Identification of Putative SNP Markers Associated with Resistance to Egyptian Loose Smut Race(s) in Spring Barley

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Abstract: Loose smut (LS) disease is a serious problem that affects barley yield. Breeding of resistant cultivars and identifying new genes controlling LS has received very little attention. Therefore, it is important to understand the genetic basis of LS control in order to genetically improve LS resistance. To address this challenge, a set of 57 highly diverse barley genotypes were inoculated with Egyptian loose smut race(s) and the infected seeds/plants were evaluated in two growing seasons. Loose smut resistance (%) was scored on each genotype. High genetic variation was found among all tested genotypes indicating considerable differences in LS resistance that can be used for breeding. The broad-sense heritability (H2) of LS (0.95) was found. Moreover, genotyping-by-sequencing (GBS) was performed on all genotypes and generated in 16,966 SNP markers which were used for genetic association analysis using single-marker analysis. The analysis identified 27 significant SNPs distributed across all seven chromosomes that were associated with LS resistance. One SNP (S6_17854595) was located within the HORVU6Hr1G010050 gene model that encodes a protein kinase domain-containing protein (similar to the Un8 LS resistance gene, which contains two kinase domains). A TaqMan marker (0751D06 F6/R6) for the Un8 gene was tested in the diverse collection. The results indicated that none of the Egyptian genotypes had the Un8 gene. The result of this study provided new information on the genetic control of LS resistance. Moreover, good resistance genotypes were identified and can be used for breeding cultivars with improved resistance to Egyptian LS.

Keywords: Hordeum vulgare L.; Ustilago nuda; genetic analysis; single-nucleotide polymorphism; candidate genes

1. Introduction

Barley (Hordeum vulgare L.) is one of the most important cereal grains grown in temperate climates. It has been used for animal feed (as a grain or as fodder) and human nutrition (as a component of various foods). During its life cycle, barley suffers from many biotic and abiotic stresses which affect its yield. Loose smut (LS) (caused by the basidiomycete pathogen Ustilago nuda (Jens.) Rostr. (U. nuda)) is one of these important diseases. This disease has been reported in 1.3–11.7% of fields in Egypt [1], 50–70% of fields in Western Canada [2], and it is also common in the Great Plains of the United States [3].
Certified barley seed is highly recommended to be treated with fungicide to prevent this disease. During grain development, the smutted (diseased) heads disperse spores by wind or rain to healthy heads. After harvest, the infected seeds with LS appear normal. However, in the next season, after germination, the fungus will infect and grow within the plant and replace the head with spores at the flowering time [4]. A reduction in yield is directly associated with the percentage of infected tillers. Climate change is expected to increase the pathogen spread by increasing temperatures, changing precipitation patterns, changing extreme weather events, and reducing water availability [5]. Therefore, the chances of agriculture crops encountering both biotic and/or abiotic stresses are expected to be more frequent than in the past. This change will result in a significant deterioration in staple crops such as barley and wheat (*Triticum aestivum* L.).

Loose smut is a particular problem for organic and low-input agriculture as fungicidal seed treatments are not applied. Therefore, breeding resistant barley cultivars for LS is the main approach to reducing the devastating effects of LS in these cropping systems. It is equally important that the new cultivars combine both good agronomic performance and LS resistance. [6,7] reported *Un* as the first resistance gene in the cv. ‘Trebi’ in the 1940s. Thereafter, 15 resistance genes were found to be associated with true loose smut resistance [8–12]. Of the 15 genes, *Un8* showed durable resistance against all true loose smut isolates that were observed in Western Canada [13].

Traditional plant breeding programs have been reported as a very successful tool for developing cultivars showing resistance to diseases. However, plant breeding is a time-consuming and labor-expensive or intensive process as it largely depends on field evaluation and selection [14]. Molecular marker-assisted selection (MAS) for true loose smut resistance is an alternative approach and can be useful in accelerating breeding for resistance, as well as improving agronomic performance. Significant progress has been made in identifying molecular markers that show a strong linkage with loose smut-resistant genes like *Un8* [15]. SSR and RFLP markers were used to initially tag *Un8* [3,16]. However, when single nucleotide polymorphism (SNP) maps became available for barley, it was feasible to develop additional markers for *Un8*. Recently, Zang [17] identified two flanking SNP markers located very close to *Un8*. Since SNP markers provide valuable and more reliable tagging than other types of DNA markers, SNP markers close to *Un8* can be used for breeding resistance to true loose smut. However, SNP markers associated with the other 15 barley loose smut resistance genes are lacking. To date, there is no information on DNA markers associated with resistance against 98 Egyptian loose smut races.

More information is needed to understand the genetic control of LS resistance in barley globally. In our study, barley resistance to LS under Egyptian conditions was investigated.

The objectives of this study were to (1) examine the genetic variation among Egyptian and spring barley genotypes in their resistance to Egyptian LS races, (2) identify putative SNP markers and genomic regions associated with LS resistance to be used in marker-assisted selection for loose smut resistance in barley, and (3) select the most resistant genotypes which could be used in breeding programs to genetically improve LS resistance for Egyptian barley production.

2. Materials and Methods

2.1. Plant Materials

A total of 62 highly diverse spring barley genotypes were included in this study. The seeds of 60 genotypes were obtained from the Agriculture Research Center (ARC), Egypt, and two from the United States Department of Agriculture (USDA), USA. The complete list of the genotypes and the pedigree is represented in Supplementary Table S1. The tested genotypes included 2-rows, 6-rows, and naked and hulled barley genotypes. The genotypes were divided based on their population structure into two subpopulations based on their population structure, genetic diversity, and the possibility of using them to improve different traits relevant to Egyptian growing conditions [18].
2.2. Evaluation of LS Resistance

The genotypes were inoculated with the Egyptian race(s) of LS in the greenhouse of the Plant Pathology Department, Assiut University, Egypt, in 2017/2018. An inoculation liquid was prepared by mixing 1 gm of *U. nuda* spores collected from the Egyptian fields in the previous years with 1 L of distilled water [19]. For each tested genotype, 6–8 heads were inoculated using a 3 mL syringe. After harvest, inoculated spikes from each genotype were collected. The LS resistance was evaluated in the 2018/2019 and 2019/2020 growing seasons at the heading stage, according to Eckstein et al. [16]. The infected seeds were planted in the greenhouse in a randomized complete block design (RCBD) with three replications. The infected and health spikes for each genotype were counted and the resistance was calculated as a percentage of the infected spike divided by the total number of spikes. The level of resistance was determined based on [20] using a scale as follows: percentage of infected heads 0% = immune genotype, 0.1–5% (highly resistant) genotype, 5.1–10% (resistant), 10.1–20% (moderately resistant), 20.1–30% (moderately susceptible), 30.1–50% (susceptible), and 50.1–100% (highly susceptible).

2.3. Statistical Analysis of LS Resistance

An analysis of variance (ANOVA) of LS resistance was conducted using R software [21]. Data from both years were combined and the following model was used:

\[ Y_{ijk} = \mu + g_i + r_j + y_k + gy_{ik} + e_{ijk}, \]

where \( Y_{ijk} \) is the disease reaction of genotype \( i \) in replication \( j \) planted in year \( k \), \( \mu \) is the general mean; \( g_i \), \( r_j \), and \( y_k \) are the main effects of genotypes (fixed effects), replications, and years (random effects), respectively. \( gy_{ik} \) is genotype \( \times \) year interaction and; \( e_{ijk} \) is the error. The broad-sense heritability \((H^2)\) was calculated as follows:

\[ H^2 = \frac{\sigma^2_G}{\sigma^2_G + \frac{\sigma^2_R}{ry}} \]

where \( \sigma^2_G \) and \( \sigma^2_R \) are the variance of the lines and the residuals, \( r \) is the number of replicates within the experiment, and \( y \) is the number of years.

2.4. DNA Extraction and Genotyping-by-Sequencing (GBS)

DNA was extracted from all the 60 genotypes by collecting 4–5 leaves from five-day-old barley seedlings. The extraction protocol was conducted as described in Mourad et al. [22]. DNA concentration was measured using spectrophotometry (Gen5TM microplate reader and image software with Take3TM micro-volume plates (BioTek, Winooski, VT, USA). DNA of each genotype was digested using two restriction enzymes, *PstI* and *MspI*, for genotyping-by-sequencing as described in [23]. Single nucleotide polymorphism (SNP) calling used the TASSEL 5.0 v2 GBS pipeline [24]. Identification of SNP markers, their physical position, and localization was carried out using the Barley cv. MorexV2 genome assembly. A set of 25,700 SNPs was obtained from the GBS data that was filtered for minor allele frequency (MAF >0.05), maximum missing sites per SNP <20%, and maximum missing sites per genotype <20%. Heterozygous loci were then marked as missing to obtain better estimates of marker effects (Peter Bradbury, personal communication) and the filtration was repeated based on the previous criteria.

2.5. Single Marker Analysis of LS Resistance and Linkage Disequilibrium

The phenotypic data (loose smut resistance, %), as well as the available SNP markers, were used to perform a single-marker-analysis (SMA) to identify SNP markers significantly
associated with resistance. Single marker analysis was performed using PowerMarker software V 3.25 [25] using the following model:

\[ Y = \mu + f(\text{marker}) + \text{error}, \]

where \( Y \) is equal to the trait value, \( \mu \) is equal to the population mean, and \( f(\text{marker}) \) is a function of the significant markers. The phenotypic variation explained by each significant SNP marker was estimated using TASSEL 5.0 software [24].

For significant SNP markers located on the same chromosome, linkage disequilibrium (LD, \( r^2 \)) was calculated using TASSEL 5.0 and visualized as a heatmap using “Ldheatmap” R package [26]. The genetic distance among immune genotypes was calculated based on their resistance alleles using a simple matching approach. The genetic distance calculation was performed using the ade4 R package [27].

2.6. Gene Models Underlying Significant SNPs and Their Validation

To further confirm the SMA results, we investigated if any of the significant SNPs were located within gene models identified in the reference genome assembly published by the International Barley Genome Consortium (IBSC V.2) available on Ensemblplants (http://plants.ensembl.org/Hordeum_vulgare/Info/Index, accessed on 12 June 2022). The functional annotation of the identified gene models was retrieved from the genome annotation provided by IBSC_V2 and examined for their association with disease resistance. The expression of candidate genes was investigated from the Expression Atlas database (https://www.ebi.ac.uk/gxa/home, accessed on 12 June 2022).

2.7. Screening of the Genotypes for the Presence of the Un8 LS Resistance Gene

The 0751D06 F6/R6 TaqMan® assay, located within the Un8 gene, was developed by Zang et al. [17] to screen for the presence of Un8 resistance or susceptible alleles. The 60 barley genotypes were genotyped using this marker at the Department of Plant Sciences, the University of Saskatchewan. TaqMan® SNP genotyping was performed with the ABI StepOnePlus™ Real-Time [17]. Five genotypes were used as checks for the Un8 gene. TR12135 and AC Metcalfe were characterized as resistant to loose smut and carried the Un8 gene, while TR14150, Harrington and Xena were characterized as susceptible and lack the Un8 gene [17]. The seeds of checks were obtained from the Department of Plant Sciences, University of Saskatchewan, Saskatoon, SK, Canada.

3. Results

3.1. Phenotypic Variation in Loose Smut Resistance in the Tested Genotypes

The analysis of variance for loose smut resistance is presented in Table 1. The ANOVA revealed highly significant differences among the genotypes for loose smut resistance. No significant differences were found within the main effect of the year or within the genotypes × replications and genotypes × years interaction effects. Broad-sense heritability was high based on the average of the two years (\( H^2_B = 0.95 \)).

Table 1. Analysis of variance for loose smut resistance in the 60 genotypes comprising the Egyptian spring barley collection.

| Source         | d.f | M.S.  |
|----------------|-----|-------|
| Years (Y)      | 1   | 13.79 |
| Replications (R)| 2   | 216.30* |
| Genotypes (G)  | 56  | 1231.26** |
| GR             | 112 | 51.91 |
| GY             | 56  | 58.33 |
| GYR            | 110 | 46.53 |

Heritability 0.95

* \( p < 0.05 \), ** \( p < 0.01 \).
The 57 genotypes displayed different degrees of infected heads ranging from 0 to 64% based on the average of both years. All genotypes could be classified into six groups: immune (20 genotypes), highly resistant (12 genotypes), resistant (seven genotypes), moderately resistant (nine genotypes), moderately susceptible (four genotypes), susceptible (three genotypes), and highly susceptible (two genotypes) (Figure 1a). A set of 22 genotypes were immune in the 2019 experiment. Whereas in the 2020 trial, this number increased to 37 genotypes with 0% infected heads (Figure 1c). Twenty genotypes had no infected spikes in the two years. A positive significant correlation was found between the two years ($r = 0.78 \text{ **}$) for LS resistance (Figure 1c). The highly resistant genotypes showed only healthy spikes (Figure 2a), the susceptible genotypes had fully infected spikes (Figure 2b), while the moderately resistant genotypes showed partially infected spikes (Figure 2c,d).

**Figure 1.** (a) Distribution of loose smut reaction among the 60 genotypes comprising the Egyptian spring barley collection, (b) Venn diagram illustrating the identification of immune genotypes in 2019 and 2020, (c) Correlation of loose smut reactions measured in 2019 and 2020 among the genotypes demonstrating an immune reaction in at least one year. **$p < 0.01$.**

**Figure 2.** The effect of the Egyptian loose smut race on barley spikes (a) health spikes (b) susceptible spikes, (c,d) different degrees from few partially infected spikes found in some resistant genotypes.
3.2. Single-Marker Analysis of Loose Smut Resistance against the Egyptian Race and Linkage Disequilibrium between the Significant SNPs

Genotyping-by-sequencing generated a set of 25,700 SNPs. After filtration, the number decreased to 16,966 SNPs used for SMA and further genetic analysis. Due to the absence of the genotypes x year interaction, average LS values obtained from both years were used in the SMA analysis. The SMA identified 27 SNP markers significantly associated with the resistance ($p$-value < 0.001) (Table 2, Figure 3a). These significant SNPs were distributed among the seven chromosomes of the barley genome, with the highest number on chromosome 3H (13 SNPs) and the lowest number on chromosomes 5H and 6H (one SNP each). The phenotypic variation of the significant SNPs ($R^2$) ranged from 0.55% for SNP S3_4158424 (3H) to 47.70% for SNP S2_6903688 (2H). The four significant SNPs identified on chromosome 2H were found to have the highest $R^2$, with values ranging from 16.12% to 47.70%. The target alleles of the significant SNPs decrease the symptoms of loose smut in the genotypes with a percentage ranging from 1.71% for SNP S3_4158424 to 23.57% for SNPs S3_546788379, S3_546788391, and S3_546788395.

Table 2. Single marker analysis (SMA) for loose smut resistance in the 60 barely genotypes comprising the Egyptian spring barley collection.

| SNP ID   | Chrom. | p-Value         | F-Value | $R^2$   | Target Allele | Allele Effect |
|----------|--------|-----------------|---------|---------|---------------|---------------|
| S1_523594 | 1H     | 0.000652344     | 13.86   | 27.256  | G/A           | -15.54%       |
| S2_6903688 | 2H     | $3.37 \times 10^{-5}$ | 24.62   | 47.70   | C/G           | -14.57%       |
| S2_6903698 | 2H     | 0.000697591     | 14.65   | 35.17   | G/C           | -15.57%       |
| S2_71163470| 2H     | 0.000146541     | 18.71   | 37.643  | C/T           | -21.92%       |
| S2_481116363| 2H     | 0.000366765     | 16.12   | 34.949  | A/G           | -13.15%       |
| S3_4158424 | 3H     | 0.000906049     | 8.28258173 | 0.545 | C/T           | -1.71%        |
| S3_4158440 | 3H     | 0.00081791      | 8.42495764 | 1.085 | C/T           | -3.23%        |
| S3_51209627| 3H     | 0.000847857     | 8.37494083 | 17.162 | T/C           | -9.52%        |
| S3_346822835| 3H     | 0.00072923      | 8.85119232 | 24.637 | C/T           | -17.09%       |
| S3_425653844| 3H     | 0.000956134     | 8.20841245 | 19.239 | T/C           | -9.89%        |
| S3_543138543| 3H     | 0.000282644     | 9.941050754 | 14.42 | G/A           | -8.31%        |
| S3_546418133| 3H     | 0.000758585     | 8.529945709 | 27.916 | T/C           | -17.86%       |
| S3_546728405| 3H     | $1.85 \times 10^{-5}$ | 14.18761801 | 26.1  | G/A           | -19.87%       |
| S3_546728420| 3H     | 0.000332253     | 9.705460245 | 27.021 | A/G           | -13.42%       |
| S3_546788379| 3H     | 0.000127966     | 11.12149677 | 33.878 | T/A           | -23.57%       |
| S3_546788391| 3H     | 0.000127966     | 11.12149677 | 33.878 | A/G           | -23.57%       |
| S3_546788395| 3H     | 0.000127966     | 11.12149677 | 33.878 | C/A           | -23.57%       |
| S3_557269134| 3H     | 0.000761867     | 8.523916211 | 24.861 | T/A           | -10.69%       |
| S4_510236504| 4H     | 0.000617005     | 8.819871045 | 16.644 | A/G           | -8.94%        |
| S4_510236509| 4H     | 0.000617005     | 8.819871045 | 16.644 | A/G           | -8.94%        |
| S5_8264909 | 5H     | 0.000755518     | 8.53560527 | 22.878 | A/G           | -10.61%       |
| S6_17854959 | 6H     | 0.000986781     | 8.164849355 | 5.432  | A/G           | -7.81%        |
| S7_10286228 | 7H     | 0.000586585     | 8.960296029 | 7.383  | G/A           | -8.22%        |
| S7_27629275 | 7H     | 0.000429506     | 9.335043364 | 0.776  | T/C           | -2.17%        |
| S7_50465300 | 7H     | 0.000980333     | 8.173896443 | 3.264  | A/G           | -3.92%        |
| S7_50465325 | 7H     | 0.000980333     | 8.173896443 | 3.264  | A/G           | -3.92%        |
| S7_54495415 | 7H     | 0.000855639     | 8.36225696 | 7.226  | C/G           | -7.61%        |
Figure 3. (a) Manhattan plot showing the significant SNPs (red circles) associated with LS resistance and their locations on barley chromosomes. The red line indicated the significant threshold at 0.001, (b) LD ($r^2$) among significant SNPs located on chr 3H.

LD ($r^2$) between each pair of significant SNPs located on the same chromosome was calculated. The highest number of high LD genomic regions was found on chromosome 3H among the 13 significant SNPs (Figure 3b). Complete LD was found among S3_546788395, S3_546788391, and S3_546788379. The entire LD block was divided by SNP S3_546728420 due to missing data associated with this SNP. In addition, high LD was found between S3_546728420 and S3_51209627, and between S3_546788379 and S3_543138543. Two SNPs located on chromosome 4H were incomplete LD (Supplementary Figure S1c). No significant LD was found between the significant SNPs located on chromosome 2H nor 7H (Supplementary Figure S1a,b).

3.3. Genes Underlying Significant SNPs and Their Validation

To further understand the genetic association between the significant SNP markers and loose smut resistance in the tested genotypes, gene models harboring these SNPs and their functional annotation were investigated (Table 3). Gene models that were located within or near significant SNPs were determined. Eleven of the SNPs were not located near or within any gene models. The HORVU3Hr1G002060 gene model was found to be 455 and 439 bp from two SNPs, S3_4158424 and S3_4158440, respectively, located on chromosome 3H. This gene model encodes a protein belonging to the O-acyltransferase (WSD1-like) family. The S6_17854595 SNP located on chromosome 6H was found to be located within a gene model HORVU6Hr1G010050 which encodes a protein kinase domain-containing protein (Figure 4).
Table 3. The genomic location and annotation of gene models harboring SNP markers significantly associated with loose smut resistance in the 60 Egyptian genotypes.

| SNP ID      | Chrom. | Gene Model                  | Gene Model Position (bp) | Pos. (Gene Model) | GD (Gene Model) | Gene Annotation                                      |
|-------------|--------|-----------------------------|--------------------------|------------------|----------------|-----------------------------------------------------|
| S1_523594   | 1H     | HORVU1Hr1G000190            | 514755-520291            | after            | 3033           | Fatty acyl-CoA reductase                             |
| S2_6903688  | 2H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S2_6903698  | NA     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S2_71163470 | NA     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S2_48111636 | NA     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S3_4158424  | 3H     | HORVU3Hr1G002060            | 415879-4162702           | before           | 455            | O-acyltransferase (WSD1-like) family                |
| S3_4158440  | 3H     | HORVU3Hr1G002060            | 415879-4162702           | before           | 439            | O-acyltransferase (WSD1-like) family protein        |
| S3_51209627 | 3H     | HORVU3Hr1G019140            | 51126683-51134627        | after            | 75000          | receptor kinase 1                                   |
| S3_346828235| 3H     | HORVU3Hr1G049410            | 34682885-346863988       | before           | 34650          | Disease resistance protein RPM1                     |
| S3_42565344 | 3H     | HORVU3Hr1G056930            | 425800602-425801141      | before           | 146798         | Protein H                                           |
| S3_54313854 | 3H     | HORVU3Hr1G071870            | 543136922-543138097      | NA               | NA             | Uncharacterized protein                             |
| S3_54641813 | 3H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S3_54672840 | 3H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S3_54672842 | 3H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S3_54678879 | 3H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S3_54678891 | 3H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S3_54678895 | 3H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S3_55726913 | 3H     | HORVU3Hr1G074130            | 557216044-557217268      | after            | 51866          | Holliday junction DNA helicase                      |
| S4_51023650 | 4H     | HORVU4Hr1G060790            | 510209263-510210332      | after            | 26172          | unknown function                                   |
| S4_51023659 | 4H     | HORVU4Hr1G013970            | 510209263-510210332      | after            | 27246          | unknown function                                   |
| S5_8264909  | 5H     | HORVU1Hr1G010050            | 17851141-17927010        | within           | -              | Protein kinase domain-containing protein            |
| S6_17854595 | 6H     | HORVU1Hr1G007880            | 10273667-10273950        | after            | 12278          | HAT family dimerisation domain containing protein, expressed |
| S7_10268228 | 7H     | HORVU1Hr1G020360            | 27617705-27618357        | after            | 9368           | Undescribed protein                                 |
| S7_27627925 | 7H     | HORVU1Hr1G028130            | 5050827-50501771         | before           | 35527          | NA                                                  |
| S7_54453325 | 7H     | NA                          | NA                       | NA               | NA             | NA                                                  |
| S7_54459415 | 7H     | HORVU1Hr1G029150            | 54451633-54453868        | after            | 41547          | Ubiquitin-like domain-containing protein            |
3.4. Genotyping with Un-8 Taqman Marker

The presence or absence of the Un8 gene was tested using the 0751D06 F6/R6 TaqMan® assay. To confirm the genotyping results, two positive checks and three negative checks for the Un8 gene were used along with the 62 genotypes. The results revealed that none of the genotypes used in this study possessed the Un8 gene (Supplementary Table S1).

3.5. Selection of Superior Loose Smut Resistance Genotypes

Out of the 20 immune genotypes (Supplementary Table S1), only 16 genotypes had available sequence data after filtration criteria. To genetically confirm the superior resistance in these 16 highly resistant genotypes, the number of targeted alleles (favorable alleles) of the 27-significant SNP markers was investigated in each resistant genotype (Figure 5a, Supplementary Table S2). This number ranged from 4 to 19 SNP markers. Two genotypes, SC2-19 and SCYT-33 contained the highest number of targeted SNPs. The PNBYT7 cultivar was found to be carrying the lowest number of targeted SNP alleles (four alleles). Furthermore, the presence of the targeted alleles of the two SNP markers with the highest allele effect (S2_6903688 and S2_71163470) was investigated in the genome of these resistant genotypes. Most of the resistant genotypes (12 genotypes) were found to contain the targeted allele of S2_71163470 SNP. However, only five of these genotypes carried the targeted allele of S2_6903688 SNP.
Figure 5. (a) number of resistant alleles in the most resistant genotypes and (b) genetic distance and dendrogram analyses among the most resistant genotypes based on the presence of resistance alleles.

To further understand the possible improvement of loose smut resistance in spring barley using these genotypes, the genetic distance between each pair of resistant genotypes was calculated based on the significant SNPs (Supplementary Table S3) and presented as a dendrogram cluster (Figure 5b). Based on the dendrogram cluster, the two genotypes with the highest number of targeted SNPs, “SCYT-33 and SC2-19”, were located in the same cluster. These two genotypes differed in six alleles. The PNYTI7 genotype was located in a separate cluster from the remaining 15 genotypes.

4. Discussion

As loose smut is one of the major diseases that affect the production of barley globally, it is very important to understand the genetic basis of resistance. No previous studies related to resistance against Egyptian LS race(s) has been conducted. Up to our knowledge, our study could be considered as the first report on the genetics of LS resistance against Egyptian LS. Understanding such an important point will accelerate the production of barley, not only in Egypt but also in the different parts of the world where LS disease occurs, and organic agriculture is practiced. In the current study, the presence of highly susceptible genotypes (with the percentage of infected heads >50%) in both years indicated the successfulness of the artificial infection and the validity of the results.

4.1. Genetic Variation in Loose Smut Resistance

The presence of highly significant differences among the tested genotypes in their response to loose smut indicated our artificial inoculation and phenotypic assay were successful. The resistant genotypes would be very useful in improving resistance to this disease when grown in Egyptian environments. The high correlation in disease reaction response of the tested genotypes and the absence of G × Y interaction to LS in the two years indicated that consistent response to LS. The high degree of broad-sense heritability ($H^2_B = 0.95$) also indicated that the phenotypic variation of LS resistance in the studied materials is stable and mainly due to genotypic variation. As a result, improving LS resistance in spring barley could be possible due to the successful selection of highly resistant genotypes, which will be very useful, especially for low input and organic fields where seed treatment with fungicides is not practiced. The different degrees of the partially infected spikes (Figure 1a) may be indicated by the presence of different loose smut races, as those genotypes presented different infection types on their spikes. In the partially infected spikes, the basal parts of the infected spike are smutted, with the distal parts producing normal grains [28].

Twenty of the evaluated genotypes were found to be immune to LS in both years. Most of these genotypes were 6-rows (12 genotypes) or naked (14 genotypes), with six of them having both traits (naked 6-row type). It was known that 6-row barley genotypes are highly
fertile and high-yielding due to the formation of kernels from all the spikelet’s flowers. Naked barley genotypes were also preferred as they are linked directly to dietary food uses [29]. The presence of LS resistance in naked 6-row barley will allow the improvement of this important barley type for Egyptian barley production. Diamond from Canada and Wocus 71 (CIho 15554) from the USA had the highest number of infected spikes and were considered highly susceptible to LS. Therefore, these two genotypes can be used as susceptible checks against the Egyptian race of LS in future loose smut experiments in barley to validate and calibrate the successful inoculation with the fungus.

4.2. Genetic Analysis of LS Resistance

To fully utilize the resistant genotypes identified in this study, it is very important to understand the genetic control of LS. In this study, the genotyping was undertaken using 16,966 SNP markers which were highly polymorphic and distributed across all genotypes. The genetic association can be carried out using different kinds of analysis, such as genome-wide association study (GWAS) and single-marker analysis (SMA). In this study, the 60 genotypes used were not appropriate for GWAS studies, which require at least 100 genotypes to be precise [30]. However, SMA can be carried out using any number of genotypes, hence, it would be appropriate for this study [31,32]. Using the same set of genotypes and SNP, the single-marker analysis was successful to identify genomic regions associated with other target traits in barley, such as drought tolerance [33] and heat tolerance [34]. Single-marker analysis identified a set of 27 SNP markers associated with LS resistance in the studied genotype materials based on the average of both years data. The chromosomal location of the identified SNPs indicated the presence of multiple genomic regions controlling resistance (Figure 3). Based on the $R^2$ of the significant SNPs, all SNPs with $R^2$ less than 10% were considered as marking minor effects (chromosomes 6H and 7H). The SNPs located on chromosomes 1H, 2H, 4H, and 5H could be considered as marking major effect loci ($R^2 > 10%$) [35–37]. Two regions on chromosome 3H, as determined by low LD between two distinct groups of SNPs, were associated with LS resistance, with the majority considered as marking a major QTL. The different degrees of LD between each pair of significant SNPs on the same chromosome indicated the presence of multiple QTLs/genes. High LD SNPs were found on the 3H chromosome and indicated that these genomic regions tend to co-inherited together. As a result, we can conclude the presence of multiple genes/QTLs controlling the resistance against the Egyptian race of LS. The resistance alleles identified by these SNPs were associated with a decrease in loose smut symptoms from 1.7 to 23.9%. Chromosome 3H had a high LD genomic region of three SNPs (S3_546788379, S3_546788391, and S3_546788395) that decreased loose smut resistance by 23.92%. Therefore, these markers could be useful for breeding loose smut resistance in barley.

To further understand the genetic association between the identified significant SNPs and LS resistance, gene models harboring or located near significant SNPs were investigated. The annotation revealed 15 gene models. Out of the 15 gene models, three were identified to be located very near the significant SNPs on chromosome 1H (one gene model) and 3H (two gene models). The gene model identified on chromosome 1H was functionally annotated as fatty acyl-CoA reductase (FAR), an enzyme that is responsible for alcohol and fatty acid formation. Fatty acids were reported to be a very important regulator of plant defense against different microbes [38,39], while the functional annotation of the two gene models on chromosome 3H encodes O-acyltransferase (WSD1-like) family. The HORVU3Hr1G002060 gene model was located near S3_4158424 and S3_4158440. These two SNPs were located before the gene model at a distance of 439 and 455 bp, respectively. O-acyltransferase (WSD1-like) family plays an important role in synthesizing different types of waxes [40]. These wax components are transported from the endoplasmic reticulum to the plasma membrane via the Golgi and consequently transported out of the plant cell to the cuticle layer. The plant cuticle consists of a cutin scaffold covered with cuticular waxes [41,42] that were reported to play a vital role in plant defense against pathogens by acting as a plant’s physical barrier, regulating the plant–pathogen interactions [9,43].
Of all significant SNPs identified, one SNP on chromosome 6H is located within a gene model of HORVU3Hr1G002060. This gene encodes a protein kinase domain-containing protein. Protein kinase domains can be divided into two phosphorylating groups, phosphorylating either serine/threonine or tyrosine residues on target proteins [44]. Serine/threonine protein kinases have been associated with plant disease resistance pathways [41]. Such proteins have previously been associated with resistance to powdery mildew (Blumeria graminis f. sp. tritici) in wheat [41], stem rust resistance in barley [42], bacterial blight disease resistance in rice [12], and stripe rust resistance in wheat [43]. HORVU6Hr1G010050 was found to be highly expressed 1.8-fold under stem rust resistance compared to the control (Figure 4, Supplementary Table S4). Moreover, it also had a higher expression under powdery mildew than under control. Therefore, the S6_1785495 located on the 6H chromosome could be very useful to target the HORVU6Hr1G010050 gene. The result of gene annotation of this SNP supports the power of genetic association performed in this study in detecting putative SNP associated with loose smut resistance. A KASP marker targeting this SNP can be designed and used for validation in other genetic backgrounds before using it for marker-assisted selection to improve resistance to LS in barley. The expression of the other candidate gene models did not associate with disease resistance.

As mentioned previously, among the 15 different LS resistance genes reported, Un1 to Un15, only the Un8 gene has a defined chromosomal location [17]. Due to the lack of enough available information about the LS resistance genes in barley, it is very difficult to determine if the known resistance genes might exist in this current studied material. Hence, more studies should be conducted to provide a deep understanding of the genetic control of LS resistance under Egyptian conditions.

4.3. Genotyping of Un8 LS Resistant Gene

The Un8 resistant gene was reported as an important gene that is very effective against all LS races in Europe and Canada [17,45]. There is no available information about the effectiveness of this gene against Egyptian LS races, therefore, it was important to test the effectiveness of this gene against Egyptian races. The 0751D06 F6/R6 Un8-specific marker (A/G) was used to evaluate the 60 Egyptian genotypes, along with two resistant checks (AC Metcalfe and TR12135). All genotypes were found to be non-carriers of the Un8 gene. The immune genotypes thus appear to carry other genes for LS resistance. Further trials are needed to evaluate the effectiveness of this gene against the Egyptian race, as integrating this gene into the Egyptian barley gene pool could improve resistance to Egyptian LS races.

4.4. Selection of Superior Genotypes to LS Resistance

According to the phenotypic and genotypic criteria used in our selection, the best genotypes that can be used for crossing would be SCYT-33 and PNBYT15 for the following reasons; (I) the genetic distance between these two resistant genotypes was 0.88, (II) they were found to have 19 and 6 favorable marker alleles, respectively, and (III) a set of 19 different favorable alleles were found between them and shared three favorable alleles. Therefore, cultivars having 22 favorable alleles may be produced from crossing between these two genotypes. Although SCYT-33 and SC2-19 had the highest number of target alleles, they had the lowest genetic distance (0.43). The number of different alleles between the two genotypes was 9 and they shared 13 favorable alleles and were located in the same subpopulation [18]. Therefore, crossing these two genotypes could not be as effective as crossing between SCYT-33 and PNBYT15. Crossing between two far genetic distance genotypes with as many as target alleles could produce lines with distinct alleles controlling the trait and high combining ability [46]. An additional concern that should be taken into account when using SCYT-33 and SC2-19 as parents in breeding for LS resistance is that both genotypes have hulled grain heads which are not preferred in the food industry. In addition, the lowest number of target alleles markers were found in PNBYT17 (four alleles’ markers). This genotype was found to have a high genetic distance from SCYT-33 and SC2-19, indicating that crossing between these two genotypes and PNBYT17 could improve
5. Conclusions

The results of this study provide valuable information on the genetic improvement of loose smut resistance in barley, especially in the context of Egyptian barley production. A set of 27 SNPs were reported to be associated with loose smut resistance. Of the 27 SNPs, one SNP located on chromosome 6H could be a promising marker as it is within a gene model encoding a protein kinase previously associated with resistance to other plant pathogens. Moreover, the 0751D06 F6/R6 marker targeting the Un8 resistance genes demonstrated that this gene was not present in the Egyptian genotypes. Promising resistant genotypes were identified that could be used for future breeding programs to improve loose smut resistance in barley.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/genes13061075/s1, Table S1: Detailed information for the 60 genotypes comprising the Egyptian spring barley collection; Table S2: Presence of favorable alleles in the 16 immune genotypes; Table S3. Genetic distance calculated among the 16 immune genotypes with available sequence data; Table S4. Expression analysis of HORVU6Hr1G010050 gene model under different fungal diseases; Figure S1. Linkage disequilibrium between each pair of significant SNPs located on chromosomes 2H (a), 7H (b), and 4H (c).

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