionary history of WTK1 was inferred using the neighborjoining algorithm⁵⁸ based on evolutionary distances computed by the Poisson correction method⁵⁹. The quality of the derived phylogeny of WTK1 was assayed using a bootstrap test with 10,000 replicates⁶⁰. The analysis was performed with MEGA7 (https://www.megasoftware.net/).

Distribution of WTK1 among various Triticeae species. A set of two gene-specific, diagnostic markers, one for KinI and one for KinII (Supplementary Table 9), was developed and then used to test the distribution of *Wtk1* among various *Triticeae* species. These dominant PCR markers were designed to identify the presence of the *Wtk1* functional allele, amplifying a 1 kb PCR product for KinII and 2 kb for KinII.

Subcellular localization of the WTK1 protein. Barley protoplasts (8×10^5) were isolated 61 from H. vulgare cv. Bomi plants and electroporated with 20–40 μg DNA at 300 V/cm essentially as described before 61 , diluted into Gamborg's B-5 Basal Medium with minimal organics that contained 10% glucose, and then transferred into glass-bottomed 35 mm microwell dishes (MatTek Corporation, Ashland, MA, USA) for microscopy.

Electroporated protoplasts were cultured overnight at 22 °C before imaging under an inverted confocal laser scanning microscope (Leica TCS SP5 II; Leica Microsystems, Wetzlar, Germany) with a ×63 water immersion objective. The following light ranges were used: GFP (Ex 488 nm, Em 500–543 nm); 4',6-diamidino-2-phenylindole (Ex 405 nm, Em 430–550 nm); chlorophyll autofluorescence (Ex 488 nm, Em 673–725 nm).

A WTK1 cDNA clone was prepared as described under Isolation and sequencing of full-length WTK1 cDNA. All expression vectors were created with the Multisite Gateway Three-Fragment Vector Construction Kit (Invitrogen). The 5' entry clone was prepared by the BP reaction between pDONRP4-P1R and a 35S promoter fragment amplified from pBI221 (GenBank accession #AF502128.1) using primers attB4F35S and attB1R35S (Supplementary Table 9). The entry clone for the full-length WTK1 was created by carrying out the BP reaction between pDONR221 and the PCR product amplified from the WTK1 cDNA clone with primers attB1FKinase and attB2RYr15 (Supplementary Table 9). The entry clone for the N-terminal and C-terminal kinase-like domains were similarly prepared using primers attB1FKinase and attB2Rkinase, and attB1Freg and attB2RYr15 (Supplementary Table 9), respectively. The 3' entry clone was prepared by carrying out the BP reaction with pDONRP2RP3 and a GFPnos fragment amplified from pVEC8_GFP (GenBank: FJ949107.1) by PCR using primers attB2FGFPnos and attB3RGFPnos (Supplementary Table 9). The final clones used for transient transformation and expression in protoplasts were created by the LR reaction between the respective entry clones and the destination vector pDESTR4-R3 according to the instructions of the Gateway Kit.

Analysis of WTK1 kinase-like domains. A BLASTP search of the NCBI non-redundant protein database was used to assign the WTK1 kinase-like domains to a specific kinase superfamily and to search for proteins similar to WTK162. Multiple alignments of WTK1 kinase domains with 23 different plant kinase domains were performed using Clustal Omega with default parameters⁶³. In this analysis, we included the 4 closest kinases identified by BLASTP (WKS1, PTO, and two of WAK5), the 2 kinase domains of RPG1³⁰, the 2 kinase domains of *H. vulgare* MLOC_38442.1³¹, and 15 kinase domains from known and putative plant pattern recognition receptors, of which 14 are non-RD kinases and 1 is an RD kinase²⁹ (Supplementary Fig. 9). A phylogenetic tree was computed with RaxML⁶⁴ and drawn with iTOL (https://itol.embl.de/) to visualize the relationships between the different groups of kinases (Supplementary Fig. 10).

Analysis of the TKP family in plants. A search for predicted proteins with a TKP structure was conducted in whole-genome assemblies of various plant taxa: O. sativa (http://rapdb.dna.affrc.go.jp/download/irgsp1.html), Zea mays (http://ensembl.gramene.org/Zea_mays/), H. vulgare (http://webblast.ipk-gatersleben.de/barley_ibsc/), Secale cereale (http://webblast.ipk-gatersleben.de/ryeselect/), T. aestivum¹¹4, S. bicolor (https://phytozome.jgi.doe.gov/pz/portal.html#!info? alias=Org_Sbicolor), Arabidopsis thaliana (https://www.arabidopsis.org/download/index-auto.jsp?dir=/download_files/Proteins), Physcomitrella patens (https://plants.ensembl.org/Physcomitrella_patens/), Solanum tuberosum (http://plants.ensembl.org/Solanum_tuberosum/), Brassica napus (http://www.genoscope.cns.fr/brassicanapus/data/), and Populus trichocarpa (https://phytozome.jgi.doe.gov/pz/portal.html#!info?alias=Org_Ptrichocarpa). Kinomes were extracted according to functional annotations of the gene models of the above genome assemblies. Conserved kinase domains were identified using ProSITE (https://prosite.expasy.org/) and CD-Search (https://www.ncbi.nlm.nih.gov/Structure/cdd/

wrpsb.cgi), the predicted proteins with putative tandem kinases were extracted for further analysis. Predicted tandem kinase proteins with incomplete kinase domains were excluded from further analysis.

In total, 89 predicted protein sequences were selected for further analyses: *T. aestivum* (38 proteins), *H. vulgare* (8), *Secale cereale* (3), *O. sativa* (10), *Z. mays* (8), *S. bicolor* (12), *A. thaliana* (1), *Physcomitrella patens* (1), *Solanum tuberosum* (1), *Brassica napus* (5), and *P. trichocarpa* (2). In addition, the sequences of WTK1 from G25, RPG1³⁰, and MLOC_38442.1³¹ were included in the analysis to represent the tandem kinase structure, for which a function was assigned or proposed. Predicted protein sequences were aligned with MUSCLE (https://www.ebi.ac.uk/Tools/msa/muscle/). The evolutionary relationships between predicted proteins having a tandem kinase structure was inferred by the maximum likelihood method based on the JTT matrix-based model⁶⁵. The bootstrap consensus tree was calculated from 1000 replications⁶⁰. Branches were merged when a node was found in <50% of the bootstrap replicates (Supplementary Fig. 11a). The analysis was performed with MEGA7 (https://www.megasoftware.net/).

Graphical representation of amino acid conservation based on sequence logos was performed using WebLogo 3 (http://weblogo.threeplusone.com/). The presence of key conserved residues²⁷ was used to predict the catalytic activity of each domain.

Data availability

Sequences have been deposited in GenBank under accession numbers MG649384 and MG674157. The authors declare that all other data supporting the findings of this study are available within the manuscript and its supplementary files or are available from the corresponding author upon request.

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Author contributions

E.Y., L.H., D.R., V.K., L.F., A.F., Z.F., H.S., and A.D. developed markers. E.Y., L.H., D.R., V.K., and L.F. developed recombinants, conducted *Pst* phenotyping, and constructed genetic maps. D.R., Z.F., T.F., and A.B.K. constructed the CS 1BS physical map and identified the scaffold spanning the *Yr15* gene region. E.Y., L.H., and B.C. constructed and screened the G25 non-gridded BAC library. L.P. and P.L. sequenced the G25 BAC clones and assembled the G25 contigs. L.H., T.F. and A.F. identified the candidate genes. L.H., L.F., G.L., T.K., H.B., and T.F. developed EMS mutants. S.C. and J.D. performed the transgenic experiments. L.H. and W.C. isolated and sequenced full-length cDNA. W.C. and M.J.J. conducted the cellular localization experiment. V.K., T.K., S.C., and L.H. conducted expression analysis. V.K. carried out microscopic observations and quantification of fungal biomass upon *Pst* inoculation. K.S. performed confocal microscopy observations. A.F., C.S., and D.R. performed bioinformatics analyses of genomic and RNA-Seq data. A.F. and V.K. analyzed the predicted sequences of plant TKP family. V.K.

and L.H. conducted the *WTK1* distribution study. C.K.S. and M.S.H. phenotyped *Yr15/Yr5* ILs with a set of Pst isolates. T.F. conceived the project; T.F., A.H.S. and A.B.K. provided project leadership and planned the positional cloning strategy. V.K., T.F., and L. H. co-wrote the manuscript with input from all co-authors.

Additional information

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Competing interests: The University of Haifa and The Natural Resources Institute Finland (Luke) have filed two patent applications (numbers 62575688, 62575670) on the use of the sequences of the stripe rust resistance gene and the use of diagnostic markers that are based on the results described here, in which T.F., A.B.K., L.H., D.R., V.K., Z.F., A.F., A.H.S., and E.Y. are listed as inventors. The remaining authors declare no competing interests.

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