High-Density Mapping of Triple Rust Resistance in Barley Using DArT-Seq Markers

Peter M. Dracatos**1, Rouja Haghdoust†1, Ravi P. Singh2,3, Julio Huerta Espino2,3, Charles W. Barnes1, Kerrie Forrest5, Matthew Hayden5, Rients E. Niks6, Robert F. Park1 and Davinder Singh1

1 Plant Breeding Institute Cobbitty, Sydney Institute of Agriculture, The University of Sydney, Sydney, NSW, Australia, 2 International Maize and Wheat Improvement Center, Texcoco, Mexico, 3 Campo Experimental Valle de México, INIFAP, Chapingo, Mexico, 4 Instituto Nacional de Investigaciones Agropecuarias (INIA), Quito, Ecuador, 5 Agriculture Victoria, AgriBio, Centre for AgriBioscience, La Trobe University, Melbourne, VIC, Australia, 6 Plant Breeding, Wageningen University & Research, Wageningen, Netherlands

The recent availability of an assembled and annotated genome reference sequence for the diploid crop barley (Hordeum vulgare L.) provides new opportunities to study the genetic basis of agronomically important traits such as resistance to stripe [Puccinia striiformis f. sp. hordet (Psh)], leaf [P. hordei (Ph)], and stem [P. graminis f. sp. tritici (Pgt)] rust diseases. The European barley cultivar Pompadour is known to possess high levels of resistance to leaf rust, predominantly due to adult plant resistance (APR) gene Rph20. We developed a barley recombinant inbred line (RIL) population from a cross between Pompadour and the leaf rust and stripe rust susceptible selection Biosaline-19 (B-19), and genotyped this population using DArT-Seq genotyping by sequencing (GBS) markers. In the current study, we produced a high-density linkage map comprising 8,610 (SNP and in silico) markers spanning 5957.6 cM, with the aim of mapping loci for resistance to leaf rust, stem rust, and stripe rust. The RIL population was phenotyped in the field with Psh (Mexico and Ecuador) and Ph (Australia) and in the greenhouse at the seedling stage with Australian Ph and Pgt races, and at Wageningen University with a European variant of Psh race 24 (PshWUR). For Psh, we identified a consistent field QTL on chromosome 2H across all South American field sites and years. Two complementary resistance genes were mapped to chromosomes 1H and 4H at the seedling stage in response to PshWUR, likely to be the loci rpsEm1 and rpsEm2 previously reported from the cultivar Emir from which Pompadour was bred. For leaf rust, we determined that Rph20 in addition to two minor-effect QTL on 1H and 3H were effective at the seedling stage, whilst seedling resistance to stem rust was due to QTL on chromosomes 3H and 7H conferred by Pompadour and B-19, respectively.

Keywords: high-density linkage map, DArT-Seq markers, rust resistance, QTL, barley

INTRODUCTION

Plant pathogens of the Puccinia genus are some of the most feared and damaging diseases of cereal crops (Dean et al., 2012). Since Biblical times, cereal rust pathogens have plagued farmer's fields causing significant yield losses and in severe cases crop failure and famine (Kislev, 1982). Barley is the fourth most important cereal crop in the world and is mainly used for malt production, animal
feed and, in some regions, human consumption. Three main rust diseases currently threaten barley production: wheat stem rust, barley leaf rust and barley stripe rust, caused by *Puccinia graminis* f. sp. *tritici* (*Pgt*), *P. hordei* (*Ph*), and *P. striiformis* f. sp. *hordei* (*Psh*), with the disease abbreviated hereafter as WSR, BLR, and BYR, respectively. All three rust diseases affect malting quality through reductions in kernel plumpness, kernel weight, and germination, resulting in economic losses to producers as premiums are paid for malting grade barley (Roelfs, 1978; Dill-Macky et al., 1990; Roelfs et al., 1992; Steffenson, 1992).

BYR, caused by *Psh*, is usually a problem in cooler, wetter climates that often prevail at higher altitudes, and it is therefore renowned as a cold-temperature rust disease. Despite widespread global crop losses *Psh* has not colonized all barley growing regions, predominantly due to geographic isolation (Dantuma, 1964; Macer and Van den Driessche, 1966; Nover and Scholz, 1969; Chen et al., 1995; Chen, 2007). While *Psh* has not been detected in Australia, offshore testing of Australian barley cultivars in greenhouse seedling tests or in field adult plant tests suggest between 60 and 70% are susceptible to this pathogen (Dracatos P.M. et al., 2015; Wellings, 2007). This implies a *Psh* incursion for Australia poses a significant threat to the Australian barley industry. In contrast, leaf rust (BLR) occurs globally and is frequently detected on a seasonal basis and can cause up to 60% yield losses in susceptible varieties (Park et al., 2015). The *Ph* rust fungus has systematically evolved virulence for widely deployed leaf rust resistance genes, typically via stepwise mutation, or in regions where the alternate host *(Ornithogalum umbellatum)* is present via sexual recombination (Wallwork et al., 1992). Despite their comparatively infrequent occurrence, epidemics of WSR usually have devastating effects for both wheat and barley production (Park, 2007). Due to the involvement of the same causal pathogen as wheat (*Pgt*), early sown wheat and Triticale crops threaten late-sown barley. Furthermore, studies suggest that globally >95% of barley accessions are susceptible to the widely publicized virulent races of *Pgt* derived from Eastern Africa such as the Ug99 lineage (TTKSK) (Steffenson et al., 2017).

Genetic resistance is the most environmental friendly and economically efficient control method to reduce yield losses due to rust diseases. In contrast to wheat, where more than 80 cataloged stripe rust resistance genes exist, fewer stripe rust resistance genes have been characterized for barley. Despite this, numerous genetic studies have been performed to characterize the inheritance of stripe rust resistance in barley, especially that present in the 12 international standard differential barley genotypes (Nover and Scholz, 1969; Chen and Line, 1992). Diverse inheritance patterns mainly involving recessive resistance genes were reported by Chen and Line (1999) among the 12 international standard differential barley genotypes. They observed varying inheritance patterns including: two complementary recessive genes (Emir, Varunda, and Trumpf), two independent recessive genes (Trumpf), and single recessive (BBA 2890, Grannelose Zweizeilige, I5 and PI 548708) or both recessive and dominant genes (Abyssinian 14 and Stauffers Obersulzter). A similar situation also exists for stem rust, where there is a lack of available resistance that can be effectively deployed in barley breeding programs. The cloned *rpg4/Rpg5* gene complex is the only known resistance effective against the Ug99 lineage, although virulent races exist within North America. More recently, other resistances have been identified and characterized (Steffenson et al., 2017). In a very recent study, WSR adult plant resistance (APR) genes *Rpg2* and *Rpg3* were mapped in barley on chromosomes 2H and 5H, respectively (Case et al., 2018). Recent WSR epidemics in Sicily (Patpour et al., 2015) and Germany (Olivera Firpo et al., 2017) have highlighted the importance of diversifying resistance in barley germplasm, and also emphasize that Ug99 is not the only threat to cereal production.

In contrast, BLR resistance in barley is widely available, including 26 designated resistance genes and numerous QTLs for partial resistance (Qi et al., 1998, 1999; Park et al., 2015; Kavanagh et al., 2017; Yu et al., 2018). As previously mentioned, there are numerous examples of resistance gene breakdown (mainly those with major phenotypic effect) due to dynamic and rapid evolution in prevailing pathogen populations. Only a few leaf rust resistance genes have remained durable across different regions/environments, one of which is the APR gene *Rph20*. *Rph20* was first identified as a QTL (Rph4q) in the Dutch barley cultivar Vada (Qi et al., 1998) and was later characterized in Pompadour (Golegaonkar et al., 2010; Liu et al., 2011) and the Australian cultivar Flagship (Hickey et al., 2011) where it was mapped to chromosome 5HS (Hickey et al., 2011; Liu et al., 2011). *Rph20* was found to be present at high frequency in European barley germplasm and expressed as early as the third leaf stage (Wang et al., 2010; Singh et al., 2013). However, in some accessions with very high levels of APR, the resistance was hypothesized to be due to *Rph20* and the presence of a 2nd or 3rd genetic component (Golegaonkar et al., 2010; Wang et al., 2010; Singh et al., 2013; Rothwell et al., 2019).

Recent genomic advancements in barley have improved the ability to develop physical scaffolds and utilize sequence information for marker development. Nevertheless, whole genome sequencing of populations and/or multiple accessions for non-model crop species with large genomes such as barley is still relatively expensive. Genotyping-by-sequencing (GBS) has been used as an alternative to whole genome sequencing due to: 1/lower cost, 2/generation of high quality genetic markers, and 3/suitability of the markers for genomic prediction/selection. DArT-Seq™ technology combines the DArT array hybridization complexity reduction method (Wenzl et al., 2004) with next generation sequencing and can be optimized for any species. DArT-Seq has been used across numerous crop species for genomic diversity analysis (Baloch et al., 2017), genome-wide association studies (GWAS) (Singh et al., 2018; Visioni et al., 2018) and QTL mapping (Haghdoust et al., 2018). We have previously reported the utilization of DArT-Seq to map leaf rust (Dracatos et al., 2014; Singh et al., 2017, 2018) and stripe rust (Dracatos P.M. et al., 2015; Haghdoust et al., 2018) resistance in barley using marker-trait association and QTL mapping. In these studies, both the genetic and physical position of each marker was determined based on the Bowman consensus map and Morex physical reference assembly, respectively. To overcome possible differences in gene/marker order between the parents of the mapping population with the reference, and to improve
the accuracy of mapping genes of interest, the use of trait-specific genetic maps is the preferred approach. In this study, we constructed a high-density linkage map using DArT-Seq markers for the Pompadour x Biosaline-19 (P/B-19) RIL population and precisely mapped QTLs for resistance to stripe, leaf and stem rust across different developmental stages and environments.

MATERIALS AND METHODS

Plant Material
To study the inheritance of rust resistance at the seedling and adult plant stage, we used the F$_3$-derived recombinant inbred line (RIL) Pompadour x Biosaline-19 (P/B-19) mapping population (98 RILs) developed at the Plant Breeding Institute, University of Sydney as described by Haghdoust et al. (2018). B-19 is widely susceptible to numerous $P$. striiformis f. sp. tritici races including Ph$_{mex}$, Ph$_{24}$, Ph$_{19}$, and Ph$_{10}$, and Dutch cultivar Vada were included in every tray along with the resistant parent Pompadour. Inoculation took place in a settling tower as described by Eyal et al. (1968) using a 1:12 mixture of urediniospores:Lycopodium spores with 3 mg of urediniospore being used to inoculate each tray. One isolate, Wageningen-derived $P$. striiformis f. sp. tritici race 24 (PshWUR), was tested in two consecutive experiments. The responses of 10 barley genotypes at the seedling stage determined that PshWUR was virulent on Topper, Astrix, Atem, Berac and the susceptible control Vada, but avirulent on Agio, Bigo, Emir, Mazurka, Hiproly, Abed Binder, and Trumpf. Phenotypes were scored on a 0-4 scale, where those RILs with infection types (ITs) equal or greater than 3 were deemed susceptible and ITs 0–2 were deemed resistant.

Greenhouse Inoculation and Phenotypic Analysis for Stripe Rust Resistance
Seedlings were grown and maintained in plant boxes as described in Niks et al. (2015) with the following exceptions. Following inoculation, the plant boxes were transferred to a dark dew chamber overnight, at 10°C. The rust susceptible B-19 parent and Dutch cultivar Vada were included in every tray along with the resistant parent Pompadour. Inoculation took place in a settling tower as described by Eyal et al. (1968) using a 1:12 mixture of urediniospores:Lycopodium spores with 3 mg of urediniospore being used to inoculate each tray. One isolate, Wageningen-derived $P$. striiformis f. sp. tritici race 24 (PshWUR), was tested in two consecutive experiments. The responses of 10 barley genotypes at the seedling stage determined that PshWUR was virulent on Topper, Astrix, Atem, Berac and the susceptible control Vada, but avirulent on Agio, Bigo, Emir, Mazurka, Hiproly, Abed Binder, and Trumpf. Phenotypes were scored on a 0-4 scale, where those RILs with infection types (ITs) equal or greater than 3 were deemed susceptible and ITs 0–2 were deemed resistant.

Phenotyping for BLR and WSR Resistance
Both seedling and field screening was performed as described by Singh et al. (2017). Only one $P$. triticina pathotype 5457 P+ (virulent on Rph1, Rph2, Rph3, Rph4, Rph6, Rph9, Rph10, Rph12, and Rph19) was used to inoculate the mapping population in the greenhouse at the seedling stage and in the field over two different seasons (2016 and 2 replicates in 2018) at Plant Breeding Institute, Cobbitty, NSW, Australia. Seedlings of the P/B-19 mapping population were inoculated and ITs recorded 10 and 12 days post inoculation using the 0–4 scale (Park and Karakousis, 2002), while the P/B-19 RILs were assessed in the field when the susceptible spreader control “Gus” reached a rating of 100S using a modified Cobb scale (Peterson et al., 1948) and 1–9 McNeal scale. The inoculation and disease assessment for stem rust resistance in the greenhouse at the seedling stage was performed using $P$. graminis f. sp. tritici races 98-1,2,3,5,6 as described by Dracatos P. et al. (2015).
Both dominant and co-dominant markers with at least 70% call rate were considered for map construction. A preliminary map was constructed in R/ASMap (Taylor and Butler, 2017) using the Kosambi mapping function at LOD 6. R/ASMap was used to identify and remove redundant markers, rectify markers with switched alleles, remove duplicate samples and lines with high (≥30%) missing data, a high number of crossovers, or high (≥10%) heterozygosity, the latter also being an indicator of a mixed sample. Markers with segregation distortion (p-value <0.01 calculated from a χ² test) that had inconsistent ordering compared to the Morex genome assembly and unlinked markers were excluded. Distorted markers were not automatically discarded as some linkage groups showed regions with high density of low-level distortion, presumably with biological significance. Following marker and sample filtering, the map was reordered in R/ASMap at LOD 8.

### QTL Mapping in the P/B-19 RIL Population

We used the same genotypic dataset previously described in Haghdoust et al. (2018). A high-density linkage map was constructed comprising 8,610 markers (SNPs and in silico DArT-Seq markers). In contrast to Haghdoust et al. (2018), we selected markers every 10–15 cM based on genetic positions from the newly constructed linkage map spanning >5,000 cM. IT data from the greenhouse for leaf, stem, and stripe rust were converted to binary scores and mapped using composite interval mapping (CIM). Field-based data for leaf and stripe rust from each disease nursery were mapped using quantitative disease measurements (CIM). Field-based data for seedling resistance to the barley grass stripe rust pathogen (*P. striiformis* f. sp. *pseudo-hordei*) from Haghdoust et al. (2018) was also used for QTL analysis using the new genetic map with the aim of comparing the position of stripe rust resistance QTLs. QTL regions associated with resistance to different *Psh* isolates in the P/B-19 RIL population were considered to be the same if there was overlap between their LOD-1 support intervals.

### RESULTS

#### Greenhouse and Field Psh Inoculations and Inheritance of Resistance

In greenhouse phenotypic assessments at the seedling stage with the *PshWUR* isolate, Pompadour was highly resistant (IT = “C”) and B-19 was very susceptible (IT = “4”) based on the “0–4” IT scale described in Haghdoust et al. (2018). To determine the genetic basis of resistance in the Pompadour parent, we screened the P/B-19 RIL population at the seedling stage in the greenhouse in two separate experimental replicates (Figure 1A). Across both experimental replicates, >68% of the RILs were highly susceptible (IT = “3+” to “4”). Three phenotypic responses were observed in the P/B-19 RIL population, viz. highly resistant similar to Pompadour, intermediate (restricted pustules) and fully compatible similar to the susceptible parent B-19 (Figure 1A). Chi Squared analysis suggested that the segregation ratio best fitted a 1R : 3S ratio (χ² = 0.154 at 1 d.f.; p = 0.695), supporting the presence of two complementary resistance genes conferring seedling resistance to *PshWUR*.

At the Ecuador field site, Pompadour and B-19 had a mean disease rating of 2.5 and 90% disease severity, respectively. Very similar results were observed in Mexico in 2015 and 2016 seasons. The frequency distribution histogram derived from field assessment data from Ecuador in 2017 was clearly skewed toward susceptibility, whilst data from Toluca in 2015 and 2016 seasons were also skewed toward susceptibility but were a closer fit to a normal distribution, respectively (Figure 2A).

#### Inheritance of Resistance to Ph and Pgt

In response to *Ph* pathotype 5457 P+, Pompadour gave an intermediate response of “1+C” at the seedling stage, and was highly resistant in the field (1R). In contrast, the B-19 parent was highly susceptible (“3+” and 90S, respectively) (Figure 1B). Barley leaf rust resistance in the RIL population segregated bimodally (1R : 1S) and was simply inherited at both the seedling (χ² = 0.013 at 1 d.f.; p = 0.909) and adult plant stages (2016 and 2018) (Figures 2B,C), with the segregation...
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FIGURE 2 | Frequency distribution of disease severity of 98 Pompadour/Biosaline-19 RILs at the adult plant stage in response to (A) 
Puccinia striiformis f. sp. hordei in Toluca Mexico in 2015 (large checker board) and in 2016 (black), and in Ecuador in 2017 (downward diagonal) using a modified Cobb scale; and (B) 
P. hordei in the field at Cobbitty, Australia in 2016 using a modified Cobb scale, and (C) two different experimental replicated in 2018 using a 1-9 McNeal scale, respectively. Disease severity values for parents are indicated by arrows.

pattern best fitting monogenic inheritance which was highly correlated to the presence of the Rph20 marker bPb-0837. In contrast to the phenotypic response to both leaf rust and stripe rust, both parental genotypes were resistant (Pompadour IT = “0;” and B-19 IT “;12C”) to Pgt race 98-1,2,3,5,6 at the seedling stage in the greenhouse. Pompadour was highly resistant and B-19 showed an intermediate response as shown in Figure 1C. The segregation for resistance in the P/B-19 mapping population best fitted a two-gene model ($\chi^2 = 0.974$ at 1 d.f.; $p = 0.324$), suggesting the involvement of resistance from both parents.

Genetic Map Construction

A total of 18,062 DArT-Seq markers showing polymorphism in the P/B-19 RIL population were considered for genetic map construction. Linkage map construction was performed using A5map LOD8 following marker curation. Markers that were redundant, had significant segregation distortion ($p < 0.01$) and/or low call rate (<70%), were removed as were duplicated samples, or those with missing data (>70%) (Supplementary Table S1). Following quality filtering, the final map consisted of 1,596 codominant SNP and 7,014 dominant DArT-Seq markers that spanned 5,957.6 cM across seven linkage groups corresponding to chromosomes 1H-7H (Supplementary Table S1). Approximately 50% of the markers could not be unambiguously assigned to a physical position in the 2017 Morex reference genome assembly; in 123 cases this was due to the presence of >2 map matches presumably due to paralogous gene families. In contrast, 4,220 markers were mapped to unique physical positions in the Morex genome. However, on numerous occasions the position determined in the P/B-19 linkage map did not correspond to that of the Morex reference. Numerous linkage blocks were identified during map construction, evidenced by the abundance of redundant markers. Nevertheless 4,220 markers could be mapped to unique physical positions in the Morex genome. Numerous regions in the linkage groups contained multiple redundant markers. Twenty-three such regions had >50 co-segregating markers including a region on chromosome 3H that had 233 redundant markers (Supplementary Table S1). This could be indicative of regions of the genome with repressed recombination or possibly an introgression from a wild Hordeum spp.

QTL Analysis for Rust Resistance

The high-density linkage map was used to precisely map QTL for rust resistance segregating in the P/B-19 mapping population. A total of 10 QTLs (exceeding the LOD threshold of 3) were mapped with distinct chromosomal locations for resistance to the four different rust diseases (leaf, stem, and stripe rust caused by both barley and barley grass-adapted formae speciales of P. striiformis) (Table 1). The LOD scores ranged from 3.11 to 17.45 for all QTLs and the percentage of variation explained by individual QTLs ranged from 11 to 36%, but was mostly lower than 20%, indicating that the mapped rust resistance at the seedling and adult plant stages were due to genes with both small and large effect (Table 1 and Figure 3).

For BYR resistance using the PshWUR isolate, the IT data from both seedling greenhouse experiments were highly correlated ($r = 0.8$) and distinct phenotypic classes (of resistant and susceptible) were easily distinguished and, therefore qualitative
assessments were taken from all RILs for the purpose of QTL mapping. We identified two consistent QTLs for resistance at the seedling stage on chromosomes 1H and 4H, both contributed by Pompadour (Table 1 and Figure 3). For both experiments, the mapping resulted in identical LOD profiles (Supplementary Figure S1A). The infection level of the P/B-19 RILs to BYR at the adult plant stage at field sites in both Mexico and Ecuador was assessed quantitatively and the segregation was continuous but skewed toward susceptibility. In 2016 at Toluca, Mexico, the same 1H QTL was identified as in the two greenhouse seedling experiments using both PshMex-1 and PshWUR isolates, respectively, while in 2015 (Toluca, Mexico) a signal with LOD 2.8. This QTL was not detected in 2017 from the Ecuador-derived dataset (Figure 3 and Supplementary Figure S1A). Consistent QTL on chromosome 2H were identified across all field environments, but was not identified in the greenhouse in response to the PshWUR isolate.

We also phenotyped the P/B-19 RIL population and mapped QTL for resistance to both leaf and stem rust. For leaf rust, we phenotyped the population in both the greenhouse under controlled conditions and in the field at Plant Breeding Institute Cobbity over three successive seasons at the adult plant stage using the same Ph pathotype (5457 P. hordei). We identified three QTLs for resistance on chromosomes 1H, 3H, and 5H at the seedling stage based on two highly correlated (r = 0.9) experimental replicates. From the field data, in all instances, we mapped QTL on the short arm of chromosome 5H corresponding to the position of Rph20, while an additional QTLs were also mapped only for the 2016 data on the short arm of chromosome 2H. The major-effect leaf rust QTLs identified on chromosome 5H are all in close proximity or co-locate with the bPb-0837 marker that is associated with the Rph20 resistance. The 1H leaf rust QTL was only effective at the seedling stage and was located near the centromere (Figure 3, Table 1, and Supplementary Figure S1B). For stem rust, two minor effect QTL (LOD score <5) for seedling resistance mapped on chromosomes 3H and 7H based on two experimental replicates (r = 0.7) (Figure 3, Supplementary Table S1, and Supplementary Figure S1C).

Co-location of QTLs against multiple rust pathogen species was identified on four different chromosomes, viz. 1H, 3H, 5H, and 7H. We remapped the phenotypic data reported in Haghdoust et al. (2018) for barley grass stripe rust resistance at the seedling stage using the genetic map created in the present study (Figure 3). The same QTL on chromosomes 1H, 3H, 5H, and 7H identified in Haghdoust et al. (2018) were also identified using the high-density genetic map. The chromosome 1H QTL co-located with the broadly effective QTL identified for BYR resistance, while the 3H and 7H QTLs co-located with both stem rust resistance loci mapped in this study (Figure 3).

**DISCUSSION**

The accuracy and speed taken to characterize disease resistance genes in plants has increased due to advancements in genomics, especially through cheaper sequencing technologies. GBS-derived complexity reduction methodologies such as DArT-Seq generate thousands of polymorphic markers that can be used to expand the size of linkage groups. We previously mapped both leaf rust (Dracatos et al., 2014; Kavanagh et al., 2017) and stripe rust (Dracatos P.M. et al., 2015) resistance in the

| Trait | Location | Stage | Pathogen isolate | Chromosome | Peak LOD | Position (cM) |
|-------|----------|-------|------------------|------------|----------|---------------|
| qField_Mex2015_3.30 | Toluca, Mexico_2015 | Adult | Psh Mex-1 | 1H | 3.3 | 37.55 |
| qField_Mex2015_5.13 | Toluca, Mexico_2015 | Adult | Psh Mex-1 | 2H | 5.13 | 327.06 |
| qField_Mex2016_5.30 | Toluca, Mexico_2016 | Adult | Psh Mex-1 | 2H | 5.3 | 338.59 |
| qField_Ecuador2017_3.95 | Ecuador_2017 | Adult | Field infection (mixed) | 2H | 3.95 | 335.59 |
| qGH_WUR_rep1_3.26 | WUR_greenhouse | Seedling | PshWUR | 1H | 3.26 | 42.37 |
| qGH_WUR_rep1_3.86 | WUR_greenhouse | Seedling | PshWUR | 4H | 3.86 | 329.64 |
| qGH_WUR_rep2_5.78 | WUR_greenhouse | Seedling | PshWUR | 1H | 5.78 | 43.37 |
| qGH_WUR_rep2_3.28 | WUR_greenhouse | Seedling | PshWUR | 4H | 3.28 | 339.64 |
| qGH_PBIC_5.29 | PBIC_greenhouse | Seedling | Papb isolate 9865566 | 1H | 5.29 | 35.55 |
| qGH_PBIC_3.14 | PBIC_greenhouse | Seedling | Psh isolate 9865566 | 3H | 3.14 | 472.8 |
| qGH_PBIC_4.26 | PBIC_greenhouse | Seedling | Psh isolate 9865566 | 5H | 4.26 | 331.07 |
| qGH_PBIC_4.51 | PBIC_greenhouse | Seedling | Psh isolate 9865566 | 7H | 4.51 | 967.33 |
| qGH_PBIC_3.91 | PBIC_greenhouse | Seedling | P. hordei 5457 P+ | 1H | 3.91 | 272.71 |
| qGH_PBIC_3.86 | PBIC_greenhouse | Seedling | P. hordei 5457 P+ | 3H | 3.86 | 761 |
| qGH_PBIC_9.67 | PBIC_greenhouse | Seedling | P. hordei 5457 P+ | 9H | 9.67 | 79.04 |
| qField_PBIC_2018_rep1_11.79 | PBIC_Australia_2018_rep1 | Adult | P. hordei 5457 P+ | 5H | 11.79 | 49.39 |
| qField_PBIC_2016_3.14 | PBIC_Australia_2016 | Adult | P. hordei 5457 P+ | 2H | 3.14 | 0.01 |
| qField_PBIC_2016_3.99 | PBIC_Australia_2016 | Adult | P. hordei 5457 P+ | 5H | 3.99 | 48.9 |
| qField_PBIC_2018_rep2_17.45 | PBIC_Australia_2018_rep2 | Adult | Pgt 98-1,2,4,5,6 | 3H | 3.11 | 443.79 |
| qGH_PBIC_3.11 | PBIC_greenhouse | Seedling | Pgt 98-1,2,4,5,6 | 7H | 4.6 | 933.03 |
Quantitative trait loci (QTL) for resistance on chromosomes 1H, 2H, 3H, 4H, 5H, and 7H in the Pompadour/Biosaline-19 RIL population to Puccinia striformis f. sp. hordei (red), P. striformis f. sp. pseudo-hordei (blue), P. graminis f. sp. tritici (green), and P. hordei (black). The name on QTL bars has three components: the provisional name of the gene at the QTL, location/isolate and the logarithm of odds (LOD) value recorded for the QTL.
barley and developed a high-density linkage map in wheat to map both components of complementary stripe rust resistance (Dracatos et al., 2016) using the DArT-Seq marker platform. To enable precise mapping of QTLs for resistance to stripe rust, leaf rust and stem rust derived from the European malting barley cv. Pompadour, we constructed a linkage map for P/B-19 RIL population reported in Haghdoust et al. (2018) using 8,610 polymorphic DArT-Seq and SNP markers spanning > 5,000 CM.

The ability of cereal rust pathogen spores to migrate both within and between continents and hence adapt to new environmental conditions and host genotypes continue to provide challenges to cereal production (Brown and Hovmøller, 2002; Ali et al., 2014). In cases where phenotypic analysis has to be performed offshore at international field and greenhouses because a pathogen is not present, as is the case for Psh in Australia, using a genetic map with high marker density to resolve the location of the R gene/QTL in the first instance can enhance the efficiency of subsequent pre-emptive breeding efforts. Phenotyping and genetic analysis for BYR resistance in the greenhouse determined that Pompadour likely carries two complementary resistance genes effective at the seedling stage in response to inoculations with PshWUR. CIM analysis identified the presence of two QTLs for stripe rust resistance at the seedling stage on chromosome 1H and 4H in both greenhouse experiments. We included differentials Emir and Agio in our tests and determined both shared a similar disease response to PshWUR as the Pompadour parent. Pedigree analysis of Pompadour parent revealed that it was derived from Emir through Agio, suggesting that the observed seedling resistance is likely rpsEm1 and rpsEm2 genes (Figure 4). Further intercrossing and subsequent rust testing is required to truly confirm the complementary gene hypothesis, following the procedures used in wheat by Dracatos et al. (2016) to map the complementary stripe rust resistance genes Yr73 and Yr74. We also assessed the P/B-19 mapping population at high altitude cooler regions in Ecuador and Mexico under field conditions over three successive seasons. The same 1H QTL was also mildly effective in Toluca, Mexico in 2016 (not significant in 2015, LOD = 2.78) at the adult plant stage in the field, suggesting it may be interactive with the 2H QTL. In contrast to the PshWUR isolate we used for mapping of seedling resistance, the isolate used in Toluca (Mex-1) was virulent with respect to the Emir (rpsEm1 and rpsEm2) resistance. Resistance to PshWUR was previously mapped by Niks et al. (2015) on chromosome 4H in the Vada x L94 mapping population and was contributed by the L94 parent and believed to be derived from the Ethiopian accession Grannelose Zweizeilige. We did not identify the 4H QTL from the data from Mexico or Ecuador, suggesting it is either ineffective to the Mex-1 isolate and Ecuadorian Psh population, or only effective at the seedling stage. A similar result was previously observed for the Lr12/Lr31 wheat leaf rust complementary gene system where only Lr12 conferred resistance in adult plants (Singh et al., 1999).

FIGURE 4 | Pedigree analysis of the Pompadour parent, accessions in orange were resistant to Psh isolates PshWUR and PshMex-1 used in this study.
stripe rust resistance providing broad-spectrum resistance under diverse environments.

Numerous barley accessions have been reported to carry the Rph20 resistance, however, most are highly susceptible as seedlings using oil-based inoculation methods (Golegaonkar et al., 2010). Two previous studies using different P. hordei isolates and inoculation methods determined that (Rphq4 – Rph20) was expressed from the 3rd leaf stage to maturity (Wang et al., 2010; Singh et al., 2013). In contrast, Wang et al. (2010) determined that a second QTL for partial resistance, Rphq2 on chromosome 2HL, prolongs latent period at the seedling stage but has almost no effect on disease resistance in adult plants. We identified three QTLs for leaf rust resistance at the seedling stage, on chromosomes 1H, 3H, and 5H. The presence of the bPb-0837 marker allele was strongly correlated with resistance in the P/B-19 RIL population, suggesting that the 5HS QTL detected is likely Rph20; however, the 1H QTL appears to be an uncharacterized QTL only expressed at the seedling stage. Our data suggests that in the Pompadour background, Rph20 is expressed earlier than previously determined. Relative to the Australian cultivar Flagship, numerous European varieties including Pompadour are more resistant (near immune) to leaf rust under field conditions under high inoculum pressure (Golegaonkar et al., 2010). Whether this is due to additional QTL for partial resistance (as observed in Vada), allelic variation at the Rph20 locus or variation in the pathogen population is yet to be determined. Our field data suggest that despite two experiment-specific minor effect QTLs, Rph20 was consistently detected across all three replicates. Whether this indicates that the Pompadour Rph20 allele confers stronger resistance or that other genetic components could not be effectively phenotyped due to environmental variation or inoculum pressure is unknown. Once Rph20 is cloned, further studies of allelic variation and their phenotypic effect will be possible, and will be relevant to breeding programs and evolutionary studies.

Two cloned resistance genes (Rpg1 and rpg4/Rpg5) have proved the most important to date for durable protection to stem rust in barley (Steffenson et al., 2017). Although rpg4/Rpg5 is still widely effective to most global Pgt isolates, virulence for Rpg1 is now common, and in some countries such as Australia, virulence is thought to be fixed. In this study, we were successful in mapping two QTLs for resistance to stem rust at the seedling stage on chromosomes 3H and 7H from greenhouse, derived from each of experiments using the parents of the P/B-19 RIL populations. A very recent study mapping QTL for resistance to stem rust in barley mapped both Rpg2 and Rpg3 on chromosomes 2H and 5H, respectively (Case et al., 2018). The same study identified multiple co-locating QTL, and a single QTL on chromosomes 3H and 7H, respectively, in response to North American isolates of Pgt at the seedling stage (Case et al., 2018). Due to the lack of common marker types the relationship between these QTL is unknown. However, the 7H QTL identified in the present study appears to provide a high level of protection at the seedling stage and is potentially a valuable source of resistance for varietal improvement.

DArT-Seq is a high-throughput complexity reduction based GBS technology enhancing genetic capabilities including: high-density linkage map construction, GWAS, QTL mapping and genetic diversity analysis. In summary, we used DArT-Seq genotypic data to generate a high-density linkage map in the barley RIL population P/B-19 to precisely map resistance QTL to stripe, leaf, and stem rust pathogens. We determined Pompadour was a rich source of resistance to all three rust species. Further use of DArT-Seq marker sequence information is ongoing for marker development for the selection of triple rust resistance in barley. In the era of rapid gene cloning methodologies, future experiments to isolate the chromosome 2H and 4H QTLs (likely in a region of low recombination) should adopt a mutant sequencing approach as recently described for barley leaf rust resistance gene Rph1 (Dracatos et al., 2019) to search for candidate gene(s) within the defined intervals confirmed in this study.

AUTHOR CONTRIBUTIONS

PD, RP, and DS conceptualized the work and designed the project. PD wrote the manuscript with contributions from RH, which was read, edited and approved by all co-authors. PD and RH performed the genetic and QTL analysis. PD, RH, DS, JHE, RN, RS, and CB performed the phenotypic analysis. MH and KF performed the genetic map construction.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2019.00467/full#supplementary-material

**FIGURE S1 |** LOD trace files for (A) stripe [green and black = WUR greenhouse rep 1 and 2, dark blue, red, and black dash = Mexico and Ecuador field sites and light blue = P. striiformis f. sp. pseudo-hordei], (B) leaf [pink = greenhouse PBI Cobbytty, dark blue dash = Field_PBCobbytty_2018rep1, light blue = Field_PBCobbytty_2018rep2, black dashed = Field_PBCobbytty_2016] and (C) stem rust [red dashed = greenhouse PBI Cobbytty] resistance QTLs identified in the Pompadour × Biosaline-19 RIL population.

**TABLE S1 |** Summarised marker information for the high-density linkage map constructed for the Pompadour/Biosaline-19 RIL population.
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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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