**A Large Bioassay Identifies *Stb* Resistance Genes that Provide Broad Resistance against Septoria Tritici Blotch Disease in the UK**

**Henry Tidd1, Jason Rudd1, Rumiana Ray2, Ruth Bryant3, Kostya Kanyuka4**

**1: Protecting Crops and the Environment, Rothamsted Research, Harpenden, AL5 2JQ, UK; 2Division of Plant and Crop Sciences, School of Biosciences, University of Nottingham, Sutton Bonington, Loughborough, LE12 5RD, UK; 3RAGT Seeds, Ickleton, CB10 1TA, UK; 4NIAB, 93 Lawrence Weaver Road, Cambridge, CB3 0LE, UK**

**Correspondence:**

**Henry Tidd (**[**henry.tidd@rothamsted.ac.uk**](mailto:henry.tidd@rothamsted.ac.uk)**)**

**Kostya Kanyuka (**[**kostya.kanyuka@niab.com**](mailto:kostya.kanyuka@niab.com)**)**

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**Abstract**

Septoria tritici blotch (STB) is one of the most damaging fungal diseases of wheat in Europe, largely due to the paucity of effective resistance genes against it in breeding materials. Currently dominant protection methods against this disease, e.g. fungicides and the disease resistance genes already deployed, are losing their effectiveness. Therefore, it is vital that other available disease resistance sources are identified, understood and deployed in a manner that maximises their effectiveness and durability. In this study, we assessed wheat genotypes containing nineteen known major STB resistance genes (*Stb1* through to *Stb19*) or combinations thereof against a broad panel of 93 UK *Zymoseptoria tritici* isolates. Five infection symptom components (days post infection to the development of first symptoms and pycnidia, percentage coverage of the infected leaf area with chlorosis/necrosis and pycnidia and spore counts from spore wash) were measured and average disease levels calculated for each genotype. The different *Stb* genes were found to vary greatly in the levels of protection they provided, with no *Z. tritici* isolate found to be virulent against all tested resistance genes. Disease resistance controlled by different *Stb* genes was associated with different levels of chlorosis, with high levels of early chlorosis in some genotypes correlated with high resistance to fungal pycnidia development. *Stb10, Stb11, Stb12, Stb16q, Stb17,* and *Stb19* were identified as contributing broad spectrum disease resistance, and synthetic hexaploid wheat lines were identified as particularly promising sources of broadly effective STB resistances. Wheat genotypes carrying multiple *Stb* genes were found to provide higher levels of resistance than expected given their historical levels of use. The knowledge obtained here will aid UK breeders in prioritising *Stb* genes for future breeding programmes. In addition, this study identified the most interesting *Stb* genes for cloning and detailed functional analysis.

**Introduction**

Septoria tritici blotch (STB), caused by the fungal pathogen *Zymoseptoria tritici*, is one of the most damaging wheat diseases across Europe, with the capacity to cause up to 50% crop losses under disease-favourable conditions (Fones and Gurr, 2015). Approximately 70% of the fungicides used in Europe can be for the purpose of preventing *Z. tritici* epidemics (Duveiller *et al.*, 2007; Torriani *et al.*, 2015). Developing methods for protecting wheat from STB is therefore a high priority for UK wheat breeders and researchers.

Traditionally, STB protection has been achieved through the widespread application of fungicides reinforced with the deployment of a small number of *Stb* resistance genes. However, the sexual reproductive cycle that *Z. tritici* undergoes around the end of the breeding season can contribute to high levels of genetic diversity in the pathogen, leading to the rapid loss of effectiveness from fungicides. Resistant strains now exist for every major fungicide group used against them (Fraaije *et al.*, 2005; Cools and Fraaije, 2008; Stammler and Semar, 2011; Hillocks, 2012; van den Berg *et al.*, 2013; Estep *et al.*, 2015). A similar lack of durability has proven an issue with *Stb* resistance genes. For example, *Stb6* and *Stb15* have both been widely used in Northern Europe and were initially highly effective however, both have since been widely broken by *Z. tritici* due to the selection pressures caused by their widespread use (Arraiano *et al.*, 2009; Stephens *et al.*, 2021). *Stb16q* has also been brought into wide use more recently in some European countries, and initially offered very broad STB resistance. However, virulent isolates of *Z. tritici* against *Stb16q* have already been reported in Ireland and France (Dalvand *et al.*, 2018; Kildea *et al.*, 2020) and are likely to spread rapidly within field populations, making this resistance gene less useful in future breeding programmes. The lack of broad spectrum STB resistance in wheat leaves agricultural systems vulnerable when major resistance genes are broken (e.g. the cultivar Gene in the USA, which was fully resistant in 1992 but widely susceptible by 1995, causing substantial crop losses (Cowger *et al.*, 2000), or Cougar, which has become unpopular due to the development of Cougar-virulent strains of *Z. tritici* in the UK (Kildea *et al.*, 2021)). Such problems will only become more frequent as effective fungicide protection options become more limited (Birr *et al.*, 2021).

It is also noteworthy that some individual major resistance genes that have been widely used in breeding so far have proved to be more durable than others. For example, *Stb1* was introduced to the grower market in the cultivar Oasis in 1975 and has been used in many other cultivars (e.g. Sullivan) since 1979 and remained effective in the field up until mid-2000’s (Cowger *et al.*, 2000; Adhikari *et al.*, 2004; Singh *et al.*, 2016). *Stb4* also proved to be reasonably durable, lasting for approximately 15 years. After its introduction to breeding programs in 1975 (in a cross between Tadorna, Cleo and Inia 66), the first cultivar containing *Stb4* underwent a commercial release in 1984 (Somasco *et al.*, 1996), and this gene remained effective until 2000 (Jackson *et al.*, 2000). However, no individual *Stb* gene so far identified appears to be completely durable. Some cultivars containing multiple resistance genes (e.g. Kavkaz-K4500) seem to maintain broad resistances in the medium to long term (Chartrain, Berry, *et al.*, 2005). Gene pyramiding may be able to mitigate this rapid breaking of disease resistance by producing additional obstacles to fungal populations in the evolution of new virulences. For example, Kavkaz-K4500 is one of the most durable sources of field resistance used for breeding and has been shown to possess at least five qualitative resistance genes, including *Stb6*, *Stb10* and *Stb12* (Chartrain *et al.*, 2005a). Interestingly, this combination of *Stb* genes seems to be sufficient to make Kavkaz-K4500 resistant to STB under field conditions despite the fact that many international *Z. tritici* isolates are virulent on it in laboratory tests (Chartrain *et al.*, 2004a; Chartrain *et al.*, 2005a) – this may suggest high genetic diversity differences between UK and international *Z. tritici* populations, or could be related to the different levels of inoculum used in laboratory vs field trials.

The currently limited availability of data on the interaction between modern *Z. tritici* isolates and wheat (due to limited numbers of isolates being tested in most studies and the fact that many older isolates are reused in many studies for example, 22 isolates from one set of plots at a single location were used in Cowger *et al.* (2000), ten isolates from a range of Iranian farms in Dalvand *et al.* (2018) and only one 1996 isolate in Ali *et al.* (2008)), along with the difficulty in comparing data from different sources, is problematic as it has limited our ability to identify useful sources of quantitative resistances to this disease (Chartrain *et al.*, 2004). This combined with the lack of historical breeding for STB resistance, has led to a dearth of cultivars with significant quantitative resistance to the disease.

Further issues arise from the lack of standardised, modern wild-type *Z. tritici* isolates among the standard model strains for this disease, which represents a significant obstacle to the development of durable STB resistance in wheat due to the difficulties it causes in designing experiments that produce useful information on the likely field efficacy of resistance genes and QTLs for breeders and can be easily compared to other work in the same field. It is therefore important that new field isolates of *Z. tritici* are collected from all regions of interest for breeders to be used in the testing of new resistance genes. A database of *Z. tritici* isolates with known virulence profiles could help identify combinations of *Stb* resistance genes that could provide several independent resistances for each tested *Z. tritici* isolate. This could allow us to identify combinations of resistance genes that would require several independent mutations in any *Z. tritici* isolate in order for that isolate to gain virulence.

This rapid breakdown of existing resistances makes it particularly important that breeders have access to novel STB resistance genes effective against local *Z. tritici* populations. Several known major resistance genes, such as *Stb5*, *Stb17* and *Stb19*, have not previously been widely used in Europe, and could perhaps be used to replace those that have already been overcome (e.g. *Stb6* and *Stb16q*). Unfortunately, little data is currently available to breeders regarding which of these genes are sufficiently broadly effective to be worth using in breeding programs.

It is therefore clear that a future priority in wheat breeding is likely to be the development of elite lines containing a greater variety of disease resistance genes. Major resistance genes are likely to be a large part of this as they can be identified easily and applied quickly in breeding programs, and major genes not yet broken will provide excellent field resistance. More than twenty *Stb* resistance genes that could be used in wheat breeding programs have thus far been identified, providing natural protection against a variety of *Z. tritici* isolates at the different stages of the wheat life cycle (referred to as seedling and adult resistance genes) (Dreisigacker *et al.*, 2015). For many of these *Stb* genes we have some information relating to their chromosomal locations, but in the majority of cases this data is imprecise.

Overall, large pathology screens are necessary to assess the effectiveness of *Stb* genes more accurately. Conducting these screens on more genetically diverse germplasm (particularly non-elite landraces and ancestor species) may help to identify novel *Stb* genes highly effective against current *Z. tritici* populations. Here we carried out a broad screen of 2015-2017 UK *Z. tritici* isolates against a panel of wheat lines of diverse origin containing known *Stb* resistance genes to produce estimates of the effectiveness of each of these genes against contemporary field populations of *Z. tritici* in the UK. Several *Stb* genes were identified as contributing broad spectrum disease resistance, and synthetic hexaploid wheat lines were identified as promising sources of broadly effective STB resistance.

**Materials and Methods**

## Library of fungal isolates

One hundred *Z. tritici* isolates were donated by Bart Fraaije (NIAB, UK). These isolates were collated from locations around the UK in the years 2015-2017. These isolates were originally drawn from many sources with different naming conventions, and were renamed for ease of use in this project – a list of the original names of these isolates on receipt is included in the supplementary data.

In preparation for use in these experiments, these isolates were grown on 7% (w/v) YPD agar (Formedium - Hunstanton, UK) plates containing 1 unit of penicillin and 1 µg/mL streptomycin (Merck Life Science UK Limited, Gillingham, UK) to remove bacterial contamination. Approximately 25 µl of original *Z. tritici* glycerol stocks were used per plate. Inoculated plates were incubated at 16⁰C for four to seven days before the fungus was harvested using a sterile loop into 50% (w/v) glycerol and stored at ‑80⁰C. This was then repeated using antibiotic free (otherwise identical) plates to ensure the fungi used were not stressed. Fungi from antibiotics-free plates were harvested and stored identically.

Where bacterial contaminants proved resistant to the antibiotics used, contaminated glycerol stock was diluted (approximately by a factor of 100, depending on concentration), allowing individual colonies to form from single cells. Suitable uncontaminated colonies were harvested into 50% glycerol and re-plated to produce pure stocks.

## Wheat lines used

Wheat lines were chosen for use in this study that collectively contained *Stb* resistance genes *Stb1-Stb19*. These lines and the *Stb* genes they contain are listed in Table 1. Taichung 29 and KWS Cashel were both included as known low-resistance susceptible controls (of these, KWS Cashel was the primary control and Taichung 29 was included as a second control in case KWS Cashel was found to be resistant to any *Z. tritici* isolates used).

## Inoculation of wheat plants

*Z. tritici* isolates used in inoculations were cultured on antibiotic-free YPD agar plates and grown for four to seven days at 16⁰C. Fungal blastospores were then harvested using sterile loops into 5mL of 0.1% Silwet L-77 surfactant (Momentive Performance Materials, Waterford, NY, USA) in H2O and diluted to a concentration of 107 spores per mL using the average of two replicated measurements from a haemocytometer. High concentrations and the presence of a surfactant are not reflective of field conditions but were included to encourage rapid infection to reduce the time needed per bioassay.

Plants were grown for approximately three weeks (adapted for variable growth rates where necessary) at 16-hour day, 8-hour night cycles under halogen or white LED lamps at a temperature of 21°C and ambient humidity. After inoculation, these plants were transferred to 17°C and the same 16-hour day, 8-hour night cycle. The second leaf was inoculated where possible, although for some cultivars (e.g. Israel 493) the third leaves were used due to their larger size.

Leaves were affixed to aluminium inoculation tables using double sided sticky tape and rubber bands, which also defined the area inoculated and scored. Cotton buds were used to inoculate each spore suspension onto leaves of three plants of each wheat line (four strokes per leaf, ensuring an even layer of moisture on leaf surface). Non-inoculated leaves were trimmed to ensure light access to inoculated leaves.

After inoculation, plants were placed in high humidity boxes (Supplementary Figure 1) for three days before the inner tray (perforated to allow for water uptake) was removed and placed in a larger plastic watering tray to minimise the risk of causing leaf damage or cross-contamination from direct watering.

Plants were maintained for 28 days after inoculation to allow symptom development. They were watered three times per week and kept trimmed to ensure light access to inoculated leaves. From ten days post inoculation (dpi), plants were checked regularly (every two days where possible) for chlorosis, necrosis and pycnidia development, and symptoms were recorded. Photographs were taken at each check for later verification.

The final screen included 973 tested interactions. Due to the large number of wheat genotype – *Z. tritici* isolate interactions tested, one replicate was normally performed for each of these interactions in the bioassay.

## Visual symptom assessments

Necrosis, chlorosis and pycnidia development symptoms were assessed visually. Assessment of the rate of symptom and pycnidia development began ten days after seedling inoculation by *Z. tritici* for each plant. Assessments were then carried out three times a week at regular intervals until 28 days after the initial inoculation date. Leaf status was recorded as no infection (i.e. clean), chlorosis present (showing yellow chlorotic tissue but which had not yet progressed to necrosis), necrosis present (where necrotic lesions were visible), chlorosis with pycnidia (chlorotic symptoms present with small black pycnidia visible on the inoculated leaf surface) or necrosis with pycnidia. The first date on which chlorosis or necrosis was seen was used to determine the “days until symptom development” symptom value, while the date on which pycnidia were first noted was used to determine the “days until pycnidia development” symptom value. Photographs were taken at each check in case needed for later verification of results.

At 28 days post infection, before leaves were harvested for the pycnidia spore count measurements, the “percentage leaf area covered by symptoms” and “percentage leaf area covered by pycnidia” were visually assessed. The values for each leaf were rounded to 0, 20, 40, 60, 80 or 100% for each leaf. Photographs were taken in case needed for later verification of results.

## Pycnidia spore counts

After 28 days post infection, inoculated leaf regions were harvested into 15 mL Falcon tubes (one for the three leaves of each line/isolate interaction). A 2 cm X 2 cm X 1.5 cm plug of absorbent cotton wool pre-wetted with 3 mL of deionised H2O was used to provide a humid environment in each tube (encouraging pycnidia to push through stomata and ooze pycnidiospores) for 2 days before measurement.

Six mL of 0.01% Tween 20 surfactant (Croda International Plc, Snaith, UK) in H2O was then added to each tube, and tubes were vortexed for 75 s to wash spores from leaf surfaces into the liquid. The optical density at 600 nm (OD600) of one mL of the resulting suspension was measured using a spectrophotometer CARY 50 (Varian, London, UK).

The spore suspensions giving the highest OD600 were used to produce standard curves to convert OD600 ratings to spores/mL. This required a series of standard dilutions (2x, 4x, 8x, 16x and 32x) of the spore suspension in 0.01% Tween 20 in H2O to be measured with the spectrophotometer and haemocytometer (two haemocytometer measurements were averaged to provide the measurement used). In most cases curves were approximately linear, so the formula of the linear trendline (generated in Microsoft Excel) was used in conversions. Supplementary Figure 2 demonstrates the relationships between OD600 and haemocytometer spore count readings for a large set of leaves.

## Statistical analysis

Statistical tests were carried out using the statistics package R (Team, 2013) to run paired Student’s *t*-tests on data from different wheat lines (results obtained using the same *Z. tritici* isolate in the same experimental set were treated as paired) using standard R commands for this function. The large numbers of *Z. tritici* isolates tested against the wheat genotypes of interest allowed for statistical assessments of the average broad resistance of each line. ANOVA tests were used when data from multiple wheat lines was to be compared, and to verify results produced from the *t*-tests – this was done using standard R and Excel Data Analysis commands.

**Results**

**Comparative Assessment of Average *Z. tritici* Resistance in Wheat Genotypes Based on Five Phenotype Assessments**

Seventeen wheat genotypes carrying no known *Stb* gene, a single *Stb* gene, or a combination of *Stb* genes were screened against up to 100 current UK *Z. tritici* isolates. The symptoms of each genotype were compared to those of KWS Cashel, used as the susceptible control. The P-values derived using a standard student’s t-test to compare the average % pycnidia coverage of inoculated leaf area and the spore count data from spore washes for each *Z. tritici* isolate-resistant wheat line to the equivalent averages from interactions with the KWS Cashel susceptible control are shown in Table 2 – these data show which lines have significantly different symptom development levels overall than KWS Cashel (P<0.05). Mean average values the full set of genotype-isolate comparisons tested on each wheat line are given in Table 3 for each of the five measured symptoms. The proportion of isolate-wheat line interactions for which disease symptoms were entirely absent for chlorosis/necrosis and for pycnidia development is shown in Table 4.

Inoculated wheat plants were assessed for five STB disease associated symptoms: the times (dpi) taken to the development of chlorosis/necrosis symptoms and fungal pycnidia, the final percentage of the inoculated leaf sections covered by chlorosis/necrosis, the final percentage of the inoculated leaf sections covered by pycnidia, and the estimated number of pycnidiospores washed off infected leaf section. There was a significant biological variation in the rates of development of chlorosis and necrosis symptoms and percentage of leaf coverage by chlorosis/necrosis within some wheat line – *Z. tritici* isolate interactions (potentially caused by variation in factors such as sunlight levels or damage done during inoculation). The percentage of leaf covered by pycnidia was more consistent for wheat line – *Z. tritici* isolate interactions, and thus was used as the primary factor used to differentiate between virulence and avirulence.

The pycnidia coverage of all other genotypes was significantly different from the susceptible KWS Cashel in two-tailed paired Student’s *t*-tests, as shown in Table 2. The average pycnidia coverage for KWS Cashel was also higher than that for any other line (including Taichung 29, which also does not have any known *Stb* genes – this difference may be due to differences in the plant leaf architecture resulting in fewer penetration events, or by low-effect resistance QTLs that may provide slightly improved non-specific resistances in Taichung 29). Therefore, all wheat lines were significantly more resistant than KWS Cashel using the symptom measurement that most reliably differentiates virulence and avirulence. The estimated spore counts for most other genotypes were also significantly different from those for KWS Cashel, as shown in Table 2. KWS Cashel also has the highest average estimated spore count of all tested lines. This indicates that all tested wheat lines except Riband have higher average resistance to UK *Z. tritici* isolates than KWS Cashel for the symptom most directly connected to these isolate’s ability to cause an epidemic in field wheat populations.

It should be emphasised that these results are calculated by averaging disease assessment scores from many individual *Z. tritici* isolates tested for each wheat genotype. Resistant genotypes, such as TE9111, Kavkaz-K4500 and Synthetic 6X were generally resistant to almost all isolates tested. However, genotypes, such as Tadinia had far more variable resistance, with some isolates inducing high infection scores across all assessment criteria while others produced no symptoms, generating intermediate average scores (Table 3). This suggests that these resistances are specific to fungal isolates carrying particular avirulence factors (a “gene-for-gene” relationship) which are each present in only some UK *Z. tritici* isolates. This also indicates that the underlying resistance mechanisms are highly effective when recognition occurs early in *Z. tritici* development, even against isolates with the potential to be highly virulent on other lines.

In most cases, wheat genotypes displayed similar symptom severity across all measurements. However, for some genotypes (e.g. Israel 493) the development rate and final percentage leaf coverage of chlorosis were high compared to the final percentage of pycnidia leaf coverage and pycnidiospore production. Similarly, early chlorosis followed by high resistance to pycnidia development were seen in Synthetic 6X and Synthetic M3, although not all *Z. tritici* isolates stimulated visible chlorosis development in these lines (e.g. RResHT-8 and RResHT-10 show 33-86% chlorosis in both Synthetic 6X and Synthetic M3, whereas RResHT-21 and RResHT-24 show 0-7% chlorosis in both lines).

The results obtained in this study demonstrate great variability between the resistances of different wheat lines to UK *Z. tritici* isolates. As expected, wheat lines containing no known *Stb* genes are by far the least resistant group, with almost all tested isolates being highly virulent against KWS Cashel and Taichung 29. This indicates the very low levels of non-specific resistance for *Z. tritici* present in most wheat lines.

Overall, in addition to the wheat genotypes Taichung 29 and KWS Cashel (no known *Stb* genes), Riband (*Stb15,* common and widely broken in Europe (Arraiano *et al.*, 2009)) was more susceptible than other lines. Estanzuela Federal (*Stb7*) also showed low resistance to most isolates tested (though higher than in fully susceptible lines for pycnidia coverage and spore counts), indicating that UK *Z. tritici* populations are virulent towards *Stb7* and *Stb15*. Tonic also showed relatively low resistance in key symptoms, although it was less susceptible than Taichung 29, KWS Cashel or Riband.

Israel 493 (*Stb3* and *Stb6*) and TE9111 (*Stb6, Stb7* and *Stb11*) showed relatively high levels of resistances, indicating that *Stb3* and *Stb11* could have high potential interest to UK breeders. The synthetic and synthetic-derived lines Synthetic 6X, Synthetic M3 and Lorikeet also demonstrated high levels of resistance, likely due to their novel *Stb* resistance genes (*Stb5*, *Stb16q* and *Stb17*, and *Stb19* respectively). Kavkaz-K4500 (*Stb6*, *Stb7, Stb10* and *Stb12*) provides good levels of resistance, likely due to the presence of *Stb10* and *Stb12* (as *Stb6* is known to be widely broken and *Stb7* has been shown to be ineffective due to the susceptibility of Estanzuela Federal).

The lines Tadinia, Balance, Synthetic M6, Bulgaria 88, Veranopolis, and Salamouni had more intermediate average levels of resistance, indicating that the genes *Stb1, Stb2, Stb4, Stb8, Stb9, Stb13, Stb14* and *Stb18* all provided partial resistance, or provided resistance to some but not all *Z. tritici* isolates tested. These *Stb* genes could also be interesting to breeders as most would take relatively little effort to move into new wheat cultivars, and are likely to produce reasonable levels of resistance under field conditions (where inoculum levels will be lower than in these screens). However, the genetic variability of *Z. tritici* in the field suggests that individually these genes are unlikely to offer stable resistance, as at least one *Z. tritici* isolate will be virulent against each. It is likely that these genes would have to be stacked to provide durable resistance, slowing and complicating the breeding process.

It was notable that Riband, Estanzuela Federal and Tonic possessed the least resistance among *Stb* gene containing genotypes. Riband showed the highest levels of pycnidia and pycnidiospores amongst the lines possessing at least one *Stb* gene. This is likely to be because *Stb15* is known to have been widely present in European wheat lines historically (Arraiano *et al.*, 2009), meaning that the local *Z. tritici* populations have adapted to its presence. Tonic had the second highest levels of pycnidiospore production and Estanzuela Federal having the second highest levels of pycnidia coverage. This suggests that the *Stb* genes found in these lines (*Stb7*, *Stb9* and *Stb15*) do not provide good resistance to most *Z. tritici* isolates present in the UK population and should be considered low priority breeding targets for UK wheat lines (although these genes may be more effective against *Z. tritici* populations in other parts of the world).

**Identification of Preferential Breeding Targets for Maximising the Durability of STB Resistance Genes**

The broadest complete resistances were found in Synthetic M3, Kavkaz-K4500, TE9111 and Lorikeet. These genotypes collectively contain *Stb6, Stb7, Stb10, Stb11, Stb12, Stb16q, Stb17,* and *Stb19*. However, the *Z. tritici* isolates used in this test were selected from a dataset of isolates known to be virulent against lines containing *Stb6*. Additionally, *Stb6* and *Stb7* were present in less resistant lines (e.g. Veranopolis and Estanzuela Federal), likely indicating that these *Stb* genes contributed minimally to the resistances of these cultivars.

In Kavkaz-K4500 and Synthetic M3, *Stb10* is paired with *Stb12* and *Stb16q* is paired with *Stb17,* respectively. As none of the genotypes tested contained these genes individually, it is difficult to determine from these results what proportion of the resistances each gene in these pairs was responsible for. It should be noted that previous experiments and field observations demonstrate that *Stb16q* provides extremely broad resistance to the UK *Z. tritici* population present in 2015-2017 (Tabib Ghaffary *et al.*, 2012; Saintenac *et al.*, 2021) whilst *Stb17* was demonstrated to act primarily in adult plants, older than the seedlings used in this study (Tabib Ghaffary *et al.*, 2012), indicating that *Stb16q* is likely to be responsible for most of the resistance seen in Synthetic M3.

Further experimentation using NIL lines containing each of these genes individually will aid determining for certain which provide the broadest resistance – until such time as this work is completed, *Stb5,* *Stb11* and *Stb19* appear to be the highest priority breeding targets found in these bioassays.

**Identification of a class of STB resistance responses associated with strong early leaf chlorosis and reduced pycnidia production**

An examination of the level of resistance to different symptoms of *Z. tritici* infection in each wheat genotype also reveals a broader category of potentially interesting *Stb* genes that show high levels of resistance to pycnidia development but are not protected from the early development and high final coverages of chlorotic and necrotic symptoms on the leaves. For example, Israel 493 (containing *Stb3* and *Stb6*) shows the sixth highest average symptom coverage score of all tested genotypes (the fourth highest amongst genotypes possessing at least one *Stb* gene), yet has negligibly low average levels of pycnidia coverage, as shown in Figure 1. This could indicate the presence of resistance genes that act specifically to disrupt the pycnidia formation stage of fungal pathogen development or the presence of resistance pathways which cause chlorosis as a side effect less damaging then allowing the fungus to grow unimpeded, although it seems unlikely that chlorosis is directly tied to the resistance mechanism as chlorosis is usually linked with cell death and *Z. tritici* is primarily necrotrophic.

This unusual combination of symptoms could indicate the activation of resistance mechanisms involving a hypersensitive response, likely involving early reactive oxygen species-producing reactions in the chloroplasts (as indicated by the early and strong chlorosis response). This resistance mechanism seems likely to be effective at preventing the spread of a *Z. tritici* epidemic in the field by preventing pycnidia development, although there may also be some loss of photosynthetic potential from individual plants. This could suggest that *Stb3* and other resistance whose action is associated with high levels of chlorosis genes could provide more durable resistance if deployed in combination with other resistance genes, not associated with chlorosis, as the two different resistance mechanisms would be difficult for any *Z. tritici* isolate to adapt to. However, the utility of these resistances is likely to depend on the level of loss of photosynthetic potential in the field, which cannot easily be estimated from this work, as the high levels of inoculum used to ensure infection here are unrealistic under normal field conditions. Additionally, it is not known which resistance response would be activated against isolates avirulent on wheat genotypes containing both resistance genes associated with chlorosis and those that do not associate with chlorosis. Further experimentation and fieldwork are needed to determine the utility of combining these two mechanistically different types of resistance genes.

**Discussion**

*Zymoseptoria tritici* is one of the most important pathogens in the wheat-based agricultural systems of Europe, and chemical defences against it do not seem likely to be durable in the long term. It is therefore vital that breeders be able to effectively utilise *Stb* resistance genes to prevent major epidemics. This study provides data that will help to target UK breeding efforts to the most effective *Stb* resistance genes.

Data provided by field trials can be difficult to standardise due to genetic differences in *Z. tritici* populations locally (Berraies *et al.*, 2013; Mekonnen *et al.*, 2020) and globally, and due to the dramatic effect of weather conditions (particularly rainfall) on STB disease development, which can cause large fluctuations in readings between years at the same sites (Ouaja *et al.*, 2020). Additional complexities are added to data analysis by wheat lines with resistance levels that change over the wheat life cycle (e.g. high seedling and low adult resistance) and by imperfect correlations between the levels of different infection symptoms (e.g. necrosis levels and pycnidia counts) (Ouaja *et al.*, 2020). This information is particularly lacking for novel STB disease resistance sources, such as synthetic hexaploid wheats. Overall, the results presented here suggest that the lines Lorikeet (containing *Stb19*) and Synthetic M3 (containing *Stb16q* and *Stb17*) should be of the greatest interest to breeders, as these genotypes were resistant to pycnidia formation from every *Z. tritici* isolate they were challenged with in our bioassays, along with Kavkaz-K4500 (containing *Stb6, Stb7, Stb10* and *Stb12*), Synthetic 6X (containing *Stb5*) and TE9111 (containing *Stb6*, *Stb7* and *Stb11*), which had very high overall resistance. However, Synthetic M3 carries two *Stb* genes, *Stb16q* and *Stb17*. Of these, previous research suggests that *Stb17* is effective only in adult plants (Tabib Ghaffary *et al.*, 2012), suggesting that the Synthetic M3 resistance is primarily due to the effect of *Stb16q*, which is known to provide broad resistance against *Z. tritici*. However, it should be noted that the resistance provided by *Stb16q* in the field is likely to be less complete than these results suggest, as the bioassays described here used UK *Z. tritici* isolates collected between 2015 and 2017. Since these dates, use of *Stb16q* in elite wheat lines has led to selection for *Z. tritici* isolates capable of virulence against lines containing this resistance, e.g. those found in Ireland and Iran (Dalvand *et al.*, 2018; Kildea *et al.*, 2020), which will likely lead to reductions in the field effectiveness of *Stb16q* over the coming years (as has previously been seen for *Stb6* and *Stb15*). This effect has not yet been noted for the resistance gene *Stb19*, which has not been used in the UK thus far. However, it seems likely that wider use of *Stb19* in elite lines would favour the development of *Z. tritici* isolates capable of breaking this resistance, leading to the loss of efficacy of this resistance gene. It is therefore important that when *Stb19* is used, it is supported by additional genes that provide broad resistance to the local *Z. tritici* population.

The results of this bioassay suggest Kavkaz-K4500 (*Stb6, Stb7, Stb10* and *Stb12*), Synthetic 6X (*Stb5*) and TE9111 (*Stb6, Stb7* and *Stb11*) as good potential sources for these protective *Stb* resistance genes. These genotypes show no pycnidia development from 98%, 96% and 95% of tested *Z. tritici* isolates respectively, with low pycnidia coverages (a maximum of 20% average) from the remaining isolates. All isolates tested against all three genotypes proved avirulent against at least one. As results from Estanzuella Federal and previous research suggest that *Stb6* and *Stb7* provide little or no resistance from UK *Z. tritici* populations (Czembor *et al.*, 2011; Makhdoomi *et al.*, 2015; Stephens *et al.*, 2021), it seems likely that *Stb5, Stb11* and either *Stb10* or *Stb12* are responsible for these resistances. As *Stb10* and *Stb12* were not available for testing in isolation, it was not possible in this study to assess proportion of the total Kavkaz-K4500 resistance was associated with each of these genes. Therefore currently *Stb5* and *Stb11* appear to be the optimal resistances to protect the durability of *Stb19* in future wide use. The long-term effectiveness of the Kavkaz-K4500 resistance despite the widespread use of this genotype in breeding suggests that such pyramids of mutually protective *Stb* genes are likely to be effective in slowing the development of virulence against them in *Z. tritici* populations.

The most useful *Stb* genes identified here are novel genes originating from synthetic hexaploid wheat lines and those that have historically been protected by the presence of multiple resistances in a single breeding line. This may cause issues during the breeding process, as synthetic-derived lines could carry undesirable genes (causing linkage drag when resistances are transferred to elite lines, possibly reducing yields) and effective resistances may be difficult to identify from wheat lines in which they coexist with several ineffective resistances. The high average resistance of novel lines aligns well with the results of (Arraiano and Brown, 2006), which found that of 238 wheat genotypes tested, the line with the highest non-specific resistance in their study was the Italian landrace Rieti. Although the resistances identified as broadly effective in this study were highly specific rather than non-specific, both results still indicate that the time given for *Z. tritici* to adapt to widely used resistances is a vital determining factor in their effectiveness. However, the (Arraiano and Brown, 2006) paper utilised IPO isolates, which are now severely outdated and several generations removed from current wild *Z. tritici* populations, along with detached leaf assays, which may cause issues with measuring symptoms such as necrosis coverage (which (Arraiano and Brown, 2006) did not attempt to monitor). This study used more recent field isolates of *Z. tritici* collected from a more localised region around the UK and tested against a smaller set of wheat genotypes, producing a dataset more optimally targeted for identifying resistance genes of interest to breeders in this area. This study also selected wheat genotypes for testing based on the presence of known major resistance genes whereas (Arraiano and Brown, 2006) aimed to test a broader set of wheat genotypes for any resistance regardless of genetic origin, which together with the more modern *Z. tritici* isolates used in the present study makes it difficult to draw direct conclusions from differences in the average resistances observed.

Resistance to *Z. tritici* is a relatively new target in wheat breeding, meaning that much of the research relating to this pathogen and its interactions with crop plants is still in the early stages and major details of the infection and resistance processes (e.g. potential *Z. tritici* effector impacts on host chloroplast function or the mechanisms of most *Stb* gene for gene resistances) are largely unknown at a molecular level. With respect to *Avr-R* interactions, only *Stb6* and *Stb16q* have been cloned (along with the corresponding fungal effector AvrStb6 recognised by *Stb6*) (Zhong *et al.*, 2017; Saintenac *et al.*, 2018; Saintenac *et al.*, 2021). Much of the research conducted thus far has utilised the model isolate held by most laboratories, IPO 323 – however, this isolate is not reflective of modern field isolates in important ways. For example, IPO 323 is naïve to all modern fungicides and avirulent on cultivars with disease resistance genes that have now been broken down by a large majority of isolates found in the field (e.g. *Stb6*). It is therefore important that novel *Stb* resistance genes be tested more broadly against collections rather than single *Z. tritici* isolates, to assess whether they act sufficiently broadly to be useful in a commercial growing context. The *Z. tritici* isolates utilised in this study were selected from UK fields between the years 2015 and 2017, and are virulent against *Stb6*. Although these isolates have not been sequenced, the range of different resistance responses they triggered in some wheat genotypes suggests a high level of genetic diversity. This is supported by the well-established genetic diversity of Z. tritici even in limited geographic regions (Berraies *et al.*, 2013; Mekonnen *et al.*, 2020; Orellana-Torrejon *et al.*, 2022) and indicates that the results identified here should be broadly applicable to UK *Z. tritici* populations.

It should also be noted that while the resistance profiles (the specific lists of *Z. tritici* isolates to which each genotype was resistant or susceptible) of resistant wheat genotypes (wheat genotypes containing broadly effective *Stb* resistance genes) were more similar to the resistance profiles of other resistant wheat genotypes than they were to those of susceptible wheat genotypes (wheat genotypes that did not contain broadly effective *Stb* resistance genes), there is otherwise little correlation between the resistance profiles of highly resistant lines (the specific *Z. tritici* isolates that were virulent or susceptible against each wheat genotype containing broadly effective *Stb* resistance genes) (Data not shown). This suggests that each *Stb* gene recognises a different fungal elicitor. Additionally, no *Z. tritici* isolate was identified that was virulent towards all tested *Stb* genes. This suggests a high level of variation among *Stb* genes, matching that of highly variable *Z. tritici* populations. Therefore, it should be possible to develop highly resistant breeding lines by stacking many *Stb* genes. Such gene pyramids would likely improve the durability of all *Stb* genes included (provided that these *Stb* genes were only used in such gene pyramids), as it is much less likely that any given isolate would gain all of the required mutations for virulence at once and thus overcome the resistance. This could be extremely useful in the long term – for example, Kavkaz-K4500 has been considered an STB resistant breeding line for many years and still appeared effective in our experiments, suggesting that combinations of resistance genes that utilise different mechanisms may not only help to increase the durability of each individual gene, but could also be broadly effective due to the collective action of these genes. The use of modern genetic markers and breeding techniques will be necessary to overcome potential obstacles to breeding such as linkage drag and epistasis effects – for example, markers could help track specific resistance genes present in breeding materials derived from genotypes containing multiple *Stb* genes, and the production of NILs using such markers could limit the effect of linkage drag on new breeding lines.

In summary, this study revealed that sufficiently diverse *Stb* genes exist to give broad and durable protection from UK *Z. tritici* isolates to new wheat lines. However, generating this protection in a sustainable form will require extensive breeding efforts. We identified suitable *Stb* genes to prioritise for pyramiding. However, further work will be necessary to identify modern high-throughput markers such as Kompetitive Allele Specific PCR (KASP) markers (Semagn *et al.*, 2014) for each *Stb* gene of interest to ensure that multiple broadly effective genes can be stacked in a single line (as otherwise epistatic effects may make their presence difficult to confirm), and to produce lines containing each *Stb* gene from highly resistant lines individually for further detailed characterization. There therefore remains much work to be done collaboratively between UK wheat breeders and the scientific community to ensure the desired level of resistance in future wheat.

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**Figure Legends**

**Figure 1:** Showing the early chlorosis development identified on several Synthetic 6X leaves after inoculation with three *Z. tritici* strains. Pycnidia symptoms are not seen.

**Tables**

**Table 1:** Wheat lines used in this study with known *Stb* genes.

|  |  |  |
| --- | --- | --- |
| **Name of Wheat Line Used** | **Known *Stb* Resistance Genes** | **Papers Reporting Presence of *Stb* Resistance Genes in Wheat Line** |
| **Taichung 29** | No *Stb* Genes Known | - |
| **KWS Cashel** | No *Stb* Genes Known | - |
| **Bulgaria 88** | *Stb1, (Stb6)* | Adhikari *et al.*, 2004 |
| **Veranopolis** | *Stb2, (Stb6)* | Liu *et al.*, 2013 |
| **Israel 493** | *Stb3, (Stb6)* | Goodwin *et al.*, 2015 |
| **Tadinia** | *Stb4, (Stb6)* | Adhikari *et al.*, 2004 |
| **Synthetic 6X** | *Stb5* | Arraiano *et al.*, 2001 |
| **Estanzuela Federal** | *Stb7* | McCartney *et al.*, 2003 |
| **Synthetic M6 (Previously W7984)** | *Stb8* | Adhikari *et al.*, 2003 |
| **Tonic** | *Stb9* | Chartrain *et al.*, 2009 |
| **Kavkaz-K4500** | *(Stb6), Stb7, Stb10, Stb12* | Chartrain, Berry, *et al.*, 2005 |
| **TE9111** | *(Stb6), Stb7, Stb11* | Chartrain, Joaquim, *et al.*, 2005 |
| **Salamouni** | *(Stb6), Stb13, Stb14* | Cowling, 2006 |
| **Riband** | *Stb15* | Arraiano *et al.*, 2007 |
| **Synthetic M3** | *Stb16q, Stb17* | Ghaffary *et al.*, 2012 |
| **Balance** | *(Stb6), Stb18* | Ghaffary *et al.*, 2011 |
| **Lorikeet** | *(Stb6), Stb19* | Yang *et al.*, 2018 |

**Table 2:** A comparison of the % pycnidia coverage of inoculated leaf area and spore count data from spore washes for each *Z. tritici* isolate-resistant wheat line interaction and the equivalent values derived from the *Z. tritici* isolate’s interactions with the KWS Cashel susceptible control. Mean averages from each interaction (calculated using the standard function in excel) were compared to those using KWS Cashel as a host using a two-tailed Student’s *t*-test from the excel data analysis tool. The P-Values resulting from this are shown below. Both symptoms for all interactions except the Spore Count symptom for the Riband genotype show significant differences to the KWS Susceptible control.

|  |  |  |
| --- | --- | --- |
| **Wheat Genotype** | **P-Value in two-tailed Student’s *t*-test against KWS Cashel for measured symptom** | |
| **Pycnidia Coverage (from % Coverage)** | **Spore Count (from Estimated Spores per Leaf)** |
| **Taichung 29** | **1.5X10-5** | **6X10-6** |
| **Riband** | **3X10-4** | **0.10** |
| **Synthetic 6X** | **4.6X10-12** | **2X10-11** |
| **Synthetic M3** | **1.6X10-10** | **2X10-11** |
| **Kavkaz-K4500** | **4X10-13** | **1X10-11** |
| **Tadinia** | **7.3X10-8** | **2X10-7** |
| **Estanzuela Federal** | **1X10-10** | **3X10-9** |
| **Israel 493** | **7.2X10-16** | **1X10-12** |
| **TE9111** | **5.8X10-18** | **6X10-14** |
| **Bulgaria 88** | **2.1X10-6** | **2X10-6** |
| **Veranopolis** | **1.9X10-6** | **2X10-5** |
| **Synthetic M6** | **4.4X10-5** | **1X10-4** |
| **Tonic** | **1.3X10-2** | **0.02** |
| **Salamouni** | **3X10-4** | **8X10-4** |
| **Balance** | **2.3X10-6** | **4X10-6** |
| **Lorikeet** | **5.9X10-7** | **7X10-7** |

**Table 3:** **The average symptoms on inoculated leaves of each wheat genotype.**



**Table 4: The proportion of *Z. tritici* isolates that did not generate symptoms on each type on each wheat genotype.**

