

1 **BED-domain containing immune receptors confer**
2 **diverse resistance spectra to yellow rust**

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19 **Introductory paragraph**

20 Crop diseases reduce wheat yields by ~25% globally and thus pose a major threat to global
21 food security¹. Genetic resistance can reduce crop losses in the field and can be selected for
22 through the use of molecular markers. However, genetic resistance often breaks down
23 following changes in pathogen virulence, as experienced with the wheat yellow (stripe) rust
24 fungus *Puccinia striiformis* f. sp. *tritici* (*Pst*)². This highlights the need to (i) identify genes
25 that alone or in combination provide broad-spectrum resistance and (ii) increase our
26 understanding of the underlying molecular modes of action. Here we report the isolation and
27 characterisation of three major yellow rust resistance genes (*Yr7*, *Yr5*, and *YrSP*) from
28 hexaploid wheat (*Triticum aestivum*), each having a distinct and unique recognition
29 specificity. We show that *Yr5*, which remains effective to a broad range of *Pst* isolates
30 worldwide, is allelic to *YrSP* and paralogous to *Yr7*, both of which have been overcome by
31 multiple *Pst* isolates. All three *Yr* genes belong to a complex resistance gene cluster on
32 chromosome 2B encoding nucleotide-binding and leucine-rich repeat proteins (NLRs) with a
33 non-canonical N-terminal zinc-finger BED domain³ that is distinct from those found in non-
34 NLR wheat proteins. We developed diagnostic markers to accelerate haplotype analysis and
35 for marker-assisted selection to enable the stacking of the non-allelic *Yr* genes. Our results
36 provide evidence that the BED-NLR gene architecture can provide effective field-based
37 resistance to important fungal diseases such as wheat yellow rust.

38

39 **Main**

40 In plant immunity, NLRs act as intracellular immune receptors that upon pathogen
41 recognition trigger a series of signalling steps that ultimately lead to cell death, thus
42 preventing the spread of infection^{4,5}. The NB-ARC domain is the hallmark of NLRs which in
43 most cases include leucine-rich repeats (LRRs) at the C-terminus. Recent *in silico* analyses

44 have identified NLRs with additional ‘integrated’ domains⁶⁻⁸, including zinc-finger BED
45 domains (BED-NLRs). The BED domain function within BED-NLRs is unknown, although
46 the BED domain from the non-NLR DAYSLEEPER protein was shown to bind DNA in
47 *Arabidopsis*⁹. BED-NLRs are widespread across Angiosperm genomes⁶⁻⁸ and this gene
48 architecture has been shown to confer resistance to bacterial blast in rice (*XaI*^{10,11}).

49

50 The genetic relationship between *Yr5* and *Yr7* has been debated for almost 45 years^{12,13}. Both
51 genes map to chromosome arm 2BL in hexaploid wheat and were hypothesized to be
52 allelic¹⁴, and closely linked with *YrSP*¹⁵. Whilst only one of >6,000 tested *Pst* isolates
53 worldwide has been found virulent to *Yr5* (Supplementary Table 1^{14,16}), both *Yr7* and *YrSP*
54 have been overcome in the field. For *Yr7*, this is likely due to its wide deployment in cultivars
55 (Supplementary Table 2, Supplementary Figure 1). This highlights the importance of
56 stewardship plans (including diagnostic markers) to deploy *Yr5* in combination with other
57 genes as currently done in the US (e.g. *Yr5+Yr15*; UC Davis breeding programme).

58

59 To clone the genes encoding *Yr7*, *Yr5*, and *YrSP*, we identified susceptible ethyl
60 methanesulfonate-derived (EMS) mutants from different genetic backgrounds carrying these
61 genes (Figure 1, Supplementary Tables 3-4). We performed MutRenSeq¹⁷ and isolated a
62 single candidate contig for each of the three genes based on nine, ten, and four independent
63 susceptible mutants, respectively (Figure 1a; Supplementary Figure 2). The three candidate
64 contigs were genetically linked to a common mapping interval, previously identified for the
65 three *Yr* loci^{15,18,19}. Their closest homologs in the Chinese Spring wheat genome sequence
66 (RefSeq v1.0) all lie within this common genetic interval (Figure 1b; Supplementary Figure
67 3).

68

69 Within each contig we predicted a single open reading frame based on RNA-Seq data. All
70 three predicted *Yr* genes displayed similar exon-intron structures (Figure 1a), although *YrSP*
71 was truncated in exon 3 due to a single base deletion that resulted in a premature termination
72 codon. The DNA sequences of *Yr7* and *Yr5* were 77.9% identical across the complete gene;
73 whereas *YrSP* was a truncated version of *Yr5*, sharing 99.8% identity in the common
74 sequence (Supplementary Files 1 and 2). This suggests that *Yr5* and *YrSP* are encoded by
75 alleles of the same gene, but are paralogous to *Yr7*. The 23 mutations identified by
76 MutRenSeq were confirmed by Sanger sequencing and all lead to either an amino acid
77 substitution or a truncation allele (splice junction or termination codon) (Figure 1a;
78 Supplementary Table 4). Taken together, the mutant and genetic analyses demonstrate that
79 *Yr5* and *YrSP* are allelic, while *Yr7* is encoded by a related, yet distinct gene.

80

81 The *Yr7*, *Yr5*, and *YrSP* proteins contain a zinc-finger BED domain at the N-terminus,
82 followed by the canonical NB-ARC domain. Unlike previously cloned resistance genes in
83 grasses (e.g. *Mla10l*²⁰, *Sr33*²¹, *Pm3*²²), neither *Yr7* nor *Yr5/YrSP* encode Coiled Coil domains
84 at the N-terminus (Supplementary Figure 4). Only *Yr7* and *Yr5* proteins encode multiple
85 LRR motifs at the C-terminus (Figure 2a; green bars), *YrSP* having lost most of the LRR
86 region due to the presence of the premature termination codon in exon 3. *YrSP* still confers
87 functional resistance to *Pst*, although with a different recognition specificity to *Yr5*. *Yr7* and
88 *Yr5/YrSP* are highly conserved in the N-terminus, with a single amino-acid change in the
89 BED domain. This high degree of conservation is eroded downstream of the BED domain
90 (Figure 2a). The BED domain is required for *Yr7*-mediated resistance, as a single amino acid
91 change in mutant line Cad0903 led to a susceptible reaction (Figure 1a). However,
92 recognition specificity is not solely governed by the BED domain, as the *Yr5* and *YrSP* alleles
93 have identical BED domain sequences, yet confer resistance to different *Pst* isolates. The

94 highly conserved Yr7 and Yr5/YrSP BED domains could function in a similar way to the
95 integrated WKRY domain in the *Arabidopsis* RRS1-R immune receptor which binds
96 unrelated bacterial effectors yet activates defense response through mechanisms involving
97 other regions of the protein²³.

98

99 We examined the allelic variation in *Yr7*, *Yr5*, and *YrSP* across eight sequenced tetraploid
100 and hexaploid wheat genomes (Supplementary Table 5). We identified *Yr7* only in Cadenza
101 and Paragon, which are identical-by-descent in this interval (Supplementary File 3,
102 Supplementary Table 6, and Supplementary Figure 5). Both cultivars are derived from the
103 original source of *Yr7*, tetraploid durum wheat (*T. turgidum* ssp. *durum*) cultivar Iumillo and
104 its hexaploid derivative Thatcher (Supplementary Figure 5). None of the three sequenced
105 tetraploid accessions (Svevo, Kronos, Zavitan) carry *Yr7* (Supplementary Table 6).

106

107 For *Yr5/YrSP*, we identified three additional alleles in the sequenced hexaploid wheat
108 cultivars (Figure 2b; Supplementary Table 7). Cultivar Claire encodes a complete NLR with
109 six amino-acid changes, including one within the NB-ARC domain, and six polymorphisms
110 in the C-terminus compared to *Yr5*. Cultivars Robigus, Paragon, and Cadenza also encode a
111 full length NLR that shares common polymorphisms with Claire, in addition to 19 amino acid
112 substitutions across the BED and NB-ARC domains. The C-terminus polymorphisms
113 between *Yr5* and the other cultivars is due to a 774 bp insertion in *Yr5*, close to the 3' end,
114 which carries an alternate termination codon (Supplementary File 2). Tetraploid cultivars
115 Kronos and Svevo encode a fifth *Yr5/YrSP* allele with a truncation in the LRR region distinct
116 from *YrSP*, in addition to multiple amino acid substitutions across the C-terminus
117 (Supplementary Table 7). This truncated tetraploid allele is reminiscent of *YrSP* and is
118 expressed in Kronos (see Methods). However, none of these cultivars (Claire, Robigus,

119 Paragon, Cadenza, Svevo, and Kronos) exhibit a *Yr5/YrSP* resistance response, suggesting
120 that these amino acid changes and truncations may alter recognition specificity or protein
121 function.

122

123 We designed diagnostic markers for *Yr7*, *Yr5*, and *YrSP* to facilitate their detection and use in
124 breeding. We confirmed their presence in the donor cultivars Thatcher and Lee (*Yr7*),
125 Spaldings Prolific (*YrSP*), and spelt wheat cv. Album (*Yr5*) (Supplementary Tables 8-9;
126 Supplementary Figures 5-6). We tested *Yr7* and *YrSP* markers in a collection of global
127 landraces²⁴ and European cultivars²⁵ released over the past one hundred years. *YrSP* was
128 absent from the tested germplasm, except for AvocetS-*YrSP* (Supplementary Table 9). *Yr7* on
129 the otherhand was more prevalent in the germplasm tested and we could track its presence
130 across pedigrees, including Cadenza derived cultivars (Supplementary Tables 8-9;
131 Supplementary Figure 5). We confirmed *Yr5* in the AvocetS-*Yr5* and Lemhi-*Yr5* lines and it
132 was not detected in the other tested lines, consistent with the fact that *Yr5* has not yet been
133 deployed within European breeding programmes (Supplementary Tables 10 and 17 and
134 Supplementary Figure 6). The *Yr5* diagnostic marker will facilitate its deployment, hopefully
135 within a breeding strategy that ensures its effectiveness long-term²⁶.

136

137 We defined the *Yr7/Yr5/YrSP* syntenic interval across the wheat genomes and related grass
138 species *Aegilops tauschii* (D genome progenitor), *Hordeum vulgare* (barley), *Brachypodium*
139 *distachyon*, and *Oryza sativa* (rice) (Supplementary files 4 and 5, Supplementary Figure 7).

140 We identified both canonical NLRs, as well as BED-NLRs across all genomes and species,
141 except for barley, which only contained canonical NLRs across the syntenic region. The
142 phylogenetic relationship based on the NB-ARC domain suggests a common evolutionary
143 origin of these integrated domain NLR proteins before the wheat-rice divergence (~50 Mya)

144 and an expansion in the number of NLRs in the A and B genomes of polyploid wheat species
145 (Figure 3a; Supplementary Figure 8). Within the interval we also identified several genes in
146 the A, B, and D genomes that encode two consecutive in-frame BED domains (named BED-I
147 and BED_II; Figure 3b-c, Supplementary Figure 7) followed by the canonical NLR. The
148 BED domains in these genes were fully encoded within a single exon (exons 2 and 3) and in
149 most cases had a four-exon structure (Figure 3c). This is consistent with the three-exon
150 structure of single BED domain genes, such as *Yr7* and *Yr5/YrSP* (BED-I encoded on exon
151 2). To our knowledge this is the first report of the double BED domain NLR protein
152 structure. The biological function of this molecular innovation remains to be determined,
153 although our data show that the single BED-I structure can confer *Pst* resistance and is
154 required for *Yr7*-mediated resistance.

155

156 Among other mechanisms, integrated domains of NLRs are hypothesised to act as decoys for
157 pathogen effector targets⁵. This would suggest that the integrated domain might be sequence-
158 related to the host protein targeted by the effector. To identify these potential effector targets
159 in the host, we retrieved all BED-domain proteins (108) from the hexaploid wheat genome,
160 including 25 BED-NLRs, and additional BED-NLRs located in the syntenic intervals
161 (Supplementary Table 11; Supplementary file 4). We also retrieved the rice *Xa1*^{10,11} and
162 ZBED proteins, the latter being hypothesized to mediate rice resistance to
163 *Magnaporthe oryzae*⁷. We used the split network method implemented in SplitsTree4²⁷ to
164 represent the relationships between these BED domains (Figure 3d; Supplementary Figure 9).
165 Overall, BED domains are diverse, although there is evidence of a split between BED
166 domains from BED-NLRs and non-NLR proteins (only 7 of 83 non-NLRs clustered with the
167 BED-NLRs). Given that the base of the split is broad, integrated BED-domains most likely
168 derive from multiple integration events. However, *Yr7* and *Yr5/YrSP* both arose from a

169 common integration event that occurred before the *Brachypodium*-wheat divergence
170 (Supplementary Figure 9, purple). This is consistent with the hypothesis that integrated
171 domains might have evolved to strengthen the interaction with pathogen effectors after
172 integration²⁸, although we cannot exclude the potential role of the BED domains in signalling
173 at this stage.

174

175 Among BED-NLRs, BED-I and BED-II constitute two major clades, consistent with their
176 relatively low amino acid conservation (Figure 3b), that are comprised solely of genes from
177 within the *Yr7/Yr5/YrSP* syntenic region. Seven non-NLR BED domain wheat proteins
178 clustered with BED-NLRs. These are most closely related to the *Brachypodium* and rice
179 BED-NLR proteins and were not expressed in RNA-Seq data from a *Yr5* time-course (re-
180 analysis of published data²⁹; Supplementary Figure 10, Supplementary Table 12). Similarly,
181 no BED-containing protein was differentially expressed during this infection time-course,
182 consistent with the prediction that effectors alter their targets' activity at the protein level in
183 the integrated-decoy model⁵. We cannot however disprove that these closely related BED-
184 containing proteins are involved in BED-NLR-mediated resistance.

185

186 BED-NLRs are frequent in Triticeae, and occur in other monocot and dicot tribes⁶⁻⁸. To date
187 a single BED-NLR gene, *Xal*, has been shown to confer resistance to plant pathogens^{10,11}. In
188 the present study, we show that the distinct *Yr7*, *Yr5*, and *YrSP* resistance specificities belong
189 to a complex NLR cluster on chromosome 2B and are encoded by two paralogous BED-
190 NLRs genes. We report an allelic series for the *Yr5/YrSP* gene with five independent alleles,
191 including three full-length BED-NLRs (including *Yr5*) and two truncated versions (including
192 *YrSP*). This wider allelic series could be of functional significance as previously shown for
193 the *Mla* and *Pm3* loci that confer resistance to *Blumeria graminis*^{22,30} in barley and wheat,

194 respectively, and the flax *L* locus conferring resistance to *Melampsora lini*³¹. Overall, our
195 results add strong evidence for the importance of the BED-NLR architecture in plant-
196 pathogen interactions. The paralogous and allelic relationship of these three distinct *Yr* loci
197 will inform future hypothesis-driven engineering of novel recognition specificities.

198 **Methods**

199 **MutRenSeq**

200 *Mutant identification*

201 Supplementary Table 3 summarises plant materials and *Pst* isolates used to identify mutants
202 for each *Yr* gene. We used an EMS-mutagenised population in cultivar Cadenza³² to identify
203 mutants in *Yr7*; whereas EMS-populations in the corresponding AvocetS-*Yr* near isogenic
204 line (NIL) were used to identify *Yr5* and *YrSP* mutants. For *Yr7*, we inoculated M₃ plants
205 from the Cadenza EMS population with *Pst* isolate 08/21 which is virulent to *Yr1*, *Yr2*, *Yr3*,
206 *Yr4*, *Yr6*, *Yr9*, *Yr17*, *Yr27*, *Yr32*, *YrRob*, and *YrSol*³³. We hypothesised that susceptible
207 mutants would carry mutations in *Yr7*. Plants were grown in 192-well trays in a confined
208 glasshouse with no supplementary lights or heat. Inoculations were performed at the one leaf
209 stage (Zadoks 11) with a talc – urediniospore mixture. Trays were kept in darkness at 10 °C
210 and 100% humidity for 24 hours. Infection types (IT) were recorded 21 days post-inoculation
211 (dpi) following the Grassner and Straib scale³⁴. Identified susceptible lines were progeny
212 tested to confirm the reliability of the phenotype and DNA from M₄ plants was used for
213 RenSeq (see section below). Similar methods were used for AvocetS-*Yr7*, AvocetS-*Yr5*, and
214 AvocetS-*YrSP* EMS-mutagenised populations with the following exceptions: *Pst* pathotypes
215 108 E141 A+ (University of Sydney Plant Breeding Institute Culture no. 420), 150 E16 A+
216 (Culture no. 598) and 134 E16 A+ (Culture no. 572) were used to evaluate *Yr7*, *Yr5*, and
217 *YrSP* mutants, respectively. EMS-derived susceptible mutants in Lemhi-*Yr5* were previously
218 identified³⁵ and DNA from M₅ plants was used for RenSeq.

219

220 *DNA preparation, resistance gene enrichment and sequencing (RenSeq)*

221 We extracted total genomic DNA from young leaf tissue using the large-scale DNA
222 extraction protocol from the McCouch Lab (https://ricelab.plbr.cornell.edu/dna_extraction)

223 and a previously described method³⁶. We checked DNA quality and quantity on a 0.8%
224 agarose gel and with a NanoDrop spectrophotometer (Thermo Scientific). Arbor Biosciences
225 (Ann Arbor, MI, USA) performed the targeted enrichment of NLRs according to the MYbaits
226 protocol using an improved version of the previously published Triticeae bait library
227 available at github.com/steuernb/MutantHunter. Library construction was performed using
228 the TruSeq RNA protocol v2 (Illumina 15026495). Libraries were pooled with one pool of
229 samples for Cadenza mutants and one pool of eight samples for the Lemhi-*Yr5* parent and
230 Lemhi-*Yr5* mutants. AvocetS-*Yr5* and AvocetS-*YrSP* wild-type, together with their respective
231 mutants, were also processed according to the MYbaits protocol and the same bait library
232 was used. All enriched libraries were sequenced on a HiSeq 2500 (Illumina) in High Output
233 mode using 250 bp paired end reads and SBS chemistry. For the Cadenza wild-type, we
234 generated data on an Illumina MiSeq instrument. In addition to the mutants, we also
235 generated RenSeq data for Kronos and Paragon to assess the presence of *Yr5* in Kronos and
236 *Yr7* in Paragon. Details of all the lines sequenced, alongside NCBI accession numbers, are
237 presented in Supplementary Tables 3 and 12.

238

239 **MutantHunter pipeline**

240 We adapted the pipeline from <https://github.com/steuernb/MutantHunter/> to identify
241 candidate contigs for the targeted *Yr* genes. First, we trimmed the RenSeq-derived reads with
242 trimmomatic³⁷ using the following parameters: ILLUMINACLIP:TruSeq2-PE.fa:2:30:10
243 LEADING:30 TRAILING:30 SLIDINGWINDOW:10:20 MINLEN:50 (v0.33). We made *de*
244 *novo* assemblies of wild-type plant trimmed reads with the CLC assembly cell and default
245 parameters apart from the word size (-w) parameter that we set to 64 (v5.0,
246 <http://www.clcbio.com/products/clc-assembly-cell/>) (Supplementary Table 14). We then
247 followed the MutantHunter pipeline detailed at <https://github.com/steuernb/MutantHunter/>.

248 For Cadenza mutants, we used the following MutantHunter program parameters to identify
249 candidate contigs: -c 20 -n 6 -z 1000. These options require a minimum coverage of 20x for
250 SNPs to be called; at least six susceptible mutants must have a mutation in the same contig to
251 report it as candidate; small deletions were filtered out by setting the number of coherent
252 positions with zero coverage to call a deletion mutant at 1000. The -n parameter was
253 modified accordingly in subsequent runs with the Lemhi-*Yr5* datasets (-n 6).

254

255 To identify *Yr5* and *YrSP* contigs from Avocet mutants, we followed the MutantHunter
256 pipeline with all default parameters, except in the use of CLC Genomics Workbench (v10)
257 for reads QC, trimming, *de novo* assembly of Avocet wild-type and mapping all the reads
258 against *de novo* wild-type assembly. The default MutantHunter parameters were used except
259 that -z was set as 100. The parameter -n was set to 2 in the first run and then to 3 in the
260 second run. Two *Yr5* mutants were most likely sibling lines as they carried identical
261 mutations at the same position (Supplementary Figure 2, Supplementary Table 4).

262

263 For *Yr7* we identified a single contig with six mutations, however we did not identify
264 mutations in line Cad0903. Upon examination of the *Yr7* candidate contig we predicted that
265 the 5' region was likely to be missing (Supplementary Figure 2). We thus annotated potential
266 NLRs in the Cadenza genome assembly available from the Earlham Institute (Supplementary
267 Table 5, http://opendata.earlham.ac.uk/Triticum_aestivum/EI/v1.1) with the NLR-Annotator
268 program using default parameters (<https://github.com/steuernb/NLR-Annotator>). We
269 identified an annotated NLR in the Cadenza genome with 100% sequence identity to the *Yr7*
270 candidate contig, which extended beyond our *de novo* assembled sequence. We therefore
271 replaced the previous candidate contig with the extended Cadenza sequence (100% sequence
272 identity) and mapped the RenSeq reads from Cadenza wild-type and mutants as described

273 above. This confirmed the candidate contig for *Yr7* as we retrieved the missing 5' region
274 including the BED domain. The improved contig now also contained a mutation in the
275 outstanding mutant line Cad0903 (Supplementary Figure 2). The Triticeae bait library does
276 not include integrated domains in its design so they are prone to be missed, especially when
277 located at the ends of an NLR. Sequencing technology could also have accounted for this:
278 MiSeq was used for Cadenza wild-type whereas HiSeq was chosen for Lemhi-*Yr5* and we
279 recovered the 5' region in the latter, although coverage was lower than for the regions
280 encoding canonical domains. In summary, we sequenced nine, ten, and four mutants for *Yr7*,
281 *Yr5*, and *YrSP*, respectively and identified for each target gene a single contig that accounted
282 for all mutants.

283

284 **Candidate contig confirmation and gene annotation**

285 We sequenced the *Yr7*, *Yr5*, and *YrSP* candidate contigs from the mutant lines (annotated in
286 Supplementary Files 1 and 2) to confirm the EMS-derived mutations using primers
287 documented in Supplementary Table 15. We first PCR-amplified the complete locus from the
288 same DNA preparations as the ones submitted for RenSeq with the Phusion® High-Fidelity
289 DNA Polymerase (New England Biolabs) following the suppliers protocol
290 (<https://www.neb.com/protocols/0001/01/01/pcr-protocol-m0530>). We then carried out
291 nested PCR on the obtained product to generate overlapping 600-1,000 bp amplicons that
292 were purified using the MiniElute kit (Qiagen). The purified PCR products were sequenced
293 by GATC following the LightRun protocol ([https://www.gatc-biotech.com/shop/en/lightrun-](https://www.gatc-biotech.com/shop/en/lightrun-tube-barcode.html)
294 [tube-barcode.html](https://www.gatc-biotech.com/shop/en/lightrun-tube-barcode.html)). Resulting sequences were aligned to the wild-type contig using
295 ClustalOmega (<https://www.ebi.ac.uk/Tools/msa/clustalo/>). This allowed us to curate the *Yr7*
296 locus in the Cadenza assembly that contained two sets of unknown ('N') bases in its

297 sequence, corresponding to a 39 bp insertion and a 129 bp deletion (Supplementary File 3),
298 and to confirm the presence of the mutations in each mutant line.

299 We used HISAT2³⁸ (v2.1) to map RNA-Seq reads available from Cadenza and AvocetS-
300 *Yr5*²⁹ to the RenSeq *de novo* assemblies with curated loci to define the structure of the genes.
301 We used the following parameters: --no-mixed --no-discordant to map reads in pairs only.
302 We used the --novel-splicesite-outfile to predict splicing sites that we manually scrutinised
303 with the genome visualisation tool IGV³⁹ (v2.3.79). Predicted coding sequences (CDS) were
304 translated using the ExPASy online tool (<https://web.expasy.org/translate/>). This allowed us
305 to predict the effect of the mutations on each candidate transcript (Figure 1a; Supplementary
306 Table 4). The long-range primers for both *Yr7* and *Yr5* loci were then used on the
307 corresponding susceptible Avocet NIL mutants to determine whether the genes were present
308 and carried mutations in that background (Figure 1a; Supplementary Files 1 and 2).

309

310 **Coiled coil domain prediction**

311 To determine whether *Yr7*, *Yr5*, and *YrSP* encode Coiled Coil (CC) domains we used the
312 NCOILS prediction program⁴⁰ (v1.0, https://embnet.vital-it.ch/software/COILS_form.html)
313 with the following parameters: MTK matrix with applying a 2.5-fold weighting of positions
314 a,d. We compared the profiles to those obtained with already characterised CC-NLR
315 encoding genes *Sr33*, *Mla10*, *Pm3* and *RPS5* (Supplementary Figure 4). We also ran the
316 program on *Yr7* and *Yr5* protein sequences where the BED domain was manually removed to
317 determine whether its integration could have disrupted an existing CC domain. To further
318 investigate whether *Yr7*, *Yr5*, and *YrSP* encode CC domains we performed a BLASTP
319 analysis⁴¹ with their N-terminal region, from the methionine to the first amino acid encoding
320 the NB-ARC domain, with or without the BED domain (Supplementary Figure 4).

321

322 **Genetic linkage**

323 We generated a set of F₂ populations to genetically map the candidate contigs
324 (Supplementary Table 3). For *Yr7* we developed an F₂ population based on a cross between
325 the susceptible mutant line Cad0127 to the Cadenza wild-type (population size 139
326 individuals). For *Yr5* and *YrSP* we developed F₂ populations between AvocetS and the NILs
327 carrying the corresponding *Yr* gene (94 individuals for *YrSP* and 376 for *Yr5*). We extracted
328 DNA from leaf tissue at the seedling stage (Zadoks 11) following a previously published
329 protocol⁴² and Kompetitive Allele Specific PCR (KASP) assays were carried out as described
330 in⁴³. R/qtl package⁴⁴ was used to produce the genetic map based on a general likelihood ratio
331 test and genetic distances were calculated from recombination frequencies (v1.41-6).

332

333 We used previously published markers linked to *Yr7*, *Yr5*, and *YrSP* (WMS526, WMS501
334 and WMC175, WMC332, respectively^{15,18,19}) in addition to closely linked markers WMS120,
335 WMS191, and WMC360 (based on the GrainGenes database
336 <https://wheat.pw.usda.gov/GG3/>) to define the physical region on the Chinese Spring
337 assembly RefSeq v1.0 (<https://wheat-urgi.versailles.inra.fr/Seq-Repository/Assemblies>). Two
338 different approaches were used for genetic mapping depending on the material. For *Yr7*, we
339 used the public data³² for Cad0127 (www.wheat-tilling.com) to identify nine mutations
340 located within the *Yr7* physical interval based on BLAST analysis against RefSeq v1.0. We
341 used KASP primers when available and manually designed additional ones including an
342 assay targeting the Cad0127 mutation in the *Yr7* candidate contig (Supplementary Table 15).
343 We genotyped the Cad0127 F₂ populations using these nine KASP assays and confirmed
344 genetic linkage between the Cad0127 *Yr7* candidate mutation and the nine mutations across
345 the physical interval (Supplementary Figure 3).

346

347 For *Yr5* and *YrSP*, we first aligned the candidate contigs to the best BLAST hit in an AvocetS
348 RenSeq *de novo* assembly. We then designed KASP primers targeting polymorphisms
349 between these sequences and used them to genotype the corresponding F₂ population
350 (Supplementary Table 15). For both candidate contigs we confirmed genetic linkage with the
351 previously published genetic intervals for these *Yr* genes (Supplementary Figure 3).

352

353 ***Yr7* gene-specific markers**

354 We aligned the *Yr7* sequence with the best BLAST hits in the genomes listed on
355 Supplementary Table 5 and designed KASP primers targeting polymorphisms that were *Yr7*-
356 specific. Three markers were retained after testing on a selected panel of Cadenza-derivatives
357 and cultivars that were positive for *Yr7* markers in the literature, including the *Yr7* reference
358 cultivar Lee (Supplementary Table 8, 8 and 15). The panel of Cadenza-derivatives was
359 phenotyped with three *Pst* isolates: *Pst* 08/21 (*Yr7*-avirulent), *Pst* 15/151 (*Yr7*-avirulent –
360 virulent to *Yr1*, 2, 3, 4, 6, 9, 17, 25, 32, *Rendezvous*, *Sp*, *Robigus*, *Solstice*) and *Pst* 14/106
361 (*Yr7*-virulent, virulent to *Yr1*, 2, 3, 4, 6, 7, 9, 17, 25, 32, *Sp*, *Robigus*, *Solstice*, *Warrior*,
362 *Ambition*, *Cadenza*, *KWS Sterling*, *Apache*) to determine whether *Yr7*-positive cultivars, as
363 identified by the three KASP markers, displayed a consistent specificity (Supplementary
364 Table 8). Pathology assays were performed as for the screening of the Cadenza mutant
365 population. We retrieved pedigree information for the analysed cultivars from the Genetic
366 Resources Information System for Wheat and Triticale database (GRIS,
367 www.wheatpedigree.net) and used the Helium software⁴⁵ (v1.17) to illustrate the breeding
368 history of *Yr7* in the UK (Supplementary Figure 5).

369

370 We used the three *Yr7* KASP markers to genotype (i) cultivars from the AHDB Wheat
371 Recommended List from 2005-2018 (<https://cereals.ahdb.org.uk/varieties/ahdb->

372 recommended-lists.aspx); (ii) the Gediflux collection of European bread wheat cultivars
373 released between 1920 and 2010²⁵ and (iii) the core Watkins collection, which represents a
374 global set of wheat landraces collected in the 1930s²⁴. Results are reported in Supplementary
375 Table 9.

376

377 ***Yr5* and *YrSP* gene-specific markers**

378 We identified a 774 bp insertion in the *Yr5* allele 29 bp upstream of the STOP codon with
379 respect to the Cadenza and Claire alleles. Genomic DNA from *YrSP* confirmed that the
380 insertion was specific to *Yr5*. We used this polymorphism to design primers flanking the
381 insertion and tested them on a subset of the collections mentioned above. We added 32 DNA
382 sample from diverse accessions of *Triticum dicoccoides*, the wild progenitor of domesticated
383 wheat (passport data shown in Supplementary Table 17). We included DNA from *Triticum*
384 *aestivum* ssp. *spelta* var. *album*³⁵ (*Yr5* donor) and Spaldings Prolific (*YrSP* donor) to assess
385 their amplification profiles. PCR amplification was conducted using a touchdown
386 programme: 10 cycles, -0.5 °C per cycle starting from 67 °C and the remaining 25 cycles at
387 62 °C. This allowed us to increase the specificity of the reaction. We observed three different
388 profiles on the tested varieties; (i) a 1,281 bp amplicon in *Yr5* positive cultivars, (ii) a 507 bp
389 amplicon in the alternate *Yr5* allele carriers, including AvocetS-*YrSP*, Cadenza, and Claire,
390 and (iii) no amplification in other varieties. We sequenced the different amplicons and
391 confirmed the insertion in *Yr5* compared to the alternate alleles (Supplementary File 2). The
392 lack of amplicons in some varieties most likely represents the absence of the loci in the
393 tested varieties. For *YrSP*, we aligned the *YrSP* and *Yr5* sequences to design KASP primers
394 targeting the G to C SNP between the two alleles (Supplementary File 2, Supplementary
395 Table 16). We tested the marker by genotyping selected cultivars as controls and cultivars
396 from the AHDB Wheat Recommended List from 2005-2018 (Supplementary Table 9).

397

398 ***In silico* allele mining for Yr7 and Yr5**

399 We used the Yr7 and Yr5 sequences to retrieve the best BLAST hits in the *T. aestivum* and *T.*
400 *turgdium* wheat genomes listed in Supplementary Table 5. The best Yr5 hits shared between
401 93.6 and 99.3% sequence identity, which was comparable to what was observed for alleles
402 derived from the wheat *Pm3* (>97% identity)⁴⁶ and flax *L* (>90% identity)³¹ genes. Yr7 was
403 identified only in Paragon and Cadenza (Supplementary Table 6; See Supplementary File 3
404 for curation of the Paragon sequence).

405

406 **Analysis of the Yr7 and Yr5/YrSP cluster on RefSeq v1.0**

407 *Definition of syntenic regions across grass genomes*

408 We used NLR-Annotator to identify putative NLR loci on RefSeq v1.0 chromosome 2B and
409 identified the best BLAST hits to Yr7 and Yr5 on RefSeq v1.0. Additional BED-NLRs and
410 canonical NLRs were annotated in close physical proximity to these best BLAST hits.
411 Therefore, to better define the NLR cluster we selected ten non-NLR genes located both
412 distal and proximal to the region, and identified orthologs in barley, *Brachypodium*, and rice
413 in *EnsemblPlants* (<https://plants.ensembl.org/>). We used different % ID cutoffs for each
414 species (>92% for barley, >84% for *Brachypodium*, and >76% for rice) and determined the
415 syntenic region when at least three consecutive orthologues were found. A similar approach
416 was conducted for *Triticum* ssp and *Ae. tauschii* (Supplementary file 4).

417

418 *Definition of the NLR content of the syntenic region*

419 We extracted the previously defined syntenic region from the grass genomes listed in
420 Supplementary Table 5 and annotated NLR loci with NLR-Annotator. We maintained
421 previously defined gene models where possible, but also defined new gene models that were

422 further analysed through a BLASTx analysis to confirm the NLR domains (Supplementary
423 Files 4 and 5). The presence of BED domains in these NLRs was also confirmed by CD-
424 Search (<https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi>).

425

426 **Phylogenetic and neighbour network analyses**

427 We aligned the translated NB-ARC domains from the NLR-Annotator output with
428 MUSCLE⁴⁷ using default parameters (v.3.8.31). We verified and manually curated the
429 alignment with Jalview⁴⁸ (v2.10.1). We used Gblocks⁴⁹ (v0.91b) with the following
430 parameters: Minimum Number Of Sequences For A Conserved Position: 9; Minimum
431 Number Of Sequences For A Flanking Position: 14; Maximum Number Of Contiguous
432 Nonconserved Positions: 8; Minimum Length Of A Block: 10; Allowed Gap Positions: None;
433 Use Similarity Matrices: Yes; to eliminate poorly aligned positions. This resulted in 36% of
434 the original 156 positions being taken forward for the phylogeny. We built a Maximum
435 Likelihood tree with the RAxML⁵⁰ program and the following parameters: raxmlHPC -f a -x
436 12345 -p 12345 -N 1000 -m PROTCATJTT -s <input_alignment.fasta> (MPI version
437 v8.2.10). The best scoring tree with associated bootstrap values was visualised and mid-
438 rooted with Dendroscope⁵¹ (v3.5.9). There was clear separation between NLRs belonging to
439 the two different clusters but the sub-clades had less support. One explanation would be that
440 conflicting phylogenetic signals that are due to events such as hybridization, horizontal gene
441 transfer, recombination, or gene duplication and loss might have occurred in the region. Split
442 networks allow nodes that do not represent ancestral species and can thus represent such
443 incompatible and ambiguous signals. We therefore used this method in the following part of
444 the analysis to analyse the relationship between the BED domains.

445

446 We used the Neighbour-net method⁵² implemented in SplitsTree4²⁷ (v4.16) to analyse the
447 relationships between BED domains from NLR and non-NLR proteins. First we retrieved all
448 BED-containing proteins from RefSeq v1.0 using the following steps: we used hmmer
449 (v3.1b2, <http://hmmer.org/>) to identify conserved domains in protein sequences from RefSeq
450 v1.0. We applied a cut-off of 0.01 on i-evalue to filter out any irrelevant identified domains.
451 We separated the set between NLR and non-NLRs based on the presence of the NB-ARC and
452 sequence homology for single BED proteins. BED domains were extracted from the
453 corresponding protein sequences based on the hmmer output and were verified on the CD-
454 search database. Alignments of the BED domains were performed in the same way as for
455 NB-ARC domains and were used to generate a neighbour network in SplitsTree4 based on
456 the uncorrected P distance matrix.

457

458 **Transcriptome analysis**

459 *Kronos analysis*

460 We reanalysed RNA-Seq data from cultivar Kronos⁵³ to determine whether the Kronos *Yr5*
461 allele was expressed. We followed the same strategy as that described to define the *Yr7* and
462 *Yr5* gene structures (candidate contig confirmation and gene annotation section). We
463 generated a *de novo* assembly of the Kronos NLR repertoire from Kronos RenSeq data and
464 used it as a reference when mapping read data from one replicate of the wild-type Kronos at
465 heading stage. Read depths up to 30x were present for the *Yr5* allele which allowed
466 confirmation of its expression. Likewise, the RNA-Seq reads confirmed the gene structure,
467 which is similar to *YrSP*, and the premature termination codon in Kronos *Yr5*. Whether this
468 allele confers resistance against *Pst* remains to be elucidated.

469

470 *Re-analysis of RNA-Seq data in Dobon et al., 2016*

471 We used RNA-Seq data previously published by Dobon and colleagues¹⁸. Briefly, two RNA-
472 Seq time-courses were used based on samples taken from leaves at 0, 1, 2, 3, 5, 7, 9, and 11
473 dpi for the susceptible cultivar Vuka and 0, 1, 2, 3, and 5 dpi for the resistant AvocetS-Yr5²⁹.
474 We used normalised read counts (Transcript Per Million, TPM) from Ramirez-Gonzalez et al.
475 2018 to produce the heatmap shown in Supplementary Figure 10 with the pheatmap R
476 package⁵⁴ (v1.0.8). Transcripts were clustered according to their expression profile as defined
477 by a Euclidean distance matrix and hierarchical clustering. Transcripts were considered
478 expressed if their average TPM was ≥ 0.5 TPM in at least one time point. We used the
479 DESeq2 R package⁵⁵ (v1.18.1) to conduct a differential expression analysis. We performed
480 two comparisons: (1) we used a likelihood ratio test to compare the full model \sim Cultivar +
481 Time + Cultivar:Time to the reduced model \sim Cultivar + Time to identify genes that were
482 differentially expressed between the two cultivars at a given time point after 0 dpi (workflow:
483 <https://www.bioconductor.org/help/workflows/rnaseqGene/>); (2) Investigation of both time
484 courses in Vuka and AvocetS-Yr5 independently to generate all of the comparisons between
485 0 dpi and any given time point, following the standard DESeq2 pipeline. Genes were
486 considered as differentially expressed genes if they showed an adjusted p-value < 0.05 and a
487 log₂ fold change of 2 or higher. Most BED-containing proteins and BED-NLRs were not
488 expressed in the analysed data. No pattern was observed for those that were expressed:
489 differences were observed between cultivars, but these were independent of the presence of
490 the yellow rust pathogen.

491

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629 **Author contributions**

630 CM performed the experiments to clone *Yr7* and *Yr5* and the subsequent analyses of their loci
631 and BED domains, designed the gene-specific markers, analysed the genotype data in the
632 studied panels, and designed and made the figures. JZ performed the experiments to
633 clone *YrSP*, confirm the *Yr7* and *Yr5* genes in AvocetS-*Yr7* and AvocetS-*Yr5* mutants, and
634 identified the full length of *Yr5* and *YrSP* with their respective regulatory elements. CM and
635 JZ developed the gene specific markers. PZ and RM performed the EMS treatment, isolation,
636 and confirmation of *Yr7*, *Yr5*, and *YrSP* mutants in AvocetS NILs. PF performed the
637 pathology work on the Cadenza *Yr7* mutants and the mapping populations. BS helped with
638 the NLR annotator analysis and provided the bait library for target enrichment and
639 sequencing of NLRs, NMA provided DNA samples for allelic variation studies and LB
640 provided Lemhi-*Yr5* mutants. RM, EL, PZ, BW, SB, and CU conceived, designed, and
641 supervised the research. CM and CU wrote the manuscript. JZ, PZ, RM, BW, NMA, LB and
642 EL provided edits.

643

644 **Data availability**

645 All sequencing data has been deposited in the NCBI Short Reads Archive under accession
646 numbers listed in Supplementary Table 13 (SRP139043). Cadenza (*Yr7*) and Lemhi (*Yr5*)
647 mutants are available through the JIC Germplasm Resource Unit (www.seedstor.ac.uk).

648

649 **Competing interests**

650 A patent application based on this work has been filed (United Kingdom Patent Application
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668 **Figure legends**

669 **Figure 1: *Yr5* and *YrSP* are allelic, but paralogous to *Yr7*.**

670 **a**, Left: Wild-type and selected EMS-derived susceptible mutant lines for *Yr7*, *Yr5*, and *YrSP*
671 (Supplementary Table 3 and 3) inoculated with *Pst* isolate 08/21 (*Yr7*), *Pst* 150 E16 A+
672 (*Yr5*), or *Pst* 134 E16 A+ (*YrSP*). Right: Candidate gene structures, with mutations in red,
673 and their predicted effects on the translated protein. **b**, Schematic representation of the
674 physical interval of the *Yr* loci. The *Yr7/Yr5/YrSP* locus is shown in orange on chromosome
675 2B with previously published SSR markers in black. Markers developed in this study to
676 confirm the genetic linkage between the phenotype and the candidate contigs are shown as
677 black lines underneath the chromosome. *Yr* loci mapping intervals are defined by the red
678 horizontal lines. A more detailed genetic map is shown in Supplementary Figure 3.

679

680 **Figure 2: *Yr7* and *Yr5/YrSP* encode integrated BED-domain immune receptor genes.**

681 **a**, Schematic representation of the *Yr7*, *Yr5*, and *YrSP* protein domain organisation. BED
682 domains are highlighted in red, NB-ARC domains are in blue, LRR motifs from NLR-
683 Annotator are in dark green, and manually annotated LRR motifs (xxLxLxx) are in light
684 green. Black triangles represent the EMS-induced mutations within the protein sequence. The
685 plot shows the degree of amino acid conservation (50 amino acid rolling average) between
686 *Yr7* and *Yr5* proteins, based on the conservation diagram produced by Jalview (2.10.1) from
687 the protein alignment. Regions that correspond to the conserved domains have matching
688 colours. The amino acid changes between *Yr5* and *YrSP* are annotated in black on the *YrSP*
689 protein. **b**, Five *Yr5/YrSP* haplotypes were identified in this study. Polymorphisms are
690 highlighted across the protein sequence with orange vertical bars for polymorphisms shared
691 by at least two haplotypes and blue vertical bars for polymorphisms that are unique to the

692 corresponding haplotype. Matching colours across protein structures illustrate 100%
693 sequence conservation.

694 **Figure 3: BED domains from BED-NLRs and non-NLR proteins are distinct.**

695 **a**, Numbers of NLRs in the syntenic regions across grass genomes (see Supplementary Figure
696 7), including BED-NLRs. **b**, WebLogo (<http://weblogo.berkeley.edu/logo.cgi>) diagram
697 showing that the BED-I and BED-II domains are distinct, with only the highly conserved
698 residues that define the BED domain (red bars) being conserved between the two types. **c**,
699 Gene structure most commonly observed for BED-NLRs and BED-BED-NLRs within the
700 *Yr7/Yr5/YrSP* syntenic interval. **d**, Neighbour-net analysis based on uncorrected *P* distances
701 obtained from alignment of 153 BED domains including the 108 BED-containing proteins
702 (including 25 NLRs) from RefSeq v1.0, BED domains from NLRs located in the syntenic
703 region as defined in Supplementary Figure 7, and BED domains from Xa1 and ZBED from
704 rice. BED-I and II clades are highlighted in purple and blue, respectively. BED domains from
705 the syntenic regions not related to either of these types are in red. BED domains derived from
706 non-NLR proteins are in black and BED domains from BED-NLRs outside the syntenic
707 region are in grey. Seven BED domains from non-NLR proteins were close to BED domains
708 from BED-NLRs. Supplementary Figure 9 includes individual labels.

709 **Supplementary Figure 1: Deployment of *Yr7* cultivars in the field is correlated with an**
710 **increase in the prevalence of *Pst* isolates virulent on *Yr7* in the UK.**

711 Percentage of total harvested weight of wheat cultivar carrying *Yr7* (green) and the
712 proportion of *Pst* isolates that are virulent to *Yr7* (orange) from 1990 to 2016 in the United
713 Kingdom. See Supplementary Table 2 for a summary of the data.

714

715 **Supplementary Figure 2: Identification of candidate contigs for the *Yr* loci using**
716 **MutRenSeq.**

717 View of RenSeq reads from the wild-type and EMS-derived mutants mapped to the best
718 candidate contigs identified with MutantHunter for the three genes targeted in this study.
719 From top to bottom: vertical black lines represent the *Yr* loci, coloured rectangles depict the
720 motifs identified by NLR-Annotator (each motif is specific to a conserved NLR domain⁵⁶),
721 while read coverage (grey histograms) is indicated on the left, e.g. [0 - 149], and the line from
722 which the reads are derived on the right, e.g. CadWT for Cadenza wild-type. Vertical bars
723 represent the position of the SNPs identified between the reads and reference assembly – red
724 shows C to T transitions and green G to A transitions. Black boxes highlight SNP for which
725 the coverage was relatively low, but still higher than the 20x detection threshold. The top
726 view shows the *Yr7* allele annotated from the Cadenza genome assembly before manual
727 curation (Supplementary File 3). Vertical black lines illustrate the assembled candidate
728 contigs and the one that was formerly *de novo* assembled from Cadenza RenSeq data, lacking
729 the 5' region containing the BED domain and thus the Cad903 mutation. The middle view
730 illustrates the *Yr5* locus annotated from the Lemhi-*Yr5* *de novo* assembly. The results are
731 similar to those described above for *Yr7*. The full locus was *de novo* assembled. The bottom
732 view illustrates the *YrSP* locus annotated from the AvocetS-*YrSP* *de novo* assembly with the
733 four identified susceptible mutants all carrying a mutation in the candidate contig. The full
734 locus was *de novo* assembled.

735

736 **Supplementary Figure 3: Candidate contigs identified by MutRenSeq are genetically**
737 **linked to the *Yr* loci mapping interval.**

738 Schematic representation of chromosome 2B from Chinese Spring (RefSeq v1.0) with the
739 positions of published markers linked to the *Yr* loci and surrounding closely linked markers
740 that were used to define their physical position (orange rectangle). The chromosome is
741 depicted as a close-up of the physical locus indicating the positions of KASP markers that
742 were used for genetic mapping (horizontal bars, Supplementary Table 15). Blue colour refers

743 to *Yr7*, red to *Yr5*, and purple to *YrSP*. The black arrow points to the NLR cluster containing
744 the best BLAST hits for *Yr7* and *Yr5/YrSP* on RefSeq v1.0. Coloured lines link the physical
745 map to the corresponding genetic map for each targeted gene (see Methods). Genetic
746 distances are expressed in centiMorgans (cM).

747

748 **Supplementary Figure 4: *Yr7*, *Yr5* and *YrSP* proteins do not encode for a Coiled-Coil**
749 **domain in the N-terminus.**

750 Graphical outputs from the COILS prediction program in three sliding windows (14, 21,
751 and 28 amino acid, shown in green, blue, and red, respectively) for *Yr5* and *Yr7* with or
752 without the BED domain (left) and characterised canonical NLRs: Sr33²¹, Mla10²⁰, Pm3²²
753 and RPS5⁵⁷. The X axis shows the amino acid positions and the Y axis the probability of a
754 coiled coil domain formation. There was no difference in the prediction between the two Yr
755 proteins with or without their BED domain. The 14 amino acid sliding window is the least
756 accurate according to the user manual, consistent with the additional peaks observed in Sr33,
757 Mla10 and Pm3 that were not annotated as CC domains in the corresponding publications²⁰⁻
758 ²². Thus, the peak at position 1,200 in *Yr5* is unlikely to represent a CC domain. We
759 performed a BLASTP search with the N-terminal region of the *Yr5* and *Yr7* proteins (from
760 Met to the first amino-acid encoding the NB-ARC) with or without the BED domain and the
761 best hits were proteins predicted to encode BED-NLRs from *Aegilops tauschii*, *Triticum*
762 *uratu* and *Oryza Sativa* (data not shown). Based on the COILS prediction and the BLAST
763 search, we concluded that *Yr7* and *Yr5/YrSP* do not encode CC domains.

764

765 **Supplementary Figure 5: Pedigrees of selected Thatcher-derived cultivars and their *Yr7***
766 **allelic status.**

767 Pedigree tree of Thatcher-derived cultivars where each circle represents a cultivar and the
768 size of the circle is proportional to its prevalence in the tree. Colours illustrate the genotype
769 with red showing the absence of *Yr7* and yellow its presence. Cultivars in grey were not
770 tested or are intermediate crosses. *Yr7* originated from *Triticum durum* cv. Iumillo and was
771 introgressed into hexaploid wheat through Thatcher (indicated by arrow). Each *Yr7* positive
772 cultivar is related to a parent that was also positive for *Yr7*. Figure was generated using the
773 Helium software⁴⁵ (v1.17).

774

775 **Supplementary Figure 6: Diagnostic genetic marker for *Yr5*.**

776

777 The *Yr5*-specific insertion was used to generate a PCR amplification product of 1,281 bp for
778 *Yr5* or a shorter amplicon for the absence of the insertion in *YrSP*, Claire, and Paragon (507
779 bp). *Yr5* positive lines include the *Yr5* spelt donor and *Yr5* near-isogenic lines AvocetS-*Yr5*
780 and Lemhi-*Yr5*. *YrSP* donor Spaldings Prolific and *YrSP* near-isogenic lines AvocetS-*YrSP*
781 carry the shorter alternate allele, similar to the Claire, Cadenza and Paragon alleles identified
782 in Figure 2. Negative controls include AvocetS and H₂O. Size marker is shown on the left.

783

784 **Supplementary Figure 7: Expansion of BED-NLRs in the Triticeae and presence of**
785 **conserved BED-BED-NLRs across the syntenic region.**

786 Schematic representation of the physical loci containing *Yr7* and *Yr5/YrSP* homologs on
787 RefSeq v1.0 and its syntenic regions. The syntenic region is flanked by conserved non-NLR
788 genes (orange arrows). Black arrows represent canonical NLRs and purple/blue/red arrows
789 represent different types of BED-NLRs based on their BED domain and their relationship
790 identified in Figure 3 and Supplementary Figure 8. Black lines represent phylogenetically
791 related single NLRs located between the two NLR clusters illustrated in Supplementary
792 Figure 9. Details of genes are reported in Supplementary File 4.

793

794 **Supplementary Figure 8: The *Yr* loci are phylogenetically related to nearby NLRs on**
795 **RefSeq v1.0 and their orthologs.**

796 Phylogenetic tree based on translated NB-ARC domains from NLR-Annotator. Node labels
797 represent bootstrap values for 1,000 replicates. The tree was rooted at mid-point and
798 visualized with Dendroscope v3.5.9. The colour pattern matches that of Figure 3 to highlight
799 BED-NLRs with different BED domains.

800

801 **Supplementary Figure 9: Neighbour-net analysis network as shown in Figure 3 with**
802 **identifiers.**

803 Neighbour-net analysis based on uncorrected *P* distances obtained from alignment of 153
804 BED domains including the 108 BED-containing proteins (including 25 NLRs) from RefSeq
805 v1.0, BED domains from NLRs located in the syntenic region as defined in Supplementary
806 Figure 7, and BED domains from Xa1 and ZBED from rice. BED-I and II clades are
807 highlighted in purple and blue, respectively. BED domains from the syntenic regions not
808 related to either of these types are in red. BED domains derived from non-NLR proteins are
809 in black and BED domains from BED-NLRs outside the syntenic region are in grey. Seven
810 BED domains from non-NLR proteins were close to BED domains from BED-NLRs.

811

812 **Supplementary Figure 10: BED-NLRs and BED-containing proteins are not**
813 **differentially expressed in yellow rust-infected susceptible and resistant cultivars.**

814 Heatmap representing the normalised read counts (Transcript Per Million, TPM) from the
815 reanalysis of published RNAseq data²⁹ for all the BED-containing proteins, BED-NLRs and
816 canonical NLRs located in the syntenic region annotated on RefSeq v1.0. Lack of expression
817 is shown in white and expression levels increase from blue to red. Asterisks show cases
818 where several gene models were overlapping with NLR loci identified with NLR Annotator.
819 The colour pattern matches that of Figure 3 to highlight BED-NLRs with different BED
820 domains. Orange labels show the expression of the canonical NLRs located within the
821 syntenic interval. The seven non-NLR BED genes whose BED domain clustered with the
822 ones from BED-NLR proteins in Figure 3 and Supplementary Figure 9 are indicated by black
823 triangles.

824

825 **Supplementary Table 1: Summary of *Pst* isolates tested on *Yr5* differential lines from**
826 **2004 to 2017 in different regions.**

827 Overall, >6,000 isolates from 44 countries displaying >200 different pathotypes were tested
828 on *Yr5* materials and no virulence was recorded apart from one isolate from Australia, PST
829 360 E137 A+¹⁴. Data were obtained from public databases and reports on yellow rust
830 surveillance, whose references are recorded. It is important to note that we report here the
831 number of identified pathotypes for a given region and database. Similar pathotypes could
832 thus have been counted twice if identified in different regions.

833

834 **Supplementary Table 2: Harvested weight of known *Yr7* cultivars from 1990 to 2016 and**
835 ***virYr7* prevalence among UK *Pst* isolates.**

836 Proportion of harvested *Yr7* wheat cultivars in the UK from 1990 to 2016. The prevalence of
837 yellow rust isolates virulent to *Yr7* across this time period is shown in the top row. Original
838 data from NIAB-TAG Seedstats journal (NIAB-TAG Network) and the UK Cereal Pathogen
839 Virulence Survey (<http://www.niab.com/pages/id/316/UKCPVS>).

840

841 **Supplementary Table 3: Plant materials analysed for the present study with the**
842 **different *Pst* isolates used for the pathology assays.**

843

844 **Supplementary Table 4: Plant material submitted for Resistance gene enrichment**
845 **Sequencing (RenSeq).**

846 From left to right: Mutant line identifier, targeted gene, score when infected with *Pst*
847 according to the Grassner and Straib scale, mutation position, coverage of the mutation (at
848 least 99% of the reads supported the mutant base in the mutant reads), predicted effect of the
849 mutation on the protein sequence, comments. Lines with the same mutations are highlighted
850 with matching colours.

851

852 **Supplementary Table 5: Genome assemblies used in the present study.**

853 Summary of the available genome assemblies^{58,59} that were used for the *in silico* allele
854 mining and synteny analysis across rice, *Brachypodium*, barley and different Triticeae
855 accessions.

856

857 **Supplementary Table 6: *In silico* allele mining for *Yr7* and *Yr5/YrSP* in available**
858 **genome assemblies for wheat.**

859 Table presents the percentage identity (% ID) of the identified alleles and matching colours
860 illustrate identical haplotypes. Investigated genome assemblies are shown in Supplementary
861 Table 5.

862

863 **Supplementary Table 7: Polymorphisms between *Yr5* protein and its identified alleles.**

864 Positions of the polymorphic amino acids across the five *Yr5/YrSP* proteins. Polymorphisms
865 falling into the BED and NB-ARC domains are shown in red and blue, respectively.

866

867 **Supplementary Table 8: Presence/absence of *Yr7* alleles in a selected panel of Cadenza-**
868 **derivatives and associated responses to different *Pst* isolates (avirulent to *Yr7*: *Pst***
869 **15/151 and 08/21; virulent to *Yr7*: 14/106).**

870 Infection types were grouped into two categories: 1 for resistant and 2 for susceptible. We
871 used Vuka as a positive control for inoculation and absence of *Yr7*. The typical response of a
872 *Yr7* carrier would thus be 1 – 1 – 2, although some cultivars might carry other resistance
873 genes that can lead to a 1 – 1 – 1 profile (e.g. Cadenza). Cultivars that were positive for *Yr7*
874 had either one or the other profile so none of them was susceptible to a *Pst* isolate that is
875 avirulent to *Yr7*. Few cultivars (e.g Bennington, KWS-Kerrin, Brando) were susceptible to
876 one of the two isolates avirulent to *Yr7* in addition to their susceptibility to the *Yr7*-virulent
877 isolate. However, none of them carried the *Yr7* allele.

878

879 **Supplementary Table 9: Presence/absence of *Yr7* and *YrSP* in different wheat**

880 **collections.** We used Vuka, AvocetS and Solstice as negative controls for the presence of *Yr7*
881 and *YrSP* and AvocetS-*Yr* near-isogenic lines as controls for the corresponding *Yr* gene. We
882 genotyped different collections: (i) a set of potential *Yr7* carriers based on literature research,
883 (ii) a set of cultivars that belonged to the UK AHDB Recommended List
884 (<https://cereals.ahdb.org.uk/varieties/ahdb-recommended-lists.aspx>) between 2005 and 2018
885 (labelled 2005-2018-UK_RL), (iii) the Gediflux collection that includes modern European
886 bread wheat cultivars (1920-2010)²⁵, (iv) a core set of the Watkins collection, which
887 represent a set of global bread wheat landraces collected in the 1920-30s²⁴. Most of the
888 putative *Yr7* carriers were positive for all the *Yr7* markers apart from Aztec, Chablis and
889 Cranbrook. Chablis was susceptible to the *Pst* isolates that were avirulent to *Yr7* so it
890 probably does not carry the gene. Regarding the 2005-2018-UK_RL results were consistent
891 across already tested cultivars: Cadenza, Cordiale, Cubanita, Grafton and Skyfall were
892 already positive in Supplementary Table 8. Energise, Freiston, Gallant, Oakley and
893 Revelation were negative on both panels as well. Results were thus consistent across different
894 sources of DNA. *Yr7*-containing cultivars are not prevalent in the 2005-2018 Recommended
895 List set, however, this gene is present in Skyfall, which is currently one of the most harvested
896 cultivars in the UK (Supplementary Table 2). We tested the *YrSP* marker on this set and it
897 was positive only for AvocetS-*YrSP*. The frequency of *Yr7* was relatively low in the Gediflux
898 panel (4%). This is consistent with results in Supplementary Table 2: *Yr7* deployment started
899 in the UK in 1992 with Cadenza and it was rarely used prior to that date. The same was
900 observed in the subset of the Watkins collection (10%) where landraces that were positive for
901 *Yr7* all originated from India and the Mediterranean basin. *Yr7* was introgressed into
902 Thatcher (released in 1936) from Iumillo, which originated from Spain and North-Africa
903 (Genetic Resources Information System for Wheat and Triticum -
904 <http://www.wheatpedigree.net/>). Iumillo is likely to be pre-1920s and these landraces are all
905 bread wheats so they might have inherited it from another source. However, there is no
906 evidence for *Yr7* coming from another source than Iumillo in the modern bread wheat
907 cultivars.

908

909 **Supplementary Table 10: Presence/absence of *Yr5* alleles in selected cultivars.**

910 A subset of the aforementioned collection was investigated for the *Yr5* presence. “Yes” in the
911 *Yr5* column refers to amplification of the 1,281 bp amplicon with the *Yr5*-Insertion primers

912 (Supplementary Figure 6). “Yes” in the *Yr5* alternate alleles column refers to the
913 amplification of the 507 bp amplicon that was identified for AvocetS-*YrSP*, Claire, Cadenza
914 and Paragon in Supplementary Figure 6. “Yes” in the no amplification column refers to
915 identification of a profile similar to the one found for AvocetS in Supplementary Figure 6.

916

917 **Supplementary Table 11: Identified BED-containing proteins in RefSeq v1.0 based on a**
918 **hmmer scan analysis (see Methods).**

919 Several features are added: number of identified BED domains and the presence of other
920 conserved domains present, the best BLAST hit from the non-redundant database of NCBI
921 with its description and score, and whether the BED domain was related to BED domains
922 from NLR proteins based on the neighbour network shown in Supplementary Figure 8.

923

924 **Supplementary Table 12: Transcripts per Million-normalised read counts from the re-**
925 **analysis of published RNA-Seq data²⁹ and associated differential expression analysis**
926 **performed with DESeq2.**

927

928 **Supplementary Table 13: Sequencing details of RenSeq data generated in this study.**

929

930 **Supplementary Table 14: *De novo* assemblies generated from the corresponding**
931 **RenSeq data.**

932

933 **Supplementary Table 15: Primers designed to map and clone *Yr7*, *Yr5*, and *YrSP*.**

934 Note that KASP assays require the addition of the corresponding 5' -tails for the two KASP
935 primers

936

937 **Supplementary Table 16: Diagnostic markers for *Yr7*, *Yr5*, and *YrSP*.**

938 Note that KASP assays require the addition of the corresponding 5' -tails for the two KASP
939 primers.

940

941 **Supplementary Table 17: Passport data of tested *T. dicoccoides* accessions**

942

943 **Supplementary File 1: Annotation of the *Yr7* locus in Cadenza with exon/intron**
944 **structure, positions of mutations and the position of primers for long-range PCR and**
945 **nested PCRs that were carried out prior to Sanger sequencing (Supplementary Table**

946 **15)**. The file also includes the derived CDS and protein sequences with annotated conserved
947 domains. Amino acids encoding the BED domain are shown in red and those encoding the
948 NB-ARC domain are in blue. LRR repeats identified with NLR Annotator are highlighted in
949 dark green and manually annotated LRR motifs xxLxLxx are underlined and in bold black.

950

951 **Supplementary File 2: Annotation of the *Yr5/YrSP* locus in Lemhi-*Yr5* and AvocetS-**
952 ***YrSP*, respectively, with exon/intron structure, the position of mutations and the**
953 **position of primers for long-range PCR and nested PCRs that were carried out prior to**
954 **Sanger sequencing (Supplementary Table 15).** The derived CDS and protein sequences
955 with annotated conserved domains are also shown. Amino acids encoding the BED domain
956 are shown in red and those encoding the NB-ARC domain are in blue. LRR repeats identified
957 with NLR Annotator are highlighted in dark green and manually annotated LRR motifs
958 xxLxLxx are underlined and in bold black. Design of the *Yr5* PCR marker is shown at the
959 end of the file with the insertion that is specific to *Yr5* when compared to *YrSP* and Claire.

960

961 **Supplementary File 3: Curation of the *Yr7* locus in the Cadenza genome assembly based**
962 **on Sanger sequencing results.**

963 Comments show the position of the unknown bases (“N”) in the “Yr7_with_Ns” sequence.
964 Curation based on Sanger sequencing data is shown in bold black in the “curated_Yr7”
965 sequence with the 39 bp insertion and 129 bp deletion. Allele mining for *Yr7* in the Paragon
966 assembly showed that a similar assembly issue might have occurred for this cultivar (same
967 annotation in the “Yr7_Paragon_with_Ns” sequence). This is consistent with the fact that
968 both assemblies were produced with the same pipeline (Supplementary Table 5). We used
969 RenSeq data available for Paragon and performed an alignment as described for the
970 MutRenSeq pipeline against Cadenza NLRs with the curated *Yr7* loci included. A screen
971 capture of the mapping is shown. Only one SNP was identified (75% Cadenza, 25%
972 Paragon). Across the six reads supporting the alternate base, four displayed several SNPs and
973 mapped to an additional Cadenza NLR. This provides evidence for the presence of the
974 identical gene in Paragon which is supported by phenotypic data.

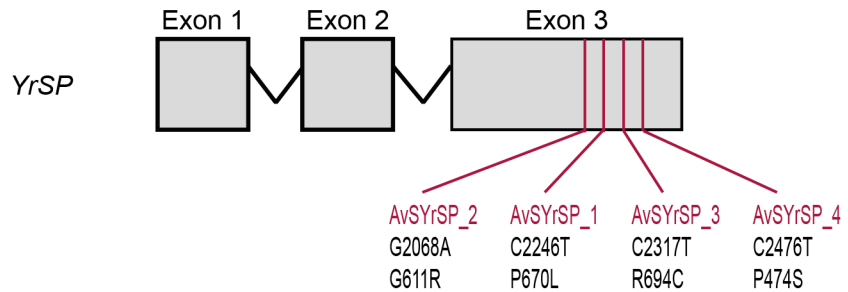
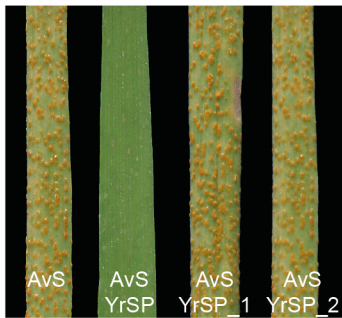
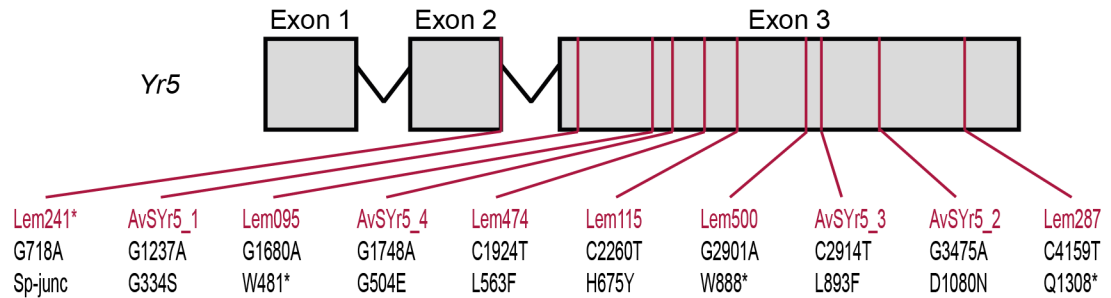
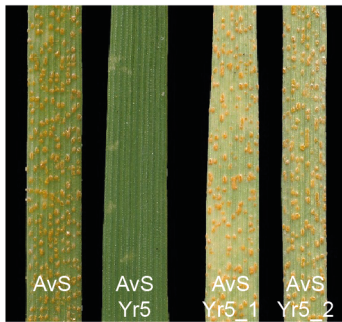
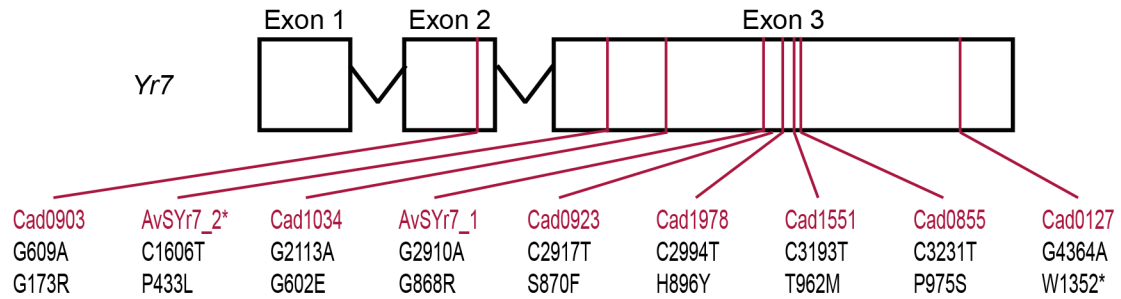
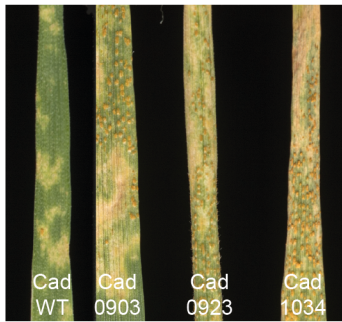
975

976 **Supplementary File 4: Syntenic region across different grasses (Supplementary Table 5)**
977 **and the NLR loci identified with NLR-Annotator.** See Methods for a detailed explanation
978 of the analysis and Supplementary Figure 7 for an illustration.

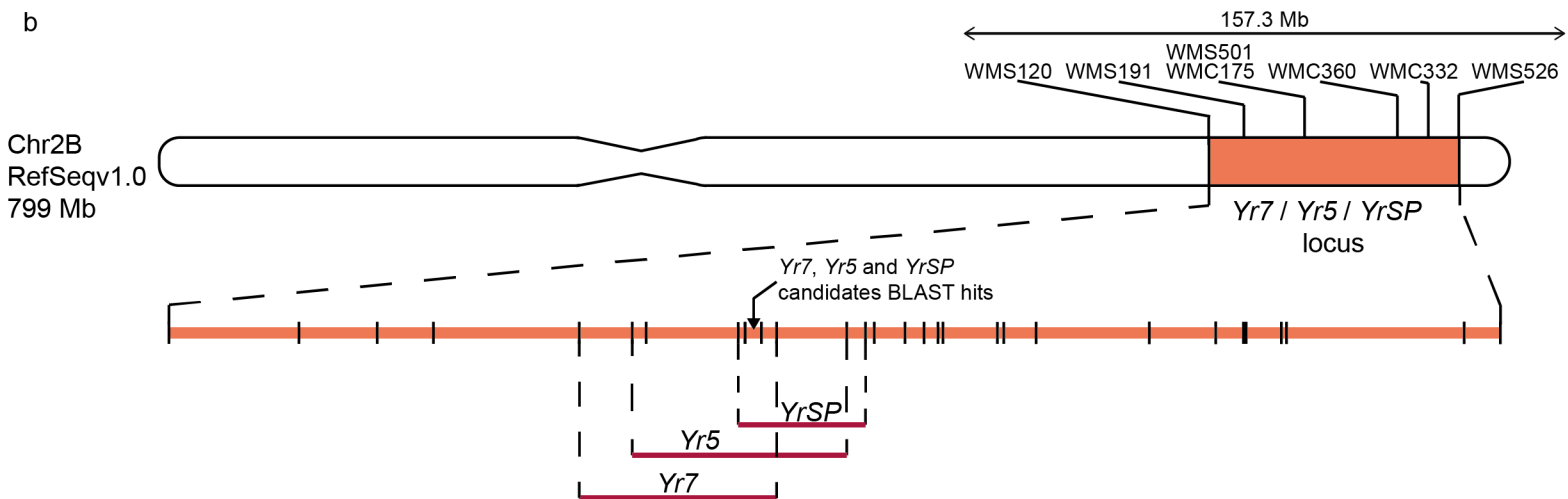
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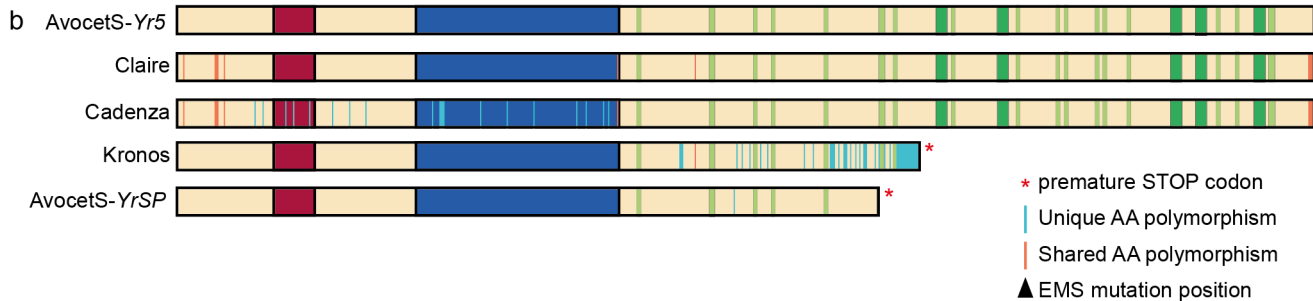
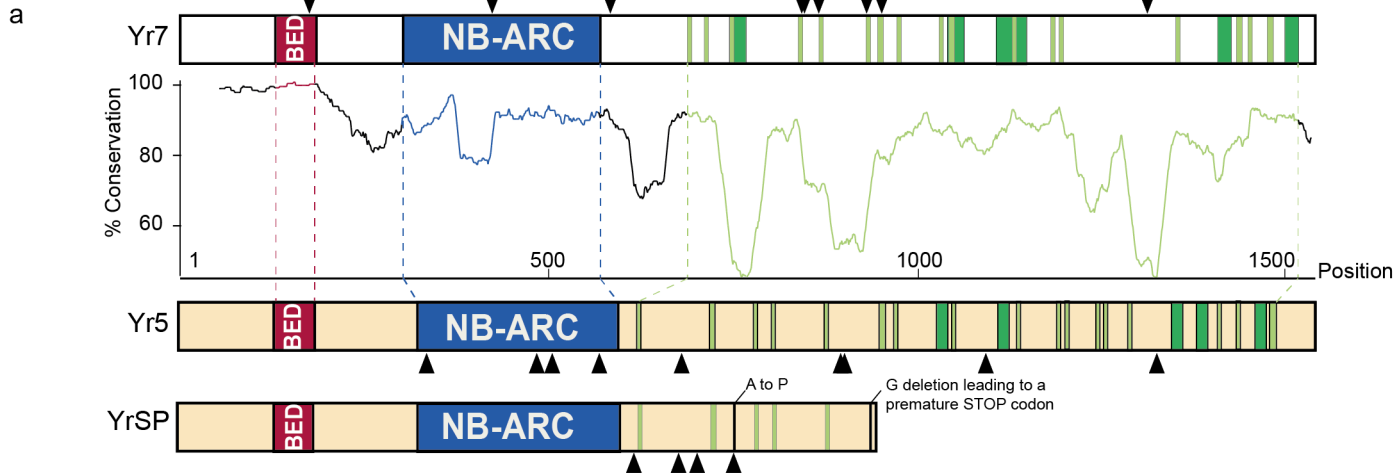
980 **Supplementary File 5: Curated sequences of BED-NLRs from chromosome 2B and**
981 **Ta_2D7.** Exons are highlighted with different colours (yellow, green, blue, pink). Amino
982 acids encoding the BED domain are shown in red and those encoding the NB-ARC domain
983 are in blue. LRR repeats identified with NLR Annotator are highlighted in dark green and
984 manually annotated LRR motifs xxLxLxx are underlined and in bold black.

a



b





a

| Syntenic interval | #NLRs | #BED | | | |
|------------------------------|-------|------|--------|---------|-----------|
| | | NLRs | NLRs-I | NLRs-II | NLRs-I-II |
| Rice | 6 | 2 | - | - | - |
| Brachypodium | 4 | 4 | 1 | 1 | - |
| Barley | 2 | - | - | - | - |
| <i>Aegilops tauschii</i> (D) | 8 | 4 | 1 | 1 | 1 |
| Hexaploid wheat (D) | 6 | 2 | 1 | - | 1 |
| Tetraploid wheat (A) | 8 | 1 | 1 | - | - |
| Hexaploid wheat (A) | 12 | 5 | 3 | 2 | - |
| Tetraploid wheat (B) | 20 | 10 | 6 | 2 | 1 |
| Hexaploid wheat (B) | 13 | 5 | 1 | 1 | 3 |

