Discovery of a novel powdery mildew (*Blumeria graminis*) resistance locus in rye (*Secale cereale* L.)

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Powdery mildew (PM) is one of the most devastating diseases globally. In cereals, the causative agent of PM is the ascomycete fungus *Blumeria graminis* (DC.) speer (*Bg*), which is capable of inflicting severe grain yield losses (≥ 20%) and quality reduction in cereals. In periods of conducive conditions, such as frequent precipitation and low to moderate temperatures, *Bg* can cause severe epidemics by repeated infections and clonal reproduction. As an obligate biotroph, *Bg* is highly specialized to its host species, divided into several distinct ‘formae specialis’ (f. sp.) and dependent on a living host for survival and reproduction. In the later stages of the growing season, or in periods of unconducive conditions, chasmothecia structures are formed. The chasmothecia field inoculum facilitates the infection of volunteer plants or the successive autumn sown winter crop, allowing *Bg* to overwinter as dormant mycelia. Long-distance dispersal by wind of conidiospores drives the large spatial variability in the *Bg* population. In barley (*Hordeum vulgare* L.), the virulence complexity of *B. graminis* f. sp. *hordei* has been observed to increase with the prevailing wind direction from west to east in Europe. Rather than geographical origin, evidence suggests that local use of host resistance and fungicides are the primary factors influencing the European *B. graminis* f. sp. *hordei* population. These findings underline the seriousness of long-distance dispersal of *B. graminis* spores, which allows novel and aggressive pathotypes to evolve through the recombination of distinct pathotypes and rapidly spread. In rye, the causative agent of PM, *B. graminis* f. sp. *secalis*, has attracted little scientific interest in recent years.

To reduce our dependency on pesticides, host resistance constitutes a sustainable alternative for farmers to ensure crop productivity. At present, 23 major PM resistance (*R*) genes have been identified in rye (Table 1). However, it is likely that a subset of colocatalized *R* genes are allelic, as there was no evidence that these genes were distinct from existing *R* genes prior to denotation.

The majority of characterized *Pm* genes encode intracellular nucleotide-binding leucine-rich repeat (NLR) proteins that recognize pathogen effector molecules, leading to an effector-triggered immunity resistance response. Canonical NLR genes are composed of three domains. In grasses, the N-terminus is composed of a coiled-coil (CC) domain believed to be involved in signaling and the induction of cell death. In the center, a nucleotide-binding adaptor shared by APAF-1, R proteins, and the CED-4 (NB-ARC) domain functions as a...
regulatory domain determining the protein activation state. Last, in the C-terminus, a leucin-rich repeat (LRR) domain is involved in effector recognition. In rye, 770 canonical NLR genes have been identified in the ‘Lo7’ reference genome. Rye has been an important source for the improvement of PM resistance in wheat (Triticum aestivum L.) by chromosomal translocation of segments housing R genes. While PM is effectively controlled by host genetic resistance in cereals, it has historically been rapidly overcome by virulent races of Bg. Currently, several of the top-yielding hybrid rye cultivars examined in official Danish trials are susceptible to PM disease (Supplementary Table S1). Genomic-based breeding techniques have, however, dramatically accelerated the introgression and pyramiding of several R genes for enhanced resistance durability. Recent advances in genomic resources available in rye, including the 600 K SNP array and chromosomal-scale reference genome of the German inbred winter rye line ‘Lo7’ and Chinese population rye variety ‘Weining’, constitute significant milestones in rye genomic breeding. Continuous mining for the discovery of novel genetic variability in rye R genes is essential to expand the ‘toolset’ available for resistance breeding. The importance of genetic studies in rye is further supported by the possibility of introgressing novel R genes into the staple cereal wheat by chromosomal translocation lines.

In this paper, we investigate PM resistance in a less-prevalent elite Gülzow-type hybrid rye germplasm. Our objective was to I) characterize PM resistance in the assayed germplasm, II) identify PM resistance-associated SNP markers to be implemented by marker-assisted selection for breeding novel resistant hybrid rye cultivars, III) investigate whether NLR genes residing in PM resistance-associated blocks on the ‘Lo7’ and ‘Weining’ reference genomes resemble known Pm genes, and IV) develop a marker map for the 600 K high-density SNP array and validate its performance in the assayed germplasm.

Results
600 K SNP genotyping of panel. To investigate the genetics underlying powdery mildew (PM) resistance, the assayed hybrid rye breeding germplasm was genotyped on the rye 600 K SNP array. With only scaffold positional data available for the array, SNP marker sequences were anchored to the recent ‘Lo7’ rye reference genome and stringently filtered to ensure its accuracy. In total, 591,196 markers were successfully mapped to the reference genome, and the developed marker map was made available at [https://doi.org/10.5281/ZENODO.5086235](https://doi.org/10.5281/ZENODO.5086235). Quality filtration of markers for low minor allele frequency, missing markers, and missing individual scores across the panel led to the identification of 261,406 informative markers (Supplementary material 1).

| Chromosome | Arm     | Gene name | References |
|------------|---------|-----------|------------|
| 1R         | Short   | Pm1       | Pm8        | 22         |
|            |         | Pm1       | Pm17       | 25         |
|            |         | Pm1       | PmCn17     | 24         |
|            |         | Pm        |            |            |
|            |         | Pm        |            | 29         |
|            | Long    | PmSESY    |            | 30         |
|            | ND      | Pm7       |            | 31         |
| 2R         | Long    | Pm2       |            | 32         |
|            |         | – PmJZIM2RL |          | 33         |
|            |         | Pm8       |            | 34         |
|            | ND      | Pm7       |            | 35         |
|            |         | Pm8       | Pm3        | 31         |
|            |         | Pm        |            | 36         |
| 3R         | Short   | Pm3       |            | 37         |
| 4R         | Long    | Pm         |            | 38         |
|            |         | – Pm56    |            | 39         |
|            | ND      | Pm         |            | 40         |
| 5R         | Long    | Pm4       |            | 41         |
|            | Short   | – Pm56    |            | 39         |
| 6R         | Long    | Pm5       |            | 42         |
|            |         | – Pm20    |            | 40         |
| 7R         | Long    | Pm        |            | 41         |
|            |         | Pm        |            | 40         |

Table 1. Location and origin of powdery mildew resistance genes in rye (Secale cereale L.) and wheat-(Triticum aestivum L.) rye translocation lines with rye being the donor parent. After1–3. ND: Interchromosomal position not determined.
marker-to-marker distance of 25.54 kb. The largest marker-to-marker distance was 9.95 Mbp on chromosome 2R, with a mean of 4.05 Mbp across the chromosomes. As a quality parameter, the polymorphism information content (PIC) was calculated to estimate the ability of markers to detect polymorphisms within the assayed germplasm (Supplementary Table S2). Across the informative marker panel, a mean PIC of 0.234 was identified, with a mean interchromosomal PIC ranging from 0.204 to 0.249 (Table 2). Visualization of these array performance metrics along the rye genome using Circos revealed a drop in marker density and PIC across the pericentromeric region on all chromosomes (Fig. 1).

**Phenotyping of hybrid rye breeding germplasm.** To provide a comprehensive phenotypic dataset of PM resistance in the assayed Gülzow-type hybrid rye breeding germplasm, the lines were scored for their infection type (IT) against three distinct *Blumeria graminis* f. sp. *secalis* (Bgs) populations (Table 3, Supplementary Table S3). The lines scoring an IT below 1 were considered ‘resistant’ (Supplementary Fig. S1). Across the assayed germplasm, the N13 (Nienstädt, 2013) Bgs population yielded a mean IT of 2.40 ± 1.28 standard deviations (SD), with 47 resistant lines out of which 3 were restorers. The N18 (Nienstädt, 2018) Bgs population yielded a mean IT of 2.71 ± 1.11 SD with 29 resistant lines, out of which 1 was a restorer. D20 (Dyngby, 2020) Bgs population yielded a mean IT of 2.74 ± 1.10 SD with 20 resistant lines out of which 1 was a restorer. Across the assayed germplasms, 20 out of 88 non-restorer germplasm and 1 out of 92 restorer lines were consistently resistant to all three Bgs populations. Both controls, hybrid cv. KWS Binntto (‘susceptible’) and KWS Serafino (‘resistant’) were susceptible to all three Bgs populations. KWS Binntto had a mean IT of 3.56 ± 0.54 SD and KWS Serafino 3.01 ± 1.01 SD across Bgs populations.

To visualize the resistance spectrum of breeding lines, a circular neighbor-joining dendrogram was constructed, and concentric circles were added to integrate the scored IT (Fig. 2). Statistical analysis of the line infection-type distribution across Bgs population by ANOVA showed that the N13 Bgs population differed significantly from the N18 and D20 Bgs populations (*p*val < 0.00016) (Fig. 3). Nine non-restorer germplasm lines were found to exhibit a differential resistance response to the N13 Bgs population. These lines were categorized as ‘partially resistant,’ with a mean infection type of 1.52 ± 0.58 SD; they exhibited a mean infection type of 2.74 ± 0.70 and 2.80 ± 0.77 against the N18 and D20 populations, respectively.

**Genome-wide association study.** For the identification of SNP markers associated with PM resistance, GWAS using MLM and the BLINK method was used for each of the three Bgs population trials and across trials for the entire germplasm and individual parental populations (Supplementary Figs. S2, S3). Isolated single SNPs in GWAS-MLM results were removed from the analysis even if they were significantly associated, as these were interpreted as having spurious associations or incorrect mapping positions. Instead, dense peaks comprising a large number of significant associated markers within a confined region in GWAS-MLM were selected for further analysis. GWAS-MLM on the entire panel led to the identification of a haplotype block on chromosome arm 7RL that was significantly associated (−log10 = 14.6) with powdery mildew resistance spanning from 882 to 898 Mbp (Fig. 4B, C). The haplotype block harbored 244 markers exhibiting an association above the Bonferroni adjusted significance threshold (−log10 ≥ 6.7) (Supplementary Table S4). Successive GWAS-BLINK led to the identification of the top-most resistance-associated (−log10 = 37.9) marker within the haplotype block on chromosome arm 7RL at 892.09 Mbp, which explained 16.8% of the phenotypic variance (Fig. 4A).

In the non-restorer germplasm population, lines were found to carry a highly conserved haplotype across the significantly associated markers on chromosome arm 7RL (Supplementary Table S6). A resistant haplotype was conserved in 42 out of 48 resistant lines and a susceptible haplotype was found in 28 out of 31 susceptible lines, significantly associated markers on chromosome arm 7RL (Supplementary Table S6). A resistant haplotype was significantly associated markers on chromosome arm 7RL (Supplementary Table S6). A resistant haplotype was found in 28 out of 31 susceptible lines, while a nonsignificant (−log10 = 3.62) peak was identified in GWAS-MLM spanning from chromosome arm 7RL at 892.09 Mbp, which explained 16.8% of the phenotypic variance (Fig. 4A).

**Table 2.** Characteristics of informative 600 K SNP array on the Nordic Seed hybrid rye (*Secale cereale* L.) elite breeding germplasm (n = 180). Markers were positioned on the ‘Lo7’ reference genome. SD: Standard deviation, PIC: Polymorphism information content.
743.9 to 754.4 Mbp on chromosome arm 5RL (Supplementary Fig. S2, Supplementary Table S4). Gene mining by GWAS in the restorer population alone was not performed due to the low number of resistant lines (n = 5).

**Nucleotide-binding leucine-rich repeat proteins in powdery mildew resistance-associated blocks on chromosome arm 7RL.** Annotation of NLR genes in the reference genomes ‘Lo7’ and ‘Weining’ led to the identification of 770 and 1027 full-length (‘complete’) NLR genes, respectively. Positional information, annotation, and NB-ARC and NLR sequences of all ‘complete’ and ‘partial’ NLR genes for both reference genomes have been made available at https://doi.org/10.5281/zenodo.5085854. The PM resistance-associated
A block residing in the subtelomeric region of chromosome arm 7RL spanned 17 Mbp on the ‘Lo7’ reference genome and harbored 25 NLR genes (Fig. 4C, Supplementary Table S8). Out of the 244 markers residing in the block significantly associated (− log_{10}(p) > 6.72) with PM resistance, 155 were accurately positioned on the ‘Weining’ reference genome (Supplementary Table S7). The markers mapped to a site spanning 14 Mbp from 994.6 to 1,008.4 Mbp on chromosome arm 7RL and housed 16 NLR genes (Supplementary Table S8). A search in the protein database using the NCBI Blastx function showed that the majority of NLRs shared similarities with resistance gene analogs (RGAs) and Pik-2-like and Pik6-NP-like disease resistance proteins in the diploid parental species of wheat. In both reference genomes, two NLRs shared similarities with Rpp13-like disease resistance proteins in *Triticum urartu* and barley.

Figure 2. Circular neighbor-joining dendrogram of 180 hybrid rye (*Secale cereale* L.) breeding lines. Infection type (0–4) reaction against three powdery mildew populations displayed by concentric circles around the dendrogram.

Figure 3. Density plot of the infection type distribution across 180 Nordic Seed hybrid rye (*Secale cereale* L.) breeding lines against three powdery mildew populations.
Phylogenetic analysis using the NB-ARC domain of NLRs residing in the PM resistance-associated haplotype block on chromosome arm 7RL led to the finding that the majority of the NLRs were represented in both reference genomes (Fig. 5). The 'Weining' reference genome exhibited one unique NLR not present in 'Lo7', while the latter exhibited eight unique NLRs, of which five formed a distinct clade. Eight homogeneously represented NLRs from each reference genome clustered in a large clade ('clade 1').

An additional phylogenetic analysis was performed using the entire reference genome NLR repertoire including NLR genes of characterized R genes as references (Fig. 6, Supplementary Fig. S4). The NLR genes residing within the haplotype block were found to span much of the reference NLR repertoire diversity. Clade 1 remained intact in both reference genome NLR repertoire trees positioned in a section harboring four out of five reference Pm genes included, with the closest being Pm60. In both reference genome NLR repertoire trees, a single
NLR (Lo7_chr7R_nlr_94, Wei_chr7R_nlr_139) showed an evolutionary relationship with an Rpp13-like disease resistance gene and four Mla alleles.

**Characterization of an Rpp13-like NLR gene residing in the resistance-associated block on chromosome arm 7RL.** While several of the NLR genes residing within the PM resistance-associated haplotype block on chromosome arm 7RL chromosome showed evidence of an evolutionary relationship with known Pm genes, we selected the Rpp13-like NLR gene for further investigation on the basis of its close proximity. The Rpp13-like NLR gene (Lo7_chr7R_nlr_94, Wei_chr7R_nlr_139) was the closest NLR gene, residing approximately 2 Mbp from the top-most PM resistance-associated marker identified by BLINK (Fig. 4C, Supplementary Table S8).

The homologous Rpp13-like NLR genes in the 'Lo7' and 'Weining' reference genomes encoded canonical NLR proteins of 922 to 947 aa, showing 97% sequence similarity, with sequences differing by three single amino acid variants and an indel of 25 aa in the NB-ARC domain (Supplementary material 2). Protein BLAST of the NLR gene led to 88% sequence similarity with disease resistance Rpp13-like protein 4 in *T. urartu*, which encodes a 924 aa NLR protein sequence.

**Discussion**

In Denmark, the top-yielding hybrid rye cultivars and population varieties were evaluated in Danish official trials as an advisory service for farmers. In the last decade, a high level of powdery mildew infection in rye was recorded only once in the trials in 2017 at a site in southern Denmark (Supplementary Fig. S1). At the trial site, several hybrid rye cultivars displayed up to 18% leaf area covered by PM. While no recent studies have investigated the effect of powdery mildew in top-yielding hybrid rye cultivars, Matzen et al. (2019) reported a 16% yield reduction in triticale at a disease severity of ≤10% leaf area covered by powdery mildew under field conditions in 2017. It is, therefore, reasonable to presume that under certain conducive conditions, PM is capable of causing significant yield loss.
of causing substantial grain yield and quality losses in rye. Furthermore, on the basis of the trial records, we selected a hybrid cv. KWS Serafino as a resistant control in our study, as it showed less than 1% leaf area covered by PM at the site in 2017. However, under a high disease pressure, cv. KWS Serafino was found to be susceptible in the current study, suggesting a potential lower level of resistance against PM in the evaluated top-yielding hybrid cultivars than evident from the official trials.

In this study, we investigated PM resistance in the less-prevalent Gülzow-type elite hybrid rye breeding germplasm against three distinct *Blumeria graminis* f. sp. *secalis* (*Bgs*) populations from Denmark and Northern Germany. We observed a moderate level of powdery mildew resistance in the non-restorer germplasm population, and by performing a genome-wide association study (GWAS) using 261,406 informative SNP markers, we identified a strong PM resistance-associated site on the distal region of chromosome arm 7RL (Lo7–7RL PM) are colored teal.

Figure 6. Phylogenetic tree of 770 nucleotide-binding leucine-rich repeat (NLR) protein NB-ARC domains in the ‘Lo7’ rye (*Secale cereale* L.) reference genome. The NB-ARC domains of known NLR genes have been included as references. NLR genes residing in a powdery mildew resistance block in the subtelomeric region of chromosome arm 7RL (Lo7–7RL PM) are colored teal.
inbred lines yielded 362 K informative SNP markers with a mean PIC of 0.25. In this study, however, we did observe a drop in marker informativeness and density across the pericentromeric region. Similar observations were made in the study by Rabanus-Wallace, et al., who reported a considerable reduction in both genetic diversity and gene density within the pericentromeric region of the inbred rye line 'Lo7'. In barley, this region has furthermore been observed to display a 20-fold lower recombination rate. Thus, we concluded that the 600 K SNP array performed satisfactorily on the Gölzow-type germplasm in relation to the fundamental characteristics investigated. The high-density array constitutes a milestone in rye genomic resources transitioning SNP-based genomic studies and replaces the previous rye 5 K SNP array by Haseneyer, et al.

GWAS on SNP genotypic data has become an effective tool in genome-based plant breeding for the study of oligogenic traits often governed by a few genes with large effects, such as the major monogenic inherited resistance (R) genes. The use of high-density SNP typing has allowed whole-genome scans for the identification of often small haplotype blocks significantly associated with resistance. The creation of the 600 K SNP array and the chromosomal-scale reference genomes 'Lo7' and 'Weinig' in rye has significantly changed the genomic toolbox available for mining novel resistance genes. Using these recent advances in rye genomic resources, we successfully managed to identify a site on chromosome arm 7RL that was significantly associated with PM resistance. The high level of resolution provided by the dense marker panel revealed that the PM resistance-associated haplotype block spanned 17 Mbp on the distal tip of the chromosome arm 7RL subtelomeric region.

Until recently, no PM resistance gene had been identified on the rye 7R chromosome. However, during the development of translocation lines for wheat improvement using a local Chinese variety of rye 'Baill', Ren, et al. discovered a PM-resistant 7BS:7RL translocation line. With the recipient wheat parent being susceptible, the PM resistance gene traced back to the rye donor chromosome arm 7RL. Resistance phenotyping of the translocation line demonstrated that it displayed a high level of PM resistance against prevalent Bg. triticici pathotypes in China, making it very promising for the development of novel PM-resistant wheat cultivars. Although a novel finding in rye, several PM resistance genes have been identified in wheat chromosomal segments syntenic to rye chromosome arm 7RL. During Triticeae speciation, a series of recurrent translocation events gave rise to major patterns of chromosomal rearrangements. In rye, the distal region of chromosome arm 7RL, therefore, shows high homology with the wheat 2A/B/D chromosomes. Currently, more than five PM genes have been identified in wheat on the 2A/B/D chromosomes.

With the PM gene discovered by Ren, et al., originating from a forage-type population of rye in a gene pool distinct from the germplasm investigated in this study, it is reasonable to presume that the two PM genes could be either distinct or allelic. We provisionally denote the novel PM locus residing in the subtelomeric region of chromosome arm 7RL as PmNOS1.

Comparative analysis of the three Bgs populations used in the study revealed that the N13 population significantly differed from the two more recent populations, showing a less virulent composition of pathotypes. In addition, nine non-restorer germplasm breeding lines displaying a differential resistance profile were identified, showing 'partial resistance' only to the N13 population. With none of these lines carrying the resistance haplotype associated with the resistance locus on chromosome arm 7RL, our findings suggest that these do not carry the PmNOS1 locus. While this observation can be explained by the occurrence of recombination events between the PM resistance-associated marker and the causative gene, this result seems less likely due to the consistent divergence in the haplotype. Instead, it seems more likely that the differentially resistant non-restorer germplasm lines carry a distinct PM gene that lost its effect during the period 2013–2018 in northern Germany. As a result of host genetic uniformity and the high evolutionary capacity of Bg to acquire virulence and migrate rapidly over long distances, the effectiveness of PM genes is often rapidly lost. An example of this is the Mla13 PM gene introduced in a former Czechoslovakian barley cultivar, 'Koral', in 1980. After years of providing effective resistance against PM disease, virulence was observed in England in 1988 in a pathotype believed to originate from Czechoslovakia, having migrated by wind across the European continent and North Sea. Ongoing monitoring of the virulence structure of Bg was conducted to survey the effectiveness of deployed PM genes in elite cultivars. While the PM gene present in the differentially resistant non-restorer germplasm lines has seemingly been overcome, our findings suggest that PmNOS1 remains effective.

Enabled by recent advances in rye genomic resources, we investigated whether any of the NLR genes residing in the region harboring the PmNOS1 locus resembled known PM genes. In several crop species, including rye, NLR genes have been observed to accumulate at recombination hotspots in subtelomeric chromosomal regions. Additionally, we identified several large clusters of NLR genes residing in the PM resistance-associated block harboring PmNOS1 in both of the reference genomes. In addition to the PM resistance gene identified in our study, Fusarium head blight and leaf rust resistance genes have been mapped to the subtelomeric region of chromosome arm 7RL. Stem and stripe rust resistance genes reported in the 7BS:7RL translocation line developed by Ren, et al. are furthermore likely to reside in the subtelomeric region. To investigate the likely diversity of R genes residing in the PM resistance-associated block, we conducted a phylogenetic analysis using the NB-ARC domain sequence of NLR genes residing in the block. In contrast to the rapidly evolving LRR domain often exhibiting intraspecific polymorphism, the NB-ARC domain is largely conserved and suited for the study of evolutionary relationships among NLR genes. As expected, the NLR genes residing in the block represented a large proportion of the NLR repertoire diversity in rye, accentuating the evolutionary plasticity of the NLRs residing in the subtelomeric region. Intriguingly, guided by a panel of isolated NLR genes as a reference, we observed a predisposition of NLRs within the block to be in close proximity to known PM genes in wheat and barley. This evolutionary relationship could hint at a common attribute among the NLRs.

Phylogenetic analysis led to the identification of an NLR gene with an evolutionary relationship and protein sequence similarity to Rpp13-like protein 4 in T. urartu. Based on its close proximity to the marker displaying the strongest association with the PmNOS1 locus, the Rpp13-like NLR gene was selected for further characterization. In A. thaliana, Rpp13 confers resistance against Peronospora parasitica, the causative agent of downy mildew
Recent studies have, however, identified *Rpp13*-like NLR genes associated with powdery mildew resistance in cereals. In wheat, Liu et al. showed that silencing of the *Rpp13* homologous gene *TaRPP13-3* in resistant wheat cv. ‘Brock’ induced susceptibility to powdery mildew. In barley, Cheng et al. found that the expression of an *Rpp13*-like NLR gene was highly upregulated after inoculation with powdery mildew.

In rye, while the rate of decay has only been determined in a few genes related to frost response, which showed a rapid linkage decay, these genes have been demonstrated to decay after 3.76 kb, on average, across the genome in a similar hybrid breeding germplasm in maize. Due to the heterogenic nature of outcrossing species, their rate of decay is often rapid. However, in a recent population study on assayed germplasm, non-restorer germplasm was found to exhibit relatively low genetic diversity, low effective population size and high linkage disequilibrium. Consistent with their observations, we found a large conserved haplotype on chromosome arm 7RL harboring the *PmNOS1* locus. The linkage decay in the non-restorer germplasm is likely considerably reduced by the low genetic diversity and effective population size, resulting in a similarly low frequency of effective recombination events. The large amount of linkage, while beneficial for trait discovery in GWAS at lower marker density, impedes the identification of a narrow and precise genomic region that may harbor the gene of interest, even at high marker resolution. In the case of the *PmNOS1* locus, the haplotype block on chromosome arm 7RL was found to span 17 Mbp, harboring between 17 and 25 potential candidate NLR genes in the ‘Lo7’ and ‘Weining’ reference genomes. For more accurate mapping of the *PmNOS1* locus, the development of multiparent mapping populations could be conducted, allowing several generations of potential effective recombination events in the region.

Identification of the causative gene could be performed by resistance gene enrichment sequencing (RenSeq) analysis followed by transformation of a susceptible non-restorer germplasm line to validate the gene. This would in turn equally show whether the gene is present in the reference genomes and whether *PmNOS1* encodes an *Rpp13*-like NLR protein. In conclusion, our study demonstrates the immediate value of recent advances in rye genomic resources for the mining of novel resistance genes. These resources now permit accurate identification of delimited resistance-associated haplotype blocks and scanning for trait-associated genes residing within. With pathogens such as *Bgh* displaying a large evolutionary plasticity, shortening the process from identification of resistance-associated sites to isolation of the underlying *R* gene is important for the development of novel resistant cultivars. The relevance of studies in rye is accentuated by the possibility of introgressing novel *R* genes into the staple cereal wheat by chromosomal translocation lines. To aid further studies in the field, we have provided both a rye600K SNP array marker map anchored using the ‘Lo7’ reference genome and NLR repertoire information of the ‘Lo7’ and ‘Weining’ reference genomes in open-access data repositories.

### Materials and methods

#### Plant material and DNA extraction.
A panel of 180 inbred rye (*Secale cereale* L.) lines, 92 restorer and 88 non-restorer germplasm, belonging to the elite Gülzow-type hybrid rye breeding germplasm at Nordic Seed A/S (Dyngby, Denmark) were investigated. Population structure and information on the genetic characteristics of the germplasm were presented in a recent study by Vendelbo et al. The parental populations represent genetically secluded gene pools with restorer (paternal) lines carrying a dominant allele for the restoration of male fertility and non-restorer germplasm lines carrying a recessive allele and fertile cytoplasm, which is used to maintain cytoplasmic male sterile (maternal) lines. DNA extraction was performed using an adapted SDS-based method according to the USDA after Pallotta et al. on an equivalent of 75 mg of plant material collected from the coleoptiles and primary leaves of two seven-day-old seedlings per line. DNA concentration and 260/280 nm ratio of the samples were measured using an Epoch microplate spectrophotometer (Biotek), and evidence of fragmentation was obtained by size visualization on a 1.2% agarose gel.

#### Molecular marker resource and SNP genotyping.
Samples of each line containing 200 ng of high molecular weight gDNA with a ≥1.8 260/280 nm ratio were sent for single nucleotide polymorphism (SNP) genotyping at Eurofins Genomics Europe Genotyping (Aarhus, Denmark). Genotyping was performed using a 600 K SNP array with 600,843 SNP markers on an Affymetrix GeneTitan™ Scanner platform.

#### Collection and multiplication of *Blumeria graminis* f. sp. *secalis* populations.
As there was no unified information on germplasm resistance to powdery mildew, three field populations of *Bgs* were sampled in Northern Germany and Denmark in the period from 2013 to 2020 to screen the lines against a broad range of *Bgs* pathotypes prevalent in Northern Europe. Two *Bgs* populations were collected at the Nordic Seed rye multiplication site in Germany in 2013 (N13) and 2018 (N18) (52.29254°N, E9.14896°E). Ten leaves exhibiting PM disease were carefully collected and rinsed in 0.5 L of water to release *Bgs* spores. Six pots containing 15–20 susceptible 12-day-old seedlings of the restorer line R277 were then inoculated by spraying a fine mist of the spore solution using an atomizer bottle. Pots were transferred to a separate climate chamber for each population and incubated at 18 °C with 12 h of light using 400 W high-pressure Phillips SON-T Agro lamps. Populations were continuously multiplied in an overlapping two-week cycle. Each week, the tray of 2-week-old seedlings was substituted with a tray of fresh 12-day-old seedlings. Inoculation was achieved by passive dissemination of spores from the ‘older’ tray to the tray with new plants through a steel-grid shelf.

An additional *Bgs* population was collected in autumn 2020 (D20) at Nordic Seed (Dyngby, Denmark) (55.94944°N, E10.25414°E) using a mixture of hybrid cvs. KWS Binningt, KWS Bono and KWS Florano. Pots with 15 to 20 seedlings of the susceptible mixture were placed outdoors 12 days after sowing (DAS) in August-October 2020. Plants were controlled regularly, and all leaves showing PM disease within the period were collected. Prior to inoculation, leaves were placed in Petri dishes with moist filter paper and set to sporulate at room temperature in light for four to eight hours. Then, a pot containing 15–20 seedlings of the susceptible mixture was inoculated.
at 12 DAS by horizontally stroking the collected leaves across the seedlings. Inoculated pots were sprayed with a fine mist of water, placed in a container with transparent lids to ensure 100% RH and incubated in the dark at 10–15 °C for 24 h. After incubation in the dark, the pots were transferred to an isolated greenhouse cabin and incubated under 16 h of daylight at 18–24 °C and 8 h of dark at 14–16 °C.

Multiplication of trial inoculum for the three Bgs populations was performed using the same procedure as described above. For each population, two trays containing 35 pots of the susceptible mixture were inoculated by brushing 3–4 highly infected pots inoculated 20 days earlier across the tray.

**Infection and scoring.** All lines were phenotyped for the infection-type response to the Bgs populations. In each greenhouse trial, eight seeds per line were sown in a 28-hole tray using a completely randomized design with two repetitions for each of the two trial replicates. For each tray, a positive (‘susceptible’) control consisting of hybrid cv. KWS Binntto and negative (‘resistant’) control cv. KWS Serafinno was included. At 14 DAS, trays were inoculated as described above by brushing 3–4 highly infected pots inoculated 20 days earlier across the tray. After 14 days of incubation, the lines were phenotyped by scoring the infection response on the first and second leaves for each of the eight seedlings per repetition in accordance with a 9-step 0–4 scale by Torp et al.88 (Table 3).

**Data analysis.** Bioinformatic analysis of SNP marker data was performed with the R studio (v. 1.3.959) interface in R statistical software (v. 4.0.1) by applying various predesigned packages92,93.

**Mapping of 600 K SNP array markers to the ‘Lo7’ reference genome.** Positional data of the 600 K SNP markers were obtained by mapping each of the 600,843 SNP marker sequences to the rye reference genome ‘Lo7’ using the NCBI blastn (v. 2.9.0 +) function10,94. The mapping positions of SNPs were hereafter stringently filtered for I) complete SNP sequence alignment and II) a maximum of 1 mismatch to ensure accurate positioning.

**Molecular markers and characterization of 600 K SNP array performance.** Prior to analysis, markers were filtered for a marker allele frequency ≥ 0.005, missing individual score ≤ 0.2 and missing marker score ≤ 0.1. Fundamental characteristics of SNP marker informativeness, including polymorphism information content (PIC), were calculated using the SnpReady (v. 0.9.6) R package95. The interchromosomal distribution of the informative marker PIC, marker-to-marker distance and marker density in 10 mb bins on the ‘Lo7’ rye genome were visualized using Circos (v. 0.69.8) in the Galaxy online interface96,97. A Circos plot was constructed using the pipeline developed by Hillemann, et al.98.

**Analysis of phenotypic data.** The distribution of infection types against the three respective PM populations was visualized by density plots using ggplot2 (v. 3.3.3) R package86; To determine whether the population infection-type distribution differed significantly, ANOVA was conducted using R.

To correct the resistance phenotype for the effects of replication and population, we fitted the data to a linear mixed model using the lme4 (v. 1.1.26) package in R:

\[ y = \mu + P + R + I + \epsilon \]

where \( \mu \) is the general mean, \( P \) is the population, \( R \) represents the replications, \( I \) is the line id, and \( \epsilon \) is the residuals. \( P \) and \( R \) were set as fixed effects, and \( I \) was set as a random effect. The random effect and residuals were assumed to be independent normally distributed variables described as follows: \( I \sim N (0, I \sigma_i^2) \), and \( \epsilon \sim N (0, I \sigma_e^2) \). The BLUP solutions for the line effect were used in GWAS. Data were also fitted to a linear mixed model for each of the populations to correct for the effect of replication. In this model ‘\( P \)’ was removed.

**Genome-wide association study.** Discovery of PM resistance-associated SNP markers was performed by a genome-wide association study (GWAS) using the genomic association and prediction integration tool (GAPIT) (v.3) package in R100. The Manhattan plot was colorized using the RColorBrewer (v.1.1–2) R package color palette99. GWAS using a mixed linear model (MLM) was performed to identify discrete haplotype blocks associated with powdery mildew resistance. Additionally, the Bayesian-information and linkage-disequilibrium iteratively nested keyway (BLINK) method was performed to identify the top-most powdery mildew resistance-associated marker within each haplotype block101. BLINK uses a multiple loci test for MLM by combining a fixed effects model, Bayesian information content and linkage disequilibrium information to collectively improve the statistical power while simultaneously reducing the computational run time. Markers that are in linkage disequilibrium with the top-most significant marker at a site are excluded in BLINK. A standard Bonferroni-corrected threshold of \( \alpha = 0.05 \) was used as the significance threshold.

**Annotation of nucleotide-binding leucine-rich repeat proteins in the reference genome.** To investigate whether PM-associated sites in the ‘Lo7’ and ‘Weining’ reference genomes housed nucleotide-binding leucine-rich repeat (NLR) genes, annotation was performed using NLR-parser (v.3) and NLR-annotator (https://github.com/steue rnb/NLR-Annnotator)99,102.

**Characterization of nucleotide-binding leucine-rich repeat proteins.** The gene structure, coding sequence and NLR protein sequence of candidate genes were extracted from RNA-seq and de novo protein data provided in ‘Weining’ and ‘Lo7’ reference genome data repositories96,98. To investigate whether NLR genes residing in
PM resistance-associated sites resembled known genes, the NCBI blastx function was used for protein–protein searches in the online database. For functional analysis and the prediction of protein domains, InterPro Scan was used. Identification of sequence divergence between reference genome homologs was performed by multiple sequence alignment using the multiple sequence comparison by log-expectation (MUSCLE) method for coding sequences and the ‘Clustal Omega’ method for NLR protein sequences in Geneious Prime (v. 2020.2.3).

**Phylogenetic analysis.** Neighbor-joining clustering analysis of breeding lines was performed with Euclidean genetic distance measurement using the ape (v. 5.3) R package. The tree was constructed after 10,000 bootstrapping iterations with weak nodes (<80% recurrence) collapsed into multifurcations. A circular neighbor-joining tree was generated using the iTOL (v. 5) online tool (http://itol.embl.de/), enabling a colorful visualization of each line infection type spectrum against the three Bgs populations.

To investigate the sequence similarity between nucleotide-binding leucine-rich repeat (NLR) genes at PM resistance-associated sites in the ‘Lo7’ and ‘Weining’ reference genomes, neighbor-joining clustering analysis using NB-ARC (nucleotide-binding adaptor shared by APAF-1, R proteins, and CED-4) domain sequences was performed. Identification of PM resistance-associated sites on ‘Weining’ was performed by mapping PM resistance-associated markers using the same procedure as described previously for ‘Lo7’. As references, NB-ARC domain sequences of available leaf rust (Lr1, Lr10, Lr21, Lr22a), stem rust (Sr13, Sr22), yellow rust (Yr5, Yr10, Yr28), powdery mildew (Pm2, Pm3, Pm8, Pm12, Pm17, Pm60, Mla1, Mla6, Mla7, Mla10, RPP13, RPP-like-T. urartu) and resistance gene analogs (RGA1, RGA2, RGA3, RGA4, RGA5) were included. NB-ARC sequences were obtained from the UniProt online database. Phylogenetic analysis was conducted using a pipeline developed by Topparslan, et al. in R. Multiple sequence alignment of NB-ARC domain sequences was performed via msa (v. 1.20.1) using the ‘Clustal Omega’ method and pairwise genetic distance based on identity calculated with the seqfinr (4.2–8) package in R. Phylogenetic analysis was conducted using a pipeline developed by Topparslan, et al. (2020) in R. Multiple sequence alignment of NB-ARC domain sequences was performed via msa (v. 1.20.1) using the ‘Clustal Omega’ method and pairwise genetic distance based on identity calculated with the seqfinr (4.2–8) package in R. A tree was constructed for the ‘Lo7’ and ‘Weining’ NLR repertoires separately and visualized using ggtree (v. 2.2.4) R package.

**Graphical editing.** Graphs and figures were outputted from R in .svg format and manually curated using Inkscape (v. 1.1) (https://inkscape.org/).

**Ethical statement.** Experimental research and field studies on plants (either cultivated or wild), including the collection of plant material, complies with relevant institutional, national, and international guidelines and legislation.

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**References**

1. Schlegel, R. H. J. Rye genetics, breeding and cultivation (CRC Press Taylor & Francis Group, 2014).
2. Schlegel, R. H. I. & Korzun, V. Genes, markers and linkage data of rye (Secale cereale L.), 11th updated inventory, V. 0.1.21. 1–115, http://www.rye-genetics-map.de (2021).
3. Crespo-Herrera, L. A., Garkava-Gustavsson, L. & Åhman, I. A systematic review of rye (Secale cereale L.) as a source of resistance to pathogens and pests in wheat (Triticum aestivum L.). Hereditas 154, 1–9. https://doi.org/10.1186/s11686-017-0033-5 (2017).
4. Savary, S. et al. The global burden of pathogens and pests on major food crops. Nat. Ecol. Evolut. 3, 430–439 (2019).
5. Savary, S., Ficke, A., Aubertot, J.-N. & Hollier, C. Crop losses due to diseases and their implications for global food production losses and food security. Food Secur. 4, 519–537. https://doi.org/10.1007/s12571-012-0020-5 (2012).
6. Griffer, C. A., Das, M. K. & Stromberg, E. L. Effectiveness of adult-plant resistance in reducing grain yield loss to powdery mildew in winter wheat. Plant Dis. 77, 618–622 (1993).
7. Conner, R. L., Kuzyk, A. D. & Su, H. Impact of powdery mildew on the yield of soft white spring wheat cultivars. Can. J. Plant Sci. 83, 725–728 (2003).
8. Matzen, N., Heick, T. M. & Jørgensen, L. N. Control of powdery mildew (Blumeria graminis spp.) in cereals by Serenade® ASO (Bacillus amyloliquefaciens (former subtilis) strain QST 713). Biol. Control 139, 104067 (2019).
9. To Beest, D. E., Paveley, N. D., Shaw, M. W. & Van Den Bosch, F. Disease-weather relationships for powdery mildew and yellow rust on winter wheat. Phytopathology 98, 609–617 (2008).
10. Tsuchiya, A. et al. Phenotypic characterization of cereal powdery mildew: Close or distant relatives?. Mol. Plant Pathol. 15, 304–314. https://doi.org/10.1111/mpp.12093 (2014).
11. Jørgensen, J. H. Erysiphe graminis, powdery mildew of cereals and grasses. Genet. Plant Pathogenic Fungi Adv. Plant Pathol. 6, 137–157 (1988).
12. Jankovic, T. et al. New insights into the life cycle of the wheat powdery mildew: direct observation of ascospore infection in Blumeria graminis f. sp. tritici. Phytopathology 105, 797–804. https://doi.org/10.1094/PHYTO-10-14-0268-R (2015).
13. Cowger, C., Parks, R. & Kosman, E. Structure and migration of U.S. Blumeria graminis f. sp. tritici populations. Phytopathology 106, 295–304. https://doi.org/10.1094/PHYTO-03-15-0066-R (2016).
14. Dreiseitl, A. Great pathotype diversity and reduced virulence complexity in a Central European population of Blumeria graminis f. sp. hordei in 2015–2017. Europ. J. Plant Pathol. 153, 801–811. https://doi.org/10.1007/s10658-018-1593-6 (2018).
15. Zhu, J. et al. Genetic evidence of local adaption and long distance migration in Blumeria graminis f. sp. hordei populations from China. J. General Plant Pathol. 82, 69–81. https://doi.org/10.1007/s10327-016-0643-1 (2016).
16. Limpert, E., Godet, F. & Müller, K. Dispersal of cereal mildews across Europe. Agric. Forest Meterol. 97, 293–308 (1999).
17. Wolfe, M. S., Brändle, U., Koller, B., Limpert, E. & McDermott, J. barley mildew in Europe: population biology and host resistance. Euphytica 63, 125–139 (1992).
18. Brown, J. K. & Hovmoller, M. S. Aerial dispersal of pathogens on the global and continental scales and its impact on plant disease. Science 297 (2002).
19. Kast, W. K. & Geiger, H. H. Studies on the inheritance of mildew resistance in rye. I. Results from inbred lines and F1 crosses. (1982).
20. Miedaner, T., Schmidt, H. & Geiger, H. Components of variation for quantitative adult-plant resistance to powdery mildew in winter rye. *Phytopathology* **83**, 1071–1075 (1993).

21. Nelson, R., Wiesner-Hanks, T., Wisser, R. & Balint-Kurti, P. Navigating complexity to breed disease-resistant crops. *Nat. Rev. Genet.* **19**, 21–33. https://doi.org/10.1038/nr.2017.82 (2018).

22. Friebe, B., Heun, M. & Bushuk, W. Cytological characterization, powdery mildew resistance and storage protein composition of tetraploid and hexaploid 1BL/1RS wheat-rye translocation lines. *Theor. Appl. Genet.* **78** (1989).

23. Hsam, S. L. K. & Zeller, F. J. Evidence of allelism between genes pm8 and pm17 and chromosomal location of powdery mildew and leaf rust resistance genes in the common wheat cultivar "Amigo." *Plant Breed.* **116**, 119–122 (1997).

24. Ren, T. H. et al. Development and characterization of a new 1BL.1RS translocation line with resistance to stripe rust and powdery mildew of wheat. *Euphytica* **169**, 207–213. https://doi.org/10.1007/s10681-009-9934-5 (2009).

25. Han, G. Characterization of the level of resistance to powdery mildew in striped rust. *Plant Dis.* **104**, 2940–2948. https://doi.org/10.1094/PDIS-02-20-0323-R (2020).

26. He, H. et al. Characterization of a new gene for resistance to wheat powdery mildew on chromosome 1RL of wild rye Secale sylvestre. *Theor. Appl. Genet.* **134**, 887–896. https://doi.org/10.1007/s00122-020-03739-1 (2021).

27. Koebner, R. M. D. Introduction into wheat of disease resistance from related species. *Annual report of the Plant Breeding Institute*, 69–70 (1986).

28. Driscoll, C. J. & Jensen, N. F. A genetic method for detecting introduced intergenetic translocations. *Theor. Appl. Genet.* **88**, 838–847 (2006).

29. An, D. G., Li, L. H., Li, J. M., Li, H. J. & Zhu, Y. G. Introgression of resistance to powdery mildew conferred by chromosome 2R and stripe rust. *Plant Dis.* **104**, 2948–2953. https://doi.org/10.1094/PDIS-02-20-0324-R (2020).

30. Hysing, S.-C. et al. A high-quality genome assembly highlights rye genomic characteristics and agronomically important genes. *Nat. Genet.* **53**, 574–584. https://doi.org/10.1038/s41588-021-00888-z (2021).

31. Vendelbo, N. M., Sarup, P., Orabi, J., Kristensen, P. S. & Jahoor, A. Genetic structure of a germplasm for hybrid breeding in rye (Secale cereale L.). *PLOS ONE* **15**, e0239541. https://doi.org/10.1371/journal.pone.0239541 (2020).
60. Vendelbo, N. M. et al. Genomic scan of male fertility restoration genes in a 'Gülzow' type hybrid 1 breeding system of rye (Secale cereale L.). Int. J. Mol. Sci., Submitted.

61. Wu, Y. et al. Molecular characterization of CIMMYT maize inbred lines with genotyping-by-sequencing SNPs. Theor. Appl. Genet. 129, 753–765. https://doi.org/10.1007/s00122-016-2664-8 (2016).

62. Baker, K. et al. The low-recombining pericentromeric region of barley restricts gene diversity and evolution but not gene expression. Plant J. 79, 981–992. https://doi.org/10.1111/tpj.12600 (2014).

63. Liu, H. J. & Yan, J. Crop genome-wide association study: A harvest of biological relevance. Plant J. 97, 8–18. https://doi.org/10.1111/tpj.14139 (2019).

64. Brachi, B., Moris, G. P. & Borevitz, J. O. Genome-wide association studies in plants: The missing heritability is in the field. Genome Biol. 12, 1–8 (2011).

65. Genes, G., Li, W., Chalilla, G. S., Zhu, H. & Wei, W. Recurrence of chromosome rearrangements and reuse of dna breakpoints in the evolution of the triticaceae genome. G3 Genetics 6, 3837–3847. https://doi.org/10.1534/g3.116.035089 (2016).

66. Martis, M. et al. Reticulate evolution of the rye genome. Plant Cell 25, 3685–3698. https://doi.org/10.1016/j.tpc.2014.05.003 (2013).

67. Huang, X. Q. & Röder, M. S. Molecular mapping of powdery mildew resistance genes in wheat: A review. Euphytica 137, 203–223 (2004).

68. Zhang, R. et al. Pm62, an adult-plant powdery mildew resistance gene introgressed from Dasypyrum villosum chromosome arm 2VL into wheat. Theor. Appl. Genet. 131, 2613–2620. https://doi.org/10.1007/s00122-018-1037-6 (2018).

69. Ma, R., Yli-Mattila, A. & Pulli, S. Phylogenetic relationships among genotypes of worldwide collection of spring and winter ryes (Secale cereale L.) determined by RAPD-PCR markers. Hereditas 140, 210–221 (2004).

70. Andersen, J. R. & Lubberstedt, T. Functional markers in plants. Trends Plant Sci. 8, 554–560. https://doi.org/10.1016/j.tplants.2003.09.010 (2003).

71. Brown, I. J., Jessop, A. C. & Rezanoor, N. H. Genetic uniformity in barley and its powdery mildew pathogen. Proc. Royal Soc. Lond. Ser. B Biol. Sci. 246, 83–90 (1991).

72. Xue, A. G. et al. Virulence structure of Blumeria graminis f sp tritici, the causal agent of wheat powdery mildew, in Ontario, Canada, in 2018 and 2019. Canad. J. Plant Pathol. 43, https://doi.org/10.1080/07060661.2021.1915876 (2021).

73. Liu, N. et al. Virulence structure of Blumeria graminis f sp tritici and its adaptive diversity by ISSR and SSR profiling analyses. PLoS ONE 10, e0130881. https://doi.org/10.1371/journal.pone.0130881 (2015).

74. Nieri, D., Di Donato, A. & Ercolano, M. R. Analysis of tomato meiotic recombination profile reveals preferential chromosome positions for NB-LRR genes. Euphytica https://doi.org/10.1007/s10681-017-1982-5 (2017).

75. Chen, N. W. G. et al. Common bean subtelomeres are hot spots of recombination and favor resistance gene evolution. Front. Plant Sci. 9, https://doi.org/10.3389/fpls.2018.01185 (2018).

76. Gruner, P., Schmitt, A. K., Fath, K., Pieplo, H. P. & Miederaner, T. Mapping and validating stem rust resistance genes directly in self-compatible genetic resources of winter rye. Theor. Appl. Genet. https://doi.org/10.1007/s00122-021-03800-7 (2021).

77. Wehling, P. et al. Leaf-rust resistance in rye (Secale cereale L.) 1 Genetic analysis and mapping of resistance genes Pr1 and Pr2. Theor. Appl. Genet. 107, 432–438. https://doi.org/10.1007/s00122-003-1263-7 (2003).

78. Van De Weyert, A. L. et al. A species-wide inventory of NLR genes and alleles in Arabidopsis thaliana. Cell 178, 1260–1272. https://doi.org/10.1016/j.cell.2019.07.039 (2019).

79. Steuernagel, B. et al. The NLR-annotator tool enables annotation of the intracellular immune receptor repertoire. Plant Physiol. 183, 468–482. https://doi.org/10.1002/pp.19.01273 (2020).

80. Mondragon-Palomino, M., Meyers, B. C., Michelmore, R. W. & Gaut, B. S. Patterns of positive selection in the complete NBS-LRR gene family of Arabidopsis thaliana. Genome Res. 12, 1305–1315. https://doi.org/10.1101/gr.159402 (2002).

81. Stone, R. D. et al. T-RPP13-3, a CC-NBS-LRR-like gene located on chr 7D, promotes disease resistance to wheat powdery mildew in Brock. J. Phytopathol. 168, 688–699. https://doi.org/10.1111/1439.12949 (2020).

82. Cheng, J. et al. Genome-wide Identification and expression analyses of RPP13-like genes in barley. BioChip J. 12, 102–113. https://doi.org/10.1007/s13206-017-2203-y (2018).

83. Li, Y. et al. High levels of nucleotide diversity and fast decline of linkage disequilibrium in rye (secale cereale L.) genes involved in frost response. BMC Plant Biol. 11, 6 (2011).

84. Gupta, P. K., Rustgi, S. & Kuhar, P. L. Linkage disequilibrium and association studies in higher plants: present status and future prospects. Plant Mol. Biol. 57, 461–485. https://doi.org/10.1007/s11103-005-0257-5 (2004).

85. Ren, Y. et al. QTL analysis and nested association mapping for adult plant resistance to powdery mildew in two bread wheat populations. Front. Plant Sci. 8. (2017).

86. Novakazi, F. et al. You had me at “MAGIC”!: Four barley MAGIC populations reveal novel resistance QTL for powdery mildew. Genes 11, 1512 (2020).

87. Arora, S. et al. Resistance gene cloning from a wild crop relative by sequence capture and association genetics. Nat. Biotechnol. 37, 139–143. https://doi.org/10.1038/s41587-018-0007-9 (2019).

88. USDA. United States Department of Agriculture: wheat and barley DNA extraction protocol (96-well plate format), <https://www.ars.usda.gov/ARSUserFiles/60701500/SmallGrainsGenotypingLaboratory/Protocols/wheat%20%20barleyDNA%20extracti on_original.pdf> (2006).

89. Pallotta, M. A. et al. Marker assisted wheat breeding in the southern region of Australia. Proceedings of the Tenth International Wheat Genetics Symposium, 789–791 (2003).

90. RStudio Team. Rstudio: integrated development environment for R. RStudio, Inc., Boston, <http://www.rstudio.com> (2015).

91. R Core Team. R a language and environment for statistical computing. R foundation for statistical computing, Vienna, Austria, <https://www.R-project.org/> (2021).

92. NCBI. National Center for Biotechnology Information, <https://www.ncbi.nlm.nih.gov> (2021).

93. Granato, I. S. C. et al. snpReady: A tool to assist breeders in genomic analysis. Mol. Breed. 38, 102. https://doi.org/10.1007/s11032-018-0844-8 (2018).

94. Afgan, E. et al. The Galaxy platform for accessible, reproducible and collaborative biomedical analysis. Nucleic Acids Res. 46, 537–544 (2018).

95. Rahse, G. & Hileman, S. Galactic cirrus: User-friendly cirrus plots within the galaxy platform. GigaScience 9, gga06560 (2020).

96. Hileman, S., Rahse, H. & Gallardi, C. Visualization with cirrus (galaxy training materials) <https://training.galaxyproject.org/training-material/topics/visualisation/cirrus/tutorial.html> (2021).

97. Wickham, H. ggplot2: R package (2017). <https://cran.r-project.org/web/packages/ggplot2/index.html> (2021).

98. Zhang, A. E. et al. GAPT: Genome association and prediction integrated tool. Bioinformatics 36, 2397–2399. https://doi.org/10.1093/bioinformatics/btx444 (2012).

99. Huang, M., Liu, X., Zhou, Y., Summers, R. M. & Zhang, Z. BLINK: A package for the next level of genome-wide association studies with both individuals and markers in the millions. Gigascience https://doi.org/10.1093/gigascience/giy154 (2019).
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Author contributions

All authors were involved in the study design. N.M.V. performed the phenotyping, bioinformatic analysis, and visual output and wrote the manuscript. K.M. assisted in the DNA extraction of the germplasm. P.S.K oversaw the multiplication of Bgs populations. J.O. was responsible for all communication with Trait Genetics and Eurofins Genomics conducting the SNP genotyping. K.M., P.S., J.O. and A.J. were involved in the intellectual input for the study, including the interpretation of the results. All authors were involved in the conceptualization of the study and revision of the manuscript.

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Competing interests

The authors declare no competing interests.

Additional information

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