



Adding Complexity to Complexity: Gene Family Evolution in Polyploids

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Comparative genomics of non-model organisms has resurrected whole genome duplication (WGD) from being viewed as a somewhat obscure process that happens in plants to a primary driver of eukaryotic diversification. The shadow of past ploidy increases has left a strong signature of duplicated genes organized into gene families, even in small genomes that have undergone effectively complete rediploidization. Nevertheless, despite continually advancing technologies and bioinformatics pipelines, resolving the fate of duplicate genes remains a substantial challenge. For example, many important recognition processes are driven not only by allelic expansion through retention of duplicates but also by diversification and copy number variation. This creates technical difficulties with assembly to reference genomes and accurate interpretation of homology. Thus, relatively little is known about the impacts of recent polyploidization and hybridization on the evolution of gene families under selective forces that maintain diversity, such as balancing selection. Here we use a complex of species and ploidy levels in the genus *Arabidopsis* (*A. lyrata* and *A. arenosa*) as a model to investigate the evolutionary dynamics of a large and complicated gene family known to be under strong balancing selection: the receptor-like kinases, which include the female component of genetically controlled self-incompatibility. Specifically, we question: (1) How does diversity of S-receptor kinase (SRK) alleles in tetraploids compare to that in their close diploid relatives? (2) Is there increased trans-specific polymorphism (i.e., sharing of alleles that transcend speciation, characteristic of balancing selection) in tetraploids compared to diploids due to the higher number of copies they carry? (3) Do these highly variable loci show evidence of introgression among extant species/ploidy levels within or outside known zones of hybridization? (4) Is there evidence for copy number variation among paralogs? We use this example to highlight specific issues to consider when interpreting gene family evolution, particularly in relation to polyploids but also more generally in diploids. We conclude with recommendations for strategies to address the challenges of resolving such complex loci in the future, using advances in deep sequencing approaches.

Keywords: polyploidy, gene family evolution, self-incompatibility, copy number variation, trans-specific polymorphism, balancing selection, introgression

INTRODUCTION

Background and Aims

The sequencing of the human genome in 2001 (Lander et al., 2001) promised to revolutionize modern medicine and lead to a new era in understanding the complexity of genetic control of complex phenotypes. While this has certainly been true, it is really the comparative genomics of non-model organisms that has led to a complete revolution in understanding (e.g., Seeb et al., 2011; da Fonseca et al., 2016). One unexpected finding was that whole genome duplication (WGD) has been an important process contributing to the genomic history of all eukaryotes, including those with relatively small genomes, such as the yeast *Saccharomyces cerevisiae* (Wolfe and Shields, 1997) and the model plant *Arabidopsis thaliana* (Blanc and Wolfe, 2004). Although Susumu Ohno in the late 1960s had emphasized the central role of gene duplication in the evolutionary history of vertebrates (Ohno, 1970), it wasn't until after his death in 2000 that comparative genomic studies confirmed that fish had undergone multiple rounds of WGD (e.g., Meyer and Van de Peer, 2005), as he had predicted. He also had predicted that effective rediploidization following duplication was inevitable but that some duplicates would be retained to perform new or specialized functions, leaving a footprint of past duplications and organization of genes into gene families. His ideas about the fates of duplicate genes to include specialization of function (now known as “subfunctionalization”; Force et al., 1999) also have been resurrected and form the basis for understanding the history of complex genomes such as salmonids, which underwent an independent WGD after the last teleost specific duplication (Hermansen et al., 2016; Lien et al., 2016). Comparative studies of vertebrates have thus been critical for establishing polyploidization as a creative evolutionary force shaping the genomes of all eukaryotes (Van de Peer et al., 2017), as had long been recognized for plants (e.g., Soltis et al., 1992; Adams, 2007).

Nevertheless, despite recognition that duplicated genes are critical for understanding genome structure and function (Van de Peer et al., 2017), the practicalities of assembling duplicates in genomic resequencing studies, resolving orthology, and interpreting their potentially redundant effects on phenotypes remains a substantial challenge (da Fonseca et al., 2016). Retention of duplicate genes following genomic or tandem duplication is non-random (Adams, 2007) and is both constrained and promoted by achieving appropriate levels of expression (e.g., Gout and Lynch, 2015; Mattenberger et al., 2017; Rodrigo and Fares, 2018). The “gene balance” hypothesis, for example, predicts that loci involved in regulating levels of expression of integrated genetic pathways (such as transcription factors or members of signal transduction pathways) should show increased retention of duplicates to maintain coordinated function (Birchler and Veitia, 2010). Genes for which high expression is advantageous might be expected to retain expression in duplicated copies whereas divergence in patterns of expression could be advantageous for others. Genes that are retained in duplicate through one round of WGD also have been found to be preserved through later rounds (Seoighe and Gehring, 2004). Thus, not considering the role of gene copies

retained in duplicate could alter interpretation of regulatory processes associated with adaptation.

One type of adaptive process often associated with large and complex gene families is recognition of self vs. non-self, where high polymorphism is favored by continually changing selection pressures, and retention of duplicate copies could be beneficial for increasing allelic repertoire. For example, the “big bang” theory of the emergence of the adaptive immune systems in vertebrates invokes multiple rounds of WGD as the major source of this potential (Flajnik and Kasahara, 2010). Similarly, investigation of the genomic repertoire of pathogen-associated genes (R genes) in several crop plants through targeted sequence capture (Jupe et al., 2012, 2013; Giolai et al., 2016; Van Weymers et al., 2016) has revealed much more extensive gene families than was previously predicted based on whole genome resequencing studies. R genes have also been demonstrated to show signatures of adaptive introgression between closely related species of *Arabidopsis*, with extensive trans-specific sharing of alleles across species (Bechsgaard et al., 2017). An added complication for these types of gene families is that copy number can be variable even among individuals within a species (e.g., Mable et al., 2015), meaning that genome references will not always include the full complement of copies. Copy number variation has been linked to disease severity in humans (Beckmann et al., 2007; Wheeler et al., 2008) and adaptive processes in other organisms (Santenac et al., 2011; Zmienko et al., 2014; Duvaux et al., 2015; Hull et al., 2017) but methods that can reliably distinguish between lack of coverage and variation in presence of a particular gene copy are required to fully evaluate the evolutionary significance of presence/absence polymorphisms following gene duplication.

The high polymorphism expected for recognition genes means that they are prime candidates to be “lost” in genomic resequencing studies, even in diploids. For example, genes controlling sporophytically controlled self-incompatibility (SI) in plants have been found to be missing from resequencing assemblies because they are too divergent from the reference genome and so trawling in the unassembled reads is necessary to characterize these highly polymorphic genes (Mable et al., 2017). Both male and female components are members of large gene families that show extensive trans-specific polymorphism, with highly similar alleles shared across species and even genera but high divergence between functional specificities (Schierup et al., 1998; Paetsch et al., 2006; Castric and Vekemans, 2007; Busch et al., 2008; Guo et al., 2011; Tedder et al., 2011; Leducq et al., 2014). The gene controlling female specificity (S-receptor kinase, *SRK*) is part of a large family of receptor kinases, which evolved through a complex history of gene duplication and loss, followed by gene fission and fusion (Xing et al., 2013). Gene conversion between *SRK* and other members of the gene family is also thought to have contributed to expansion of functional allelic diversity (Prigoda et al., 2005; Guo et al., 2011). This creates additional challenges with interpreting which variants are parts of the functional locus regulating the SI response and which are functionally unlinked but show high sequence similarity. For sporophytic SI, the phenotype of the pollen is determined by the genotype of the diploid (or tetraploid) parent, so there can be dominance in both pollen and stigma.

Dominance is known to be complex, with non-linear interactions that can differ between pollen and stigma (Lewis, 1947; Stevens and Kay, 1989; Hatakeyama et al., 1998; Shiba et al., 2002; Mable et al., 2003; Llaurens et al., 2009; Schoen and Busch, 2009). Trans-specific polymorphism (i.e., sharing of alleles that transcends speciation) of *SRK* alleles has been well established for diploids (Charlesworth et al., 2006; Boggs et al., 2009; Castric et al., 2010), and is thought to be a key indicator of the action of balancing selection (Takahata, 1990). However, the strength of balancing selection on tetraploids has not been assessed specifically. Since tetraploid individuals can carry up to four different *SRK* alleles, there is potential for increased sharing across species, at least of recessive alleles. They can also carry multiple copies of recessive alleles (Mable et al., 2004), which could result in the maintenance of more variants within specificities than for diploids. While previous work has demonstrated that linkage and dominance works similarly in tetraploid compared to diploid *Arabidopsis lyrata* (Mable et al., 2004), the evolutionary dynamics of *S*-alleles in tetraploids has not been studied.

In addition, interpreting the fate of duplicate genes in polyploids is complicated by the fact that hybridization is often associated with WGD and so it can be difficult to disentangle the effects of combining and duplicating genomes on patterns of duplicate gene expression or dynamics of gene families (e.g., Evans, 2007; Guggisberg et al., 2009; Mable, 2013). Fortunately, rapid advances in sequencing technology and bioinformatic processing mean that the toolbox available to resolve such challenges continues to improve. Targeted sequence capture, for example, has been used effectively to investigate genomic changes in polyploids (Salmon et al., 2012; Gardiner et al., 2016; Krasileva et al., 2017). However, even with these advances in technology there are important issues to consider when resolving and interpreting evolutionary dynamics of gene families, particularly for systems in which recent polyploidization and hybridization could complicate accurate assembly into orthologs and subsequent genotyping within and between copies.

The purpose of this paper is to discuss these issues in the context of understanding the evolutionary dynamics of the *SRK* gene family in a species complex (*A. lyrata* and *A. arenosa*) that includes both diploids and tetraploids, with tetraploids showing extensive introgression in a hybrid zone in central Europe (Schmickl et al., 2010; Jørgensen et al., 2011; Schmickl and Koch, 2011; Hohmann et al., 2014; Muir et al., 2015; Novikova et al., 2016; Hohmann and Koch, 2017). In *A. arenosa*, tetraploids have been predicted to have arisen through autopolyploidisation (Arnold et al., 2015); secondary contact with *A. lyrata* during interglacial and postglacial range contractions and expansions has subsequently led to introgression between tetraploids in the two species. Our intent was to use investigation of *S*-receptor kinase evolution in this species complex as a model for understanding how balancing selection operates in polyploid genomes and to determine whether these highly polymorphic gene families could be useful indicators of hybridization and introgression. Specifically, our objectives were to question: (1) How does diversity of *SRK*-related alleles in tetraploids compare to that in their close diploid

relatives? (2) Is there increased trans-specific polymorphism of *SRK* alleles in tetraploids compared to diploids because of the increased number of copies they can carry? (3) Do these highly variable loci show evidence of introgression among extant species/ploidy levels within or outside known zones of hybridization? (4) Is there evidence for copy number variation among paralogs?

We use these questions to highlight the challenges for interpreting gene family evolution, particularly in polyploids, but also relevant to diploids. We conclude with recommendations for how some of these challenges might be overcome using deep sequencing approaches. We reiterate the recommendation from others (Salmon et al., 2012; Jupe et al., 2013; Gardiner et al., 2016; Van Weymers et al., 2016; Krasileva et al., 2017) that non-amplicon based targeted sequence capture (e.g., whole genome exon capture or targeting of particular gene families) is the most promising method for tackling the full complexity of gene family evolution in complex genomes but suggest cautionary strategies that should be considered when interpreting evolutionary patterns.

Notes on Terminology and Known Challenges Associated With the *SRK* Gene Family

A complication with understanding the evolution of complex gene families is distinguishing what is meant by an “allele.” For *SRK*, there can be sequence variation within “specificities,” which are *SRK* types that confer a specific SI phenotype (i.e., a protein expressed on the surface of the stigma that is recognized as self by the comparable protein expressed on the surface of the pollen grain). These specificities (which we will refer to as “alleles”) can be as divergent from one another as they are from other genes (which we will refer to as “loci”) in the same gene family. Moreover, phylogenetic clustering alone is not sufficient to predict which sequence variants represent *SRK* alleles because gene conversion with unlinked loci has resulted in higher similarity between paralogs than among *SRK* alleles (Prigoda et al., 2005). Diploid individuals should contain only two functional *SRK* alleles but could contain varying numbers of loci in the gene family that are not linked to the SI phenotype; since tetraploids can contain multiple copies of the same allele without altering the specificity or dominance (Mable et al., 2004), the number of *SRK* alleles expected in a polyploid cannot be predicted. Thus, assigning “sequence variants” to gene family loci or *SRK* alleles is even more complicated in polyploids than for diploids. *SRK* alleles have been grouped into four different dominance classes (A1, A2, A3, B; Prigoda et al., 2005). Polymorphisms within specificities/alleles (which we will refer to as “haplotypes”) are more apparent for recessive than dominant alleles because the former are expected to occur at higher frequency and show more sharing between populations (Bechsgaard et al., 2006; Castric and Vekemans, 2007; Castric et al., 2008, 2010; Stoeckel et al., 2008; Llaurens et al., 2009; Goubet et al., 2012). There is a single most recessive allele (S1, Class A1; Prigoda et al., 2005) that is found globally and in multiple species in the genus *Arabidopsis* (Mable et al.,

2003; Dart et al., 2004; Prigoda et al., 2005; Mable and Adam, 2007; Castric et al., 2010; Foxe et al., 2010). Alleles in Class B are recessive to all other classes except S1 but are more similar to unlinked loci (*Aly13-2* and *Aly13-7*) than to the other classes (Prigoda et al., 2005) and show more intra-allele polymorphisms than dominant alleles (Classes A2 and A3; Prigoda et al., 2005; Castric et al., 2010). The high trans-specific polymorphism also means that naming of alleles can be confusing because a variant found in certain species is often provided a specific number before discovering that it potentially represents the same specificity as an already named allele in another species (Castric et al., 2010). Thus, alleles are named with the species in which they were originally described as a prefix (e.g., *Aly* refers to *A. lyrata*, *Aha* refers to *A. halleri*, *Ath* refers to *A. thaliana*, *Aar* refers to *A. arenosa*). Finally, since the SI phenotype is determined by a combination of variants at the female *SRK* and male *SCR* genes, phenotypic specificities are labeled only “S#” (e.g., S1) for segregation analyses.

From our previous studies on the evolutionary dynamics of *SRK* alleles in diploids, we have already described challenges in generating robust data for interpreting these complex gene families in diploids, relevant for the sequencing strategies we apply here: (1) Primers designed to be general enough to recognize all *SRK* alleles also amplify the rest of the gene family, so a major challenge is assigning sequence variants to loci (Schierup et al., 2001; Charlesworth et al., 2003b; Mable et al., 2003, 2017; Mable and Adam, 2007). (2) This is complicated by the fact that, due to the extensive polymorphism in *SRK* and evidence that gene conversion has contributed to allelic repertoire, paralogs that are not linked to the SI phenotype can be more similar to “real” alleles than “real” alleles are to one another, so similarity can’t always be used to assign functionality (Schierup et al., 2001; Mable et al., 2003; Prigoda et al., 2005). (3) Amplicon-based approaches are inherently at risk of generating PCR recombinants between copies, making it difficult to distinguish errors from actual recombination, introgression in hybrids, or gene conversion between sequences. (4) It is also difficult to distinguish presence/absence of paralogs from amplification biases during PCR (Mable et al., 2017). (5) There is extensive length heterogeneity within and between members of the gene family, so it can be difficult to establish the positional homology necessary to interpret patterns of selection (Charlesworth et al., 2003a). (6) The highly polymorphic nature of *SRK* alleles means that they are sometimes too divergent from the reference genome to be assembled using standard filtering strategies; this means that these types of alleles might frequently be found in the unassembled reads for resequencing projects (Mable et al., 2017).

MATERIALS AND METHODS

Sampling and Overview of Methods

Samples were obtained from both diploid and tetraploid populations of *A. lyrata* and *A. arenosa* sampled from Central Europe (Table 1). Although current systematics suggests

separation of diploid and tetraploid *A. arenosa* into distinct species taxonomically (Koch et al., 2008), for simplicity, we will refer to both as *A. arenosa* here. We sampled individuals from 3–5 populations of each “type”: A2x refers to diploid *A. arenosa*, A4x to tetraploid *A. arenosa*, L2x to diploid *A. lyrata* and L4x to tetraploid *A. lyrata*. Tetraploid populations occurring in a hybrid zone between the two species (Schmickl, 2009; Schmickl and Koch, 2011; Hohmann et al., 2014; Muir et al., 2015; Novikova et al., 2016) were included to test for patterns of introgression. Diploids have not been found to hybridize (Jørgensen et al., 2011) and so were considered “pure” populations. To test patterns of linkage of sequence variants with the SI phenotype, we also included 104 individuals from crosses between *A. lyrata* tetraploid parents whose genotypes had been partially resolved by cloning and Sanger sequencing; we performed di-allele crosses within these families to establish SI phenotypes that could be compared to the 454 genotypes.

We used a combination of approaches to address the main research questions: (1) 454 pyrosequencing using degenerate primers (Supplementary Table 1) targeting the *SRK* gene family (Jørgensen et al., 2012) to characterize diversity and patterns of allele sharing in diploids and polyploids; (2) direct Sanger sequencing to investigate signatures of introgression in shared haplotypes and for segregation analyses to test linkage to the SI phenotype; (3) cloning and Sanger sequencing using degenerate primers (Supplementary Table 1) to obtain longer products than possible with 454 pyrosequencing to further characterize potentially new alleles; and (4) using data from a recent genomic resequencing study (Novikova et al., 2016) to search for the *SRK* gene family using novel assembly approaches, to test whether copy number variation and patterns of introgression can be mined using existing genomic data. We focused on variation in exon 1 (the S-domain) because it contains the sites used for recognition of self vs. non-self (Schierup et al., 2001; Charlesworth et al., 2003a). However, we also used the genome mining approach to determine whether we could pull out full-length sequences that include the functional kinase domain (exons 3–7).

While 454 pyrosequencing has largely been replaced by methods demonstrated to show higher accuracy such as Illumina (Schirmer et al., 2015, 2016; D’Amore et al., 2016), we use results from this study as a platform to highlight considerations for working with gene families that should apply across methods. We thus haven’t focused on attempting to resolve 454 specific problems but instead on general issues with clustering and assigning sequence variants to loci and designating allelic specificities for interpretation of gene family evolution. We include these as “challenges” in relation to the methods used to address each objective.

Detailed Methodology

Clustering and *SRK* Genotyping Strategies

To increase the probability of amplifying all variants of *SRK* present in the populations sampled, we used 454 pyrosequencing of pooled amplicons from four sets of degenerate primers but sharing a common reverse sequence, *SLGR* (Supplementary Table 1; Schierup et al., 2001). Detailed

TABLE 1 | Populations sampled, indicating the code (Pop Code) used to identify populations in our study, the population identifier (identity) from Schmickl (2009), site description, ploidy, species, country of origin, GPS coordinates (latitude and longitude), and whether the population is in the known hybrid zone in Austria, as well as sample sizes for the 454 pyrosequencing (N 454) and targeted amplicon sequencing of *SRK01* (N *SRK01*).

Pop Code	Site Description	Identity	Species	Ploidy	Country	Latitude	Longitude	Hybrid zone	N 454	N <i>SRK01</i>
A2_SVK1	Vsoky Tatry	131R	<i>A. arenosa</i>	Diploid	Slovakia	49.2325	20.1980	No	28	28
A2_SVK2	Velkra Fatra	915141	<i>A. arenosa</i>	Diploid	Slovakia	48.8242	19.0233	No	25	0 ^b
A2_SVK3	Nizke Tatry	915140	<i>A. arenosa</i>	Diploid	Slovakia	48.8843	20.2485	No	25	16
A4_AUT1	Kernhof	915142	<i>A. arenosa</i>	Tetraploid	Austria	47.8162	15.5435	Yes	25	18
A4_AUT2	Achleichten, Wachau	123R	<i>A. arenosa</i>	Tetraploid	Austria	48.4064	15.4728	Yes	26	16
A4_AUT3	Kamptal	3R	<i>A. arenosa</i>	Tetraploid	Austria	48.5306	15.6915	Yes	15	13
A4_AUT4	Scheibenbach, Wachau	89R	<i>A. arenosa</i>	Tetraploid	Austria	48.4137	15.5200	Yes	14	11
A4_GER	Wental	20R	<i>A. arenosa</i>	Tetraploid	Germany	48.7335	10.0193	No	7	0 ^b
L2_AUT1	Pernitz-Pottenstein	112R	<i>A. lyrata</i>	Diploid	Austria	47.9275	15.9861	No	25	23
L2_AUT2	Vöslauer Hütte	96R	<i>A. lyrata</i>	Diploid	Austria	47.9803	16.1650	No	25	9
L2_CZE	Oslavany, Brno	915143	<i>A. lyrata</i>	Diploid	Czech R.	49.1219	16.3244	No	9	8
L2_GER	Veldensteiner Forst	915145	<i>A. lyrata</i>	Diploid	Germany	49.6453	11.4508	No	17	17
L4_AUT1	Dürnstein, Wachau	13R	<i>A. lyrata</i>	Tetraploid	Austria	48.3970	15.5345	Yes	25	7
L4_AUT2	Mödling	915144	<i>A. lyrata</i>	Tetraploid	Austria	48.0768	16.2698	Yes	25	18
L4_AUT3	Bachamsdorf, Wachau	50R	<i>A. lyrata</i>	Tetraploid	Austria	48.3722	15.4542	Yes	25	22
L4_AUT4	Lilienfeld	116R	<i>A. lyrata</i>	Tetraploid	Austria	47.9981	15.5736	No	21	10
L4_AUT5	Rauheneck Ruin	na ^a	<i>A. lyrata</i>	Tetraploid	Austria	48.0021	16.2309	No	19	19
L4_AUT2 x L4_AUT5	Crosses		<i>A. lyrata</i>	Tetraploid	Austria			No	104	99
Total									460	334

Crosses performed between individuals sampled from Mödling and Rauheneck Ruin near Baden were used to test segregation of genotypes resolved using 454 and *SI* phenotypes.

^aNot included in Schmickl (2009) but collected from Rauheneck Ruin, near Baden.

^bInsufficient DNA remained after the 454 sequencing to screen for *AlySRK01*.

methods for the 454 analyses are described in Jørgensen et al. (2012), including estimation of error rates and the use of segregation within known families to test the reliability of genotyping. The initial paper described the strategies used for clustering reads into contigs and filtering to reduce errors. We recommended that optimal clustering was obtained with a 90% sequence similarity criterion and excluding sequences present at a frequency of <7% of the total reads for an individual; these conclusions were based on a subset of the original data that included repeated runs involving the same individuals. We also recommended that clustering should be conducted after reads were trimmed to 200 bp from the “common primer” end (*SLGR* in this case).

Although the crosses between tetraploid *A. lyrata* individuals confirmed presence of the expected *SRK* alleles known to be present in the parents, they also indicated some inaccuracy in allele calls in relation to barcodes; a number of alleles that were not in the parents were assigned to individuals from the crosses, sometimes at high read numbers (see Jørgensen et al., 2012). We concluded that this was due to tag switching between barcodes, as had been suggested from other studies (van Orsouw et al., 2007; Carlsen et al., 2012). Blank lanes (negative controls) also sometimes contained sequences matching known *SRK* alleles, again often at high read numbers. We thus modified our filtering and clustering strategies in the analysis of the full dataset.

Reads were initially assembled into contigs based on clustering to sequences from a reference database of known *SRK* alleles and known members of the gene family that have been characterized in other studies and from our unpublished data from Sanger sequencing. A second iteration then used newly sequenced reads as seeds for clustering, in order to identify putatively new alleles (generating “read-only” contigs). BLAST analyses of “read only” contigs indicated that some known alleles (both *SRK* and paralogs) had been fragmented into multiple contigs. In such cases, contigs for a particular allele were combined, sequences sorted by barcode, and read numbers counted for each individual that contained a particular sequence type. Remaining “read only” contigs that did not show at least 80% similarity to *S*-related kinases from Genbank were not considered further. Final contigs were then sorted into putative “types”: known *SRK* alleles, putatively new *SRK*-like variants, or known paralogs. Contigs assigned to *SRK* alleles whose dominance had been established previously (Prigoda et al., 2005; Goubet et al., 2012) were further sorted into the following classes: (1) A1, consisting of a single most recessive allelic specificity that has been found globally in *Arabidopsis* species (*SRK01*); (2) A2, dominant to all other classes; (3) A3, recessive only to class A2; and (4) B, recessive to all except A1 and showing high similarity to unlinked loci (*Aly13-2* and *Aly13-7*). Contigs were also inspected for clustering of more than one named *SRK* allele from the database.

The next step was to subdivide variants within contigs into individual haplotypes, in order to test patterns of trans-specific polymorphism and to assess evidence for introgression between species. In our pilot study (Jørgensen et al., 2012) we recommended that only sequence variants present in at least 7% of the reads for an individual should be “counted” as true variants. However, in the full analysis, inspection of the contigs associated with particular alleles revealed very uneven read numbers both between individuals (ranging from a minimum of a single read to a maximum of 1,126 reads in the 465 individuals screened; average 344 ± 156) and across loci (i.e., *SRK* alleles and paralogs) within individuals. Low read numbers of particular alleles were also not directly proportional to the overall read numbers in the individual. The strict 7% threshold would have excluded some alleles that amplified in multiple individuals but were only present at low read numbers within individuals. A striking example was *SRK01*: it was fragmented across multiple contigs but when reassembled, it tended to be found at very low read numbers within individuals but was found across a wide range of individuals and showed population- and species-specific variants, as expected for a recessive allele (Billiard et al., 2007; Goubet et al., 2012). Many individuals showed <20 reads but the individuals that showed high read numbers (>100) tended not to show amplification of any other alleles, suggesting competition in the PCR when other alleles were present.

For haplotype calling, we thus also considered genotype calls at thresholds of at least 4% of reads and between 0 and 4% of reads. A problem with assessing such optimization strategies when including tetraploids is that there is not a robust basis for excluding individuals based on numbers of expected haplotypes. Although we could use diploids to determine thresholds of read numbers that minimized calling of more than two *SRK* alleles per individual and predicting homozygosity only for recessive alleles, this was confounded by the difficulties of predicting linkage of newly identified alleles (Charlesworth et al., 2003b; Prigoda et al., 2005). Tetraploids are expected to have up to four copies of *SRK* per individual but they can also contain multiple copies of recessive alleles (Mable et al., 2004), precluding extrapolating “confidence thresholds” based on diploids. We thus decided on a conservative threshold of at least 20 reads for a given haplotype to make relative comparisons among populations and species in the frequency of presence of particular variants. For reconstruction of evolutionary relationships among alleles, haplotypes present in <20 reads in a single individual and individuals with <200 total reads were excluded.

Statistical Analyses

To investigate whether there were differences in sequencing quality, detection biases, or real differences in frequency of sequence variants found we used generalized linear models to test whether the variation was significantly explained by ploidy, species or their interaction. Since multiple 454 runs were used for genotyping, we included barcoding tag number and lane as random effects, to account for any variation they explained. Analyses were conducted using JMP version 10.0 (SAS Institute, Incorporated).

Reconstructing Evolutionary Relationships Among Alleles

To establish phylogenetic relationships of newly identified alleles and to predict their dominance, we aligned the 454 sequences to the reference set (**Supplementary Data Sheet 1**) and reconstructed phylogenetic trees, using MEGA 7.0 (Kumar et al., 2016). We extracted consensus sequences for each haplotype of the *SRK*-like alleles identified and initially performed multiple alignments using the online version of Clustal Omega (Sievers et al., 2011) and then optimized by eye to establish positional homology and to set the correct reading frame to minimize stop codons, using Se-al version 2.0 (Rambaut, 1996) and McClade version 4.0 (Maddison and Maddison, 2000). To assess patterns of trans-specific polymorphism, if there was an exact match of a sequence to the reference database used for clustering, we named the haplotype “REF_HAP1” but if there was no exact match we retained the database allele (just named “REF”). We also added homologs from *A. lyrata*, *A. arenosa*, *A. halleri* and *A. thaliana* from Genbank for each specificity identified among the 454 samples (e.g., *AHASRK04* and *ATH*-haplogroup A have been identified as homologs of *AlySRK37*; Bechsgaard et al., 2006). As implemented in MEGA, the best fitting substitution model was identified using ModelTest and then Maximum Likelihood was used to cluster sequences, using 1,000 bootstrap replicates. Due to the reticulate nature of evolution in this gene family, a strictly bifurcating evolutionary history is not expected but a tree-like representation is useful for identifying clusters of similar sequences. In previous studies, we have found that phylogenetic clustering is informative about dominance for Class A3 and B alleles but that Class A2 are paraphyletic based on alignments of approximately 900 bp of sequences in exon 1 of *SRK* (Prigoda et al., 2005). We thus used phylogenetic clustering to predict dominance of new specificities identified or known specificities for which dominance had not been established. We calculated genetic distances within and between dominance classes using both the best fitting substitution model and raw % similarity, using MEGA. We then mapped relative frequency of each haplotype in the four types of populations onto the tree, using Evolveview in the Evolgenius package (He et al., 2016).

Testing the Accuracy of 454 Genotyping Using Segregation Analyses

We used the 454 pyrosequencing to genotype *SRK* from 11 families raised from crosses between tetraploid *A. lyrata* individuals whose grandparents had at least partially resolved *SRK* genotypes, in order to test segregation of alleles and as an additional test of reliability of the clustering thresholds set. Given the low read numbers found for *SRK01*, we established genotypes by a combination of allele-specific Sanger sequencing for this allele with the 454 sequencing for other alleles to compare segregation of alleles within families and to aid in excluding spurious allele calls. For a subset of these crosses, we performed controlled pollinations among all pairwise combinations of individuals, in order to test linkage of the variants identified to

the SI phenotype and to predict dominance relationships (as in Mable et al., 2004).

Direct Sanger Sequencing of *SRK01*

To complement the 454 sequencing, we used targeted direct Sanger sequencing to resolve *SRK01* genotypes to be able to investigate signatures of introgression of this recessive allele. We screened all individuals raised from the crosses between tetraploid *A. lyrata* individuals to aid in segregation analyses and a subset of individuals from the population survey to confirm haplotype calls and obtain more accurate frequencies of variants within and between individuals (Table 1).

We amplified products using an allele-specific primer (qtALS*SRK01*F: TCCTACATCATCGCAG) with the general reverse primer (SLGR: ATCTGACATAAAGATCTTGACC) that had been used for 454 sequencing. The 20 μ L PCR reactions (using reagents from Invitrogen, Inc., Paisley, UK) consisted of 1 μ L template, 2 μ L 10x PCR buffer (Invitrogen Incorporated, Paisley, UK), 2 μ L 10 mM dNTPs, 1 μ L 50 mM MgCl₂x, 0.2 μ L 10 μ M of each primer, and 0.2 μ L *Taq* polymerase. The PCRs were run in MJ research thermocyclers using the following program: initial denaturing phase of 3 min at 94°C, 1 min annealing at 54°C, 2 min extension at 72°C; followed by 34 cycles of 30 s at 94°C, 30 s at 54°C, 2 min at 72°C; and a final extension step of 6 min at 72°C.

Individuals that showed amplification of products of the expected size (~500 bp) were sent for sequencing to The GenePool in Edinburgh, using the reverse primer SLGR. Chromatograms were checked for base-calling errors using Sequencher 4.7 (Gene Codes Corporation, Ann Arbor, MI) and BLAST was used to confirm sequence identity.

Sequences were aligned using Sequencher, version 4.7 and heterozygous positions were recorded using IUPAC (International Union of Pure and Applied Chemistry) ambiguity codes. The phase of heterozygous positions was resolved by matching to variants found in the 454 sequencing and to homozygous sequences found in the Sanger sequencing. Genotypes predicted based on this process were then aligned to the specific 454 sequences for each individual. Species-specific variants were identified in diploids based on private haplotypes for the two species. We used the datamonkey server (www.datamonkey.org; Delpont et al., 2010), which implements statistical tests associated with the programme HyPhy (Pond et al., 2005), to test for evidence of recombination using GARD (Genetic Algorithm for Recombination Detection; Pond et al., 2006). In addition, we manually inspected alignments for evidence of potential breakpoints and in such cases, aligned each "section" independently to the other haplotypes identified for a particular specificity. Where a putatively recombinant type showed similarity to two or more species-specific haplotypes in different regions of the sequence, they were classified as potentially introgressed. A minimum spanning network (Bandelt et al., 1999) was drawn using PopArt (Leigh and Bryant, 2015) to resolve the relationships among the *SRK01* haplotypes.

Cloning and Sanger Sequencing of Longer *SRK* Alleles

As the 454 sequences were too short to be informative for future population genetics analyses and tests for selection, we used degenerate primers (Supplementary Table 1) to amplify longer products from tetraploid *A. lyrata* and *A. arenosa* sampled from the hybrid zone in the Wachau region of Austria (~600 bp, also described in Ruiz-Duarte, 2012). We then used these products as seeds for the genome mining (see section Mining *SRK* Alleles From Genome Resequencing Data) to determine whether we could determine the genomic location of the "new" alleles found, as an indication of linkage to the S-locus.

Genomic DNA was extracted from three to four leaves from plants of tetraploid *A. lyrata* and *A. arenosa* individuals using a modified CTAB protocol (Doyle and Doyle, 1987). Degenerate primers known to amplify a number of different gene family copies and *SRK* alleles (Schierup et al., 2001) in *A. lyrata* and *A. halleri* (Forward: 13SeqF1, 5'-ccgacggtaacctgtcatcctc-3' and Reverse: SLGR, 5'-atctgacataaagatcttgacc-3') were used (Charlesworth et al., 2000). Genomic DNA was mixed with a pair of primers, 10 μ mol each, 4 μ L of 5x buffer (ready-made), 50 mM MgCl₂, 0.4 μ L of 10 mM dNTP mixtures, 0.1 μ L *Taq* DNA Polymerase (Mango *Taq*, Bioline). PCR amplification conditions were as follows: denaturation at 94°C for 2 min followed by 34 cycles of 94°C for 30 s, 50°C for 30 s, and 72°C for 30 s; a final extension at 72°C for 5 min.

PCR products were cloned into pGEM®-T Vector Systems (Promega Inc.). Colony PCR (20–30 colonies per individual) was conducted to test for inserts using SP6 and T7 primers, followed by *AluI* digestion to identify clones carrying different putative *SRK* alleles. To avoid errors that might occur during PCR amplification and sequencing, a minimum of three independent clones with the same restriction profile were sequenced at the GATC BIOTECH facility. SeqMan software (DNASTAR, Inc) was used to clean and create consensus sequences.

We created separate alignments for each allele that was found both in the 454 and the Sanger sequencing by aligning the new sequences to references from Genbank and to the 454 sequences, in order to confirm shared specificity (Supplementary Data Sheet 2).

Mining *SRK* Alleles From Genome Resequencing Data

The 454 pyrosequencing data was not appropriate for determining presence and absence of paralogs because of: (1) the difficulty of distinguishing gene copies from new alleles at the *SRK* locus; and (2) amplification biases that made it difficult to set thresholds for reliability. Several known paralogs (*Aly8*, *Aly9*, *Aly13-2/13-7*) were expected to amplify with the primer set used. Polymorphic regions like the S-locus are known to be difficult to assemble in genome resequencing studies due to divergence from the reference genome (Mable et al., 2017) but we tested whether *de novo* assemblies from a genome resequencing study (Novikova et al., 2016) could be used to assess copy number of the *SRK*-related kinase gene family. We also attempted to pull out full-length sequences that spanned the S-domain (exon

1), transmembrane (exon 2) and kinase domains (exons 3-7) (Charlesworth et al., 2003a).

There are currently 28 fully resequenced genomes available from diploid and tetraploid *A. lyrata* and *A. arenosa*, from which we selected three or four individuals from each species and ploidy level to test whether we could obtain useful information on copy number and complete gene sequences. We used our paired end read data (Genbank SRR2040821, SRR2040822, SRR2040825, SRS945917, SRS1256176, SRS1256175, SRR2020827, SRR2040828, SRR2040829, SRR2040830, SRR2040791, SRR3111440, SRR3111441) and trimmed the reads for adapter contamination using cutadapt (Martin, 2011) and the respective adapter sequences. To obtain *SRK* alleles from these data we attempted two different approaches: mapping based and *de novo* assembly, on average we used ~110 million paired end reads for the tetraploid accessions and ~60 million reads for the diploid accessions corresponding to an average coverage of 20x.

In the initial mapping strategy we used as reference the S-locus region of the *SRK* locus on scaffold 7 of the MN74 reference genome (which was originally sampled from a North American outcrossing populations and has the S13 allele of the genes AL7G32720 = *SCR*, AL7G32730 = *SRK*, AL7G32710 = *ARK3*; Mable et al., 2017). Upon mapping we intended to extract reads that mapped to *SRK* and adjacent sequences in pairs and to perform a *de novo* assembly of these sequences only. In a first attempt we mapped reads using bwa (Li and Durbin, 2009). However, this approach did not yield any or an extremely low number of reads mapping to *SRK*, while adjacent regions were covered by the expected number of sequencing reads. Since bwa expects reads to have an identity of 90% or more to the reference and *SRK* alleles show much lower similarity (as little as 70% identity), we were not successful in mapping *SRK* reads to the reference. In a second attempt we used Next Gen Mapper (Sedlazeck et al., 2013), which only requires 65% of identity between read and reference. By this approach we were able to map reads to the S-locus including *SRK* but nevertheless a *de novo* assembly of these reads into complete or partial copies of the *SRK* locus failed.

We used CLC genomics workbench (<https://www.qiagenbioinformatics.com/>) to perform *de novo* assemblies using standard settings (automatic word and bubble size, minimum contig length 500 bp, reads were mapped back to contigs setting mismatch costs, insertion costs and deletion costs to 3 and length fraction as well as similarity fraction were set to 0.9) and the scaffolding option. Resulting scaffolds/contigs were indexed as BLAST libraries. We initially used FJ867321 (the S-domain from *AlySRK30*) to BLAST against these libraries to pull out sequences predicted to be *SRK* based on more than 50% coverage of the query sequence (filtered for low complexity, expect set to 10, word size to 11, match to 2, mismatch to -3, gap existence to 5, gap extension to 2). These hits were aligned to the first exon of AL7G32730 (*AlySRK13* from the MN47 reference genome) to identify intron/exon boundaries and then trimmed if necessary. This approach yielded in total 66 sequences in the 13 accessions analyzed (Supplementary Table 10). Therefore,

our BLAST search also must have identified other S-domain encoding genes besides *SRK*.

In order to obtain an overview on the presence of S-domain encoding genes we performed another BLAST search using the first exon of the MN47 *SRK* against the MN47 reference genome. This search revealed five genes encoding proteins that have an S-domain (AL7G32730 = *SRK*, AL7G32710 = *Aly8*, AL6G48380 = *Aly3*, AL3G23610 = *Aly9*, AL2G23090 = *Aly10.2*). From this result we expected that our contigs identified in the 13 resequenced accessions should have their best BLAST hit with one of these five loci. So, we aligned the 66 contig sequences to the first exon of the MN47 *SRK* and trimmed them in length to the first exon. Then we performed a blast search of the 66 trimmed sequences against the MN47 reference genome. All of the 66 sequences had their best blast hit with one of the five loci we had identified beforehand. Typically hits for AL7G32710, AL6G48380, AL3G23610, AL2G23090 showed a very small E-value and a high score while AL7G32730 hits were characterized by a lower score and E-value due to the lower conservation for alleles of this locus.

We initially used the BLAST results to predict similarity to known *SRK* alleles and related receptor kinase gene family members available in Genbank for each of the contigs. However, since we had identified potentially new variants in this study, we also aligned sequences pulled out from the resequenced genomes to our reference database and to the sequences found using 454 and the longer Sanger sequences to confirm sequence identity (Supplementary Data Sheet 2; Supplementary Table 12). We used clustering in phylogenetic trees (reconstructed using Maximum Likelihood in MEGA 7.0) to predict *SRK* specificity and to determine presence/absence of other members of the gene family.

One of the paralogs (*Aly9*) is known to amplify in all *A. lyrata* individuals that have been tested using PCR-based screening (Mable, personal observation). We thus used identification of this locus as a control for whether it was likely that the genome-mining approach could be reliably used to detect copy number variation in highly polymorphic gene families. The approach described initially only identified this locus in three of 12 genomes so we trialed another approach, using the sequences in Supplementary Data Sheet 2, along with the 66 contigs originally identified to BLAST the *de novo* assemblies for each genome. This resulted in an additional 102 contigs, which then were aligned back to the reference database and identities confirmed using cluster analysis. In this analysis *Aly9* was resolved for all individuals and more complete genotypes were obtained for *SRK* and the other paralogs screened, so only the results from this final analysis are presented.

RESULTS AND DISCUSSION

Objectives 1 and 2: Diversity and Allele Sharing of *SRK* in Diploids and Tetraploids

After filtering and assigning variants to alleles based on sequence similarity and predicting dominance classes and linkage to the SI phenotype based on phylogenetic clustering, we identified 107

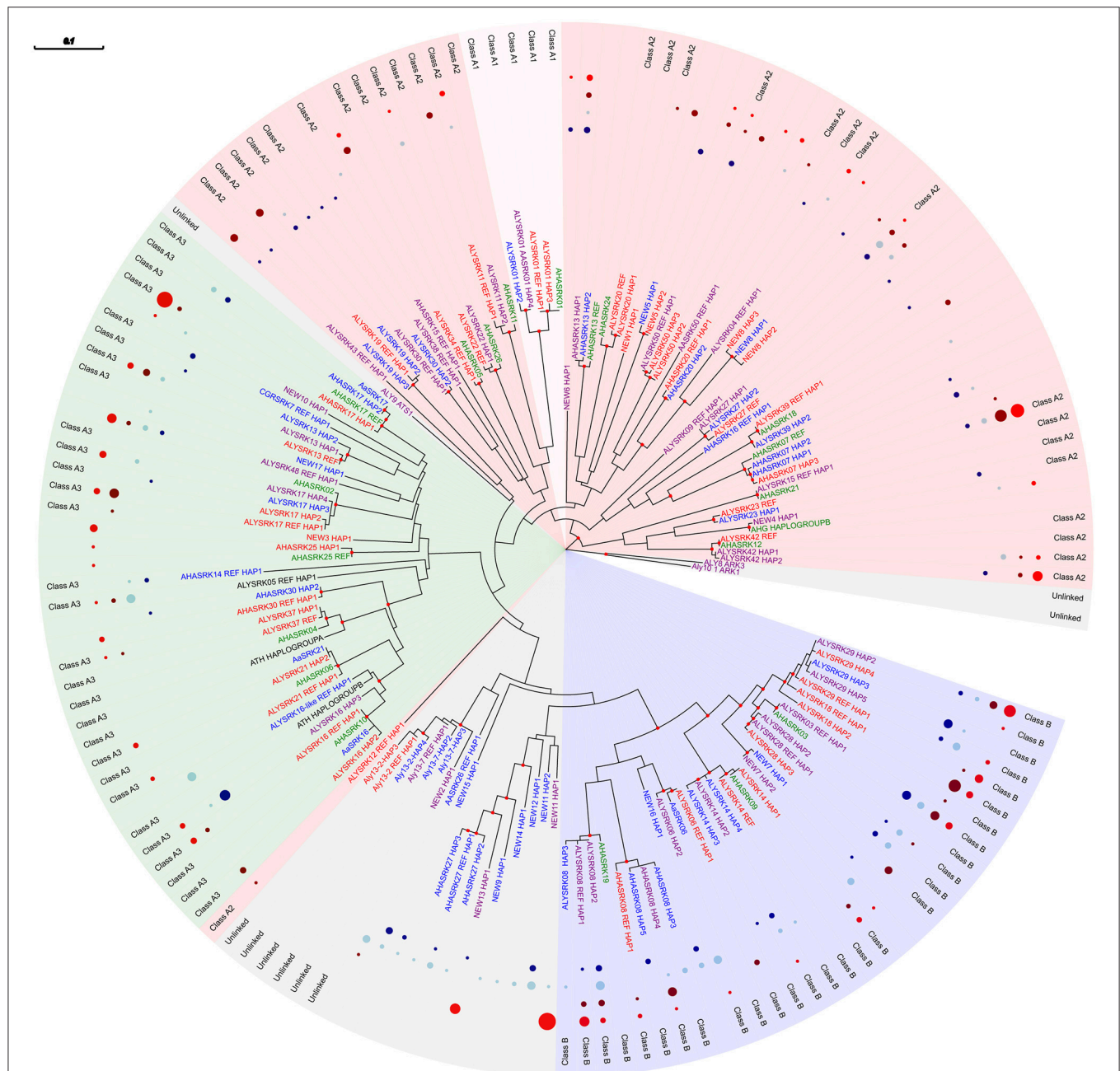


FIGURE 1 | Maximum likelihood tree based on SRK-like sequences resolved through 454 pyrosequencing, reconstructed using MEGA 7 under an HKY85 model of evolution with rate heterogeneity modeled under a gamma distribution and with proportion of invariant sites estimated. Bootstrap proportions above 70% are indicated as filled circles on nodes. The tree was rooted with the unlinked paralogs *Aly8* (*Ark3* in *A. thaliana*) and *Aly10.1* (*Ark1* in *A. thaliana*). Alleles for each SRK specificity are assigned to a dominance class based on previous studies of *A. lyrata* (Prigoda et al., 2005) and *A. halleri* (A1 = yellow; A2 = red; A3 = green; B = blue; unlinked = gray); new alleles or previously identified alleles where dominance has not been confirmed are colored according to the class predicted by their position in the tree. Tip labels are colored according to the species in which they were found in the 454 sequences (*lyrata* = red; *lyrata*+*arenosa* = purple; *arenosa* = blue) or the origin of the reference allele in cases where there was no exact match (*halleri* = green; *thaliana* = black). Also shown is the frequency of a particular haplotype in each of the four groups compared (diploid *arenosa*, A2x = dark blue; tetraploid *arenosa*, A4x = light blue; diploid *lyrata*, L2x = dark red; tetraploid *lyrata*, L4x = light red). Due to the high number of haplotypes but low read numbers for *AlySRK01* and the unlinked loci *Aly13-2* and *Aly13-7*, only a subset of haplotypes are included and frequencies are not indicated.

haplotypes (unique sequence variants) that could be grouped into 63 potential alleles (specificities) that were at least 80% similar to SRK (Figure 1; Supplementary Table 2). Seventeen

were potentially new specificities that were <90% similar to the *A. lyrata*, *A. halleri* or *A. arenosa* reference sequences included (Supplementary Table 3). However, seven of these new variants

were predicted not to be linked to SI based on phylogenetic clustering and so could represent other members of the gene family. All of the new potentially unlinked alleles were found in diploid and/or tetraploid *A. arenosa*, with two of them also occurring at high frequency in L4x populations but only a single L2x individual sharing one of the new unlinked alleles with A4x individuals. The new alleles predicted to be linked to SI were distributed more evenly among the two species.

When accounting for variation due to lane and tag as random effects using generalized linear models, we found no evidence for significant differences between species or ploidy levels or their interactions in terms of number of reads, total number of contigs resolved (indicative of the wider gene family), the number of SRK-like alleles (i.e., variants showing at least 80% similarity to known SRK sequences, so including unlinked alleles), or the number of alleles or haplotypes per individual predicted to be linked to SRK (**Supplementary Table 4**). There was a significant interaction between ploidy and species in the proportion of contigs resolved that were at least 80% similar to SRK (i.e., more reads were SRK-like than similar to other members of the gene family), with a significantly higher proportion in tetraploids compared to *A. arenosa* diploids but no significant difference compared to *A. lyrata* diploids. Since the primers used were developed based on variation within *A. lyrata* (Schierup et al., 2001; Charlesworth et al., 2003a, 2006), this could be an indication that not all SRK-like alleles were amplified for *A. arenosa* due to variation in the primer regions, resulting in resolution of more spurious contigs due to non-specific amplification. However, overall, there was very little evidence that tetraploids were fundamentally different to diploids in terms of sequence quality or the ability to resolve variants.

The 200 bp sequences produced similar resolution in phylogenetic clustering as previous studies using 600 bp (Tedder et al., 2011) and resulted in consistent patterns of polymorphism expected for dominant and recessive alleles at SRK. Examination of relative frequency distributions also generally met theoretical expectations but indicated no obvious differences in diversity between ploidy levels. There was extensive variability in relative frequencies of each haplotype, with some being restricted to certain species or populations and some being found across both species and ploidy levels (**Figure 1**; **Supplementary Table 2**). We predicted that there should be highest interspecific sharing of individual haplotypes among tetraploids due to their known introgression (Schmickl, 2009; Schmickl et al., 2010; Jørgensen et al., 2011; Schmickl and Koch, 2011) but also because they can maintain more allelic copies within individuals. We found that 23 haplotypes were shared between A4x and L4x compared to 12 between A2x and L2x, including seven that were shared among all four population types (**Supplementary Table 2**). Sharing between the two types of tetraploids was similar to that among ploidy levels within species (24 among *A. lyrata* and 22 among *A. arenosa*). The highest number of private haplotypes was also found for diploids: 19 for A2x and 15 for L2x, compared to 12 for A4x and 8 for L4x. These results are consistent with predicted patterns of introgression among the tetraploids in northeastern Austria (Wachau region and Forealps; Schmickl et al., 2010; Jørgensen et al., 2011; Schmickl and Koch, 2011).

Although it is difficult to separate increased transpecific polymorphism from this introgression, we found some evidence that there might be more differences in selection pressure or demographic history between species than between ploidy levels. Plotting allele frequency distributions for each ploidy and species combination demonstrated an excess of intermediate frequency alleles in both diploid and tetraploid *A. arenosa* (**Figure 2**), as expected for a locus under balancing selection (Mable and Adam, 2007). However, the pattern was more skewed toward low frequency alleles in *A. lyrata*, particularly in tetraploids. In North American populations of *A. lyrata*, a difference in allele frequency spectrum for SRK was found between inbreeding and outcrossing populations (Mable and Adam, 2007) but the latter showed more similar patterns as those observed for *A. arenosa* in this study. Since shifts toward intermediate frequencies are also expected for population bottlenecks (Luikart et al., 1998), it is possible that in particular diploid *A. arenosa* experienced a larger decline in population numbers since the past glaciation. What was striking in the current study was that tetraploids did not have a dramatically higher number of alleles or haplotypes within populations or alleles or haplotypes per individual than diploids, regardless of dominance class (**Table 2**). Furthermore, for neutral genes, there is a steep gradient of increasing genomic contribution of A4x found within introgressed *A. lyrata* along a transect in the hybrid zone (Schmickl, 2009; Schmickl et al., 2010; Jørgensen et al., 2011; Schmickl and Koch, 2011; Hohmann et al., 2014; Muir et al., 2015) but this is not reflected in the SRK distribution; i.e., SRK are more mixed than would be predicted based on neutral patterns, as might be expected under balancing selection. This suggests that tetraploids are not fundamentally different from diploids in their capacity for maintaining diversity of SRK, as suggested previously from segregation analyses within tetraploid families based on crosses involving one of the same tetraploid populations studied here (L4_AUT2) and a tetraploid population from Aggsbach, Austria (Mable et al., 2004).

Consistent with theory (Billiard et al., 2007), recessive alleles in diploids have been demonstrated to occur at higher frequency, to show shallower branch lengths in phylogenetic analyses, and more extensive polymorphism within specificities than dominant alleles (Llaurens et al., 2008, 2009; Castric et al., 2010; Vekemans et al., 2011; Goubet et al., 2012). In our study, Class B alleles (recessive to A2 and A3 classes) showed lower intraclass polymorphism (13% average pairwise sequence divergence, compared to 25% for Class A2 and 15% for Class A3) but more haplotypes per allele than the two dominant classes (2.56 ± 1.33 compared to 1.59 ± 0.75 in Class A2 and 1.56 ± 0.89 in Class A3, **Table 2**) and there was high divergence between classes (26–29%; **Table 3**). The paralogous locus identified in previous studies that is similar to class B alleles (*Aly13-2*) showed similar within locus variation (13%) as for class B alleles and lower divergence from class B than the other dominance classes (16% compared to at least 27% to the others). There was a higher proportion of alleles restricted to only one of the species among the dominant (29% for Class A2 and 50% for Class A3) than recessive (20% for Class B) alleles but a majority of the unlinked alleles (67%) were only found in *A. arenosa* (**Supplementary Table 2**). Thirteen alleles were found only in

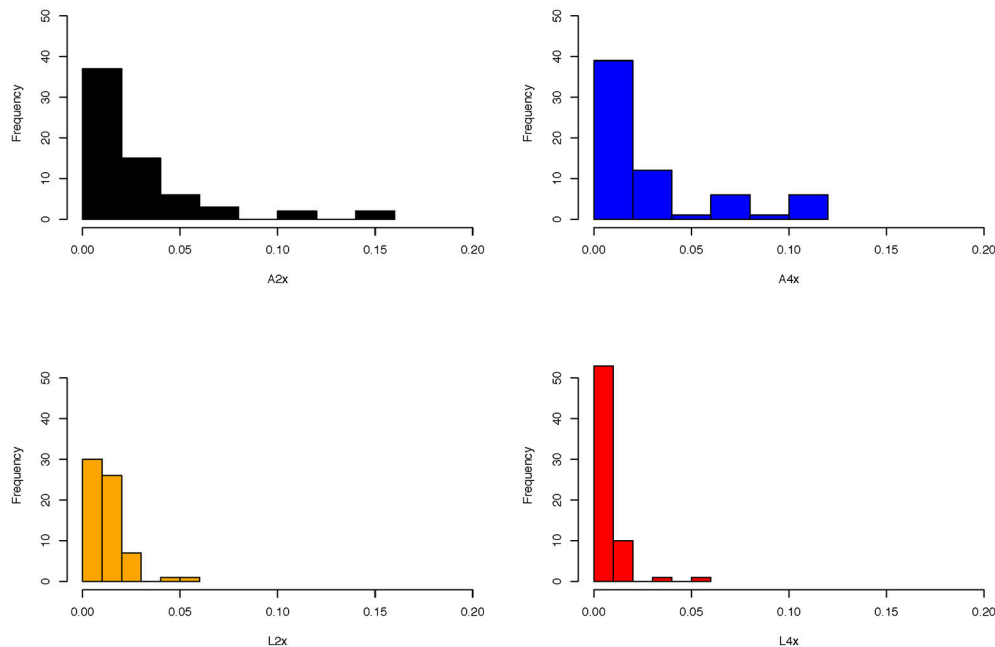


FIGURE 2 | Allele frequency distributions of *SRK* alleles identified in diploid and tetraploid populations of *A. lyrata* and *A. arenosa* using 454 pyrosequencing. Note that there appears to be an excess of intermediate frequencies in *A. arenosa* (A2x = diploids; A4x = tetraploids), with more of a skew toward low frequency alleles in *A. lyrata*, particularly in tetraploids.

tetraploids, but none were Class B and only four (three Class A2 and one Class A3) were shared between the two species. Thus, results were consistent with the increased trans-specific polymorphism expected for recessive alleles at a locus under balancing selection (Billiard et al., 2007; Llaurens et al., 2008; Castric et al., 2010; Goubet et al., 2012).

Overall, these results suggest that tetraploids do not show increased mate availability due to an increase in *S*-locus repertoire but instead might be constrained by the potential mate limitation caused by having “too many” *S*-alleles. This is similar in theory to expectations for immune genes in animals, where an optimal number of alleles has been suggested as conferring higher fitness than maximizing allelic diversity (Reusch et al., 2001; Aeschlimann et al., 2003; Wegner et al., 2003; Kalbe et al., 2009). The high allele sharing among ploidy levels precluded testing of whether there is relaxed balancing selection acting in tetraploids but this was not suggested by the site frequency distributions, which suggested a stronger species than ploidy effect. Nevertheless, there are some important caveats to consider in the interpretation of these results, due to particular challenges when working with this type of gene families (see Challenges below).

In the crosses between tetraploid *A. lyrata* individuals, we found the same three *SRK01* haplotypes using both 454 and targeted Sanger sequencing (haplotypes 1, 2, and 3). This allowed us to test the accuracy of the 454 genotyping despite the low read numbers for *SRK01* and provide more complete data for segregation analyses. For 50% of the individuals identical genotypes were predicted using the two approaches,

with 14% testing negative for the allele-specific PCR but positive using 454, compared to 10% showing the opposite pattern (Table 4). Different haplotypes were predicted by the two methods only for a single individual. However, the direct sequencing was more sensitive, resolving heterozygotes in 24% of the individuals that were predicted to be homozygous based on the 454 sequencing (compared to only 2% showing the opposite pattern). Segregation of *SRK01* genotypes in the crosses confirmed previous predictions (Mable et al., 2004) that tetraploids could harbor multiple copies of haplotypes for this recessive specificity (Table 5). These data were then combined with segregation of the haplotypes resolved using 454 pyrosequencing (Supplementary Table 5). After excluding 454 alleles not present in the parents, the majority of individuals showed four or fewer expected haplotypes. Comparison of segregation of predicted genotypes with self-incompatibility phenotypes (Figure 3; Supplementary Tables 6, 7), confirmed linkage of two alleles previously tested in other crosses (*SRK16* and *SRK29*) and one that had been identified in the grandparents but had not been deposited to Genbank (*SRK48*). However, the segregation analyses suggested that not all alleles were detected by 454 and suggested that the stringent filtering in some cases omitted alleles that must have been present based on the incompatibility phenotypes.

Challenge: Filtering Decisions for Clustering

Despite recommendations from our pilot study that a threshold of 90% similarity would be appropriate for clustering (Jørgensen et al., 2012), our analyses of the full dataset suggested that

TABLE 2 | Distribution of alleles (A) and haplotypes (B) across diploid (A2x, L2x) and tetraploid (A4x, L4x) populations (POP) for different predicted dominance classes (A2 and A3 are dominant to B), excluding Class A1, which is represented only by *SRK01*; read numbers were too low to be certain about presence or absence for that allele.

(A)									
POP	N IND	N ALLELES				N ALLELES/IND			
		A2	A3	B	ALL	A2	A3	B	ALL
A2x	75	16	9	10	35	0.21	0.12	0.13	0.47
A4x	77	15	11	13	39	0.19	0.14	0.17	0.51
L2x	70	18	7	8	33	0.26	0.10	0.11	0.47
L4x	102	14	11	10	35	0.14	0.11	0.10	0.34

(B)									
POP	N IND	N HAPLOTYPES				N HAPLOTYPES/IND			
		A2	A3	B	ALL	A2	A3	B	ALL
A2x	75	19	9	18	46	0.25	0.12	0.24	0.61
A4x	77	15	11	19	45	0.19	0.14	0.25	0.58
L2x	70	20	9	13	42	0.29	0.13	0.19	0.60
L4x	102	16	14	17	47	0.16	0.14	0.17	0.46
Total		70	43	67	180				

Alleles that did not appear to fall under any of the known dominance classes are not included, as they were predicted to be unlinked to the *SI* phenotype. Also shown is the number of alleles or haplotypes per individual.

TABLE 3 | Percent sequence divergence within and between dominance classes, for alleles identified using 454 sequencing.

Class	B	A1	A2	A3	Aly13-2
B	0.129				
A1	0.282	0.032			
A2	0.287	0.259	0.253		
A3	0.279	0.265	0.277	0.151	
Aly13-2	0.159	0.284	0.272	0.267	0.129

Divergence within classes is shown on the diagonal. An unlinked locus that shows polymorphism among haplotypes (Aly13-2) is included for comparison.

a single threshold may not be appropriate for gene families that include different levels of divergence among classes or copies; for example, in relation to dominance (Prigoda et al., 2005). In our study, BLAST analysis of “read only” contigs demonstrated that some known alleles were fragmented across multiple contigs. For recessive alleles (Class B, *SRK01*) and unlinked loci (*Aly9*, 13-2 and 13-7), combining contigs resulted in mixtures of haplotypes from different alleles (specificities), making it challenging to assign sequence variants to alleles. While several dominant alleles (*Aly16*, *Aly30*, and *Aly42*) also showed fragmentation, there was no ambiguity in assigning sequence variants to alleles. Resolving recessive alleles into unique contigs thus required more manual manipulation and sorting of variants into haplotypes. Since recessive alleles also had on average more haplotypes per allele (2.44 ± 1.42) than dominant alleles (1.57

TABLE 4 | Proportion of individuals that tested positive for *SRK01* specificity using direct Sanger and 454 sequencing, indicating the population (A2x = diploid *A. arenosa*; A4x = tetraploid *A. arenosa*; L2x = diploid *A. lyrata*; L4x = tetraploid *A. lyrata*), sample sizes (N-direct, N-454) and % of individuals that tested positive for *SRK01* in each.

Population	N-direct	% <i>SRK01</i> -direct	N-454	% <i>SRK01</i> -454
A2x	44	20.5	78	66.7
A4x	65	44.6	87	78.2
L2x	57	31.6	76	55.3
L4x	79	41.8	115	61.7
Total	245	36.6	191	59.2

± 0.74 for A2; 1.56 ± 0.89 for A3) (Supplementary Table 2), read numbers per haplotype were often lower, which made setting a single threshold for reducing spurious genotyping difficult.

Challenge: Amplicon Based Errors and Biases

From previous studies we anticipated that the single most recessive allele, *SRK01*, would be present at high frequency and would show a higher number of haplotypes than other specificities (Billiard et al., 2007; Castric and Vekemans, 2007; Llaurens et al., 2008; Castric et al., 2010; Goubet et al., 2012; Vekemans et al., 2014). In our 454 data, *SRK01* was present in all populations surveyed and we identified 15 unique variants that were present in more than one individual; however, read numbers tended to be very low (often with <10 reads per individual) and fell well below the thresholds set for considering “real” presence of a given haplotype used for other loci for most individuals. Although multiple haplotypes differing by a single or few bp are expected for recessive alleles (Castric and Vekemans, 2007), the low read numbers made it difficult to distinguish PCR errors from actual polymorphism. High read numbers were found for some individuals, but they tended to show the presence of few other sequence variants. In addition, several known paralogs that should be present in all individuals (*Aly8*, *Aly9*; Charlesworth et al., 2003b) were expected to amplify with the primer set used but this was very inconsistent. *Aly9* was present in the majority of individuals but read numbers varied dramatically from 0.5 to 92% of the total reads in an individual. There was a significant difference in the proportion of reads that were *Aly9*, with *A. arenosa* tetraploids showing a higher proportion than both diploids, which showed a significantly higher proportion than *A. lyrata* tetraploids (Supplementary Table 4). Whether this is due to an amplification bias or expansion of the gene family is difficult to distinguish. For *Aly8*, only 41/460 sequenced individuals showed any amplification and most were present at only low read numbers (maximum 15%). We thus could not assess presence or absence of other members of the gene family based on the 454 sequencing or use the paralogs to make inferences about introgression in the tetraploids to avoid the confounding effects of balancing selection. Even after correcting for chimeras, there was some evidence for recombination in some of the specificities showing polymorphism among populations (e.g., *SRK01*, some of the class B alleles) but this was difficult to

as we performed crosses before the 454 sequencing, we could not test linkage of all new variants found to the SI phenotype. It was also not feasible to determine when unlinked alleles were amplified based on the presence of “too many” haplotypes.

Objective 3: Introgression of *SRK* Alleles

For the population survey, the 454 genotyping identified 22 *SRK01* variants. Using targeted amplifications and Sanger sequencing we identified 24 haplotypes. All of these but seven had been found using the 454 pyrosequencing, but including five that matched the 454 sequences but had additional polymorphisms outside of the shared sequence region (indicated by distinct letters after the haplotype name; **Supplementary Tables 8, 9**). However, only 11 of the 22 variants found by 454 sequencing were confirmed by direct sequencing and there was a higher proportion of PCR positive results among the 454 than the Sanger sequences (**Table 4**).

Using the diploids as a guide, we identified “arenosa” and “lyrata” specific haplotypes, as well as three that appeared to be recombinants between species-specific variants (haps 7, 8, and 10; **Supplementary Data Sheet 3**), two of which were identified from a single A4x population that was predicted to be introgressed (A4X_AUT1, from Kerhnoff; Schmickl, 2009). Although analyses using GARD in the HYPHY package did not find statistical evidence for recombination breakpoints, this might have been because of the short tracts of introgression. The minimum spanning network indicated that haps 7 and 8 did in fact fall between species-specific clusters whereas hap10 was on a tip in the *A. arenosa* part of the network (**Figure 4**). What is striking is that reticulation in the network involved primarily *A. arenosa* tetraploids and that diploids had a lower diversity of *SRK01* haplotypes compared to tetraploids. There was also some haplotype sharing among tetraploids but not between the diploids. Since the crosses established that individual tetraploids could harbor up to three different *SRK01* haplotypes and many were heterozygous for two, this higher diversity among tetraploids could be because *SRK01* is effectively neutral and so could accumulate more mutations in tetraploids because of the higher copy number maintained (Mable et al., 2004). Crossing data suggest that *SRK* is functional in individuals sampled from the hybrid zone (Ruiz-Duarte, 2012), but it is also possible that selection pressure to maintain restricted recombination in the *S*-locus region (Charlesworth et al., 2006) would be relaxed with the increased copy number in tetraploids. Moreover, introgression of recessive alleles between *A. lyrata* and *A. halleri* has been found in diploids (Castric et al., 2010), suggesting that hybridization might disrupt linkage. Although the crosses we performed only included tetraploid *A. lyrata* from outside of the known hybrid zone, two individuals in one family were self-compatible (**Supplementary Table 6**). It is thus also possible that increased recombination at the *S*-locus occurs with spontaneous loss of SI in some individuals.

The presence of *A. arenosa* like haplotypes in two of the *A. lyrata* tetraploid populations and the most frequent *A. lyrata*

haplotype (hap1) in most *A. arenosa* populations from the hybrid zone (**Table 4**, **Supplementary Table 9**) could suggest more recent and secondary hybridization while the introgressed haplotypes (i.e., those that appeared to be recombinants between the species-specific variants) could reflect older events. One *A. arenosa*-like haplotype (hap2) was found in an *A. lyrata* tetraploid population in the Northeastern Austrian Forealps (L4_AUT4 from Lilienfeld), and in the crosses, which involved individuals from two peripheral *A. lyrata* populations (L4_AUT2 from Mödling and L4_AUT5 from Rauhenneck Ruin, near Baden). This could suggest undetected hybridization within these “pure” populations, as also suggested by whole-genome data (Hohmann and Koch, 2017). While these results fit with expectations based on predicted patterns of hybridization in tetraploid populations from Austria (Schmickl, 2009; Schmickl et al., 2010; Schmickl and Koch, 2011; Muir et al., 2015), there are similar caveats about the use of PCR-based genotyping as raised for the 454 sequences, as described below.

Challenge: PCR Based Approaches to Genotyping

Overall, there was not much consensus between the *SRK01* genotypes resolved using 454 and direct sequencing. While the crosses demonstrated that the latter was more sensitive to detect heterozygotes when products were amplified, the population survey revealed a potential bias against amplifying variants found in A2x populations. A much lower proportion of individuals from these populations tested positive than from other populations, and many of the haplotypes found using 454, but not direct sequencing, were from A2x populations. This potential bias reduced the sample sizes that could be used to classify haplotypes showing species-specific presence. In the segregation analyses (**Table 5**), two individuals had all three *SRK01* haplotypes segregating in the parents: one individual didn’t show presence of other alleles expected in the parents based on the 454 sequencing but showed some unexpected alleles; the other individual showed more than four expected haplotypes (**Supplementary Table 5**). Thus, we cannot rule out contamination. Moreover, interpretation of introgressed haplotypes could have been confounded by PCR-based recombination but they were found only in a stabilized hybrid population. Moreover, some haplotypes were only resolved from direct sequences of heterozygotes; in those cases cloning would be required to absolutely confirm the full range of haplotypes present. We had originally intended to also test the utility of other polymorphic members of the gene family (e.g., *Ally9*); however, since there were even more haplotypes predicted by the 454 sequencing but separated by fewer variants (data not shown), there would have been too much reliance on accurately identifying singletons.

Objective 4: Copy Number Variation in the *SRK*-Related Gene Family

Clustering of contigs resolved from the *de novo* assembly approach to genome mining of our database of *SRK* and its paralogs (i.e., all unique variants found using the 454 pyrosequencing, targeted sequencing of *SRK01*, cloning of longer

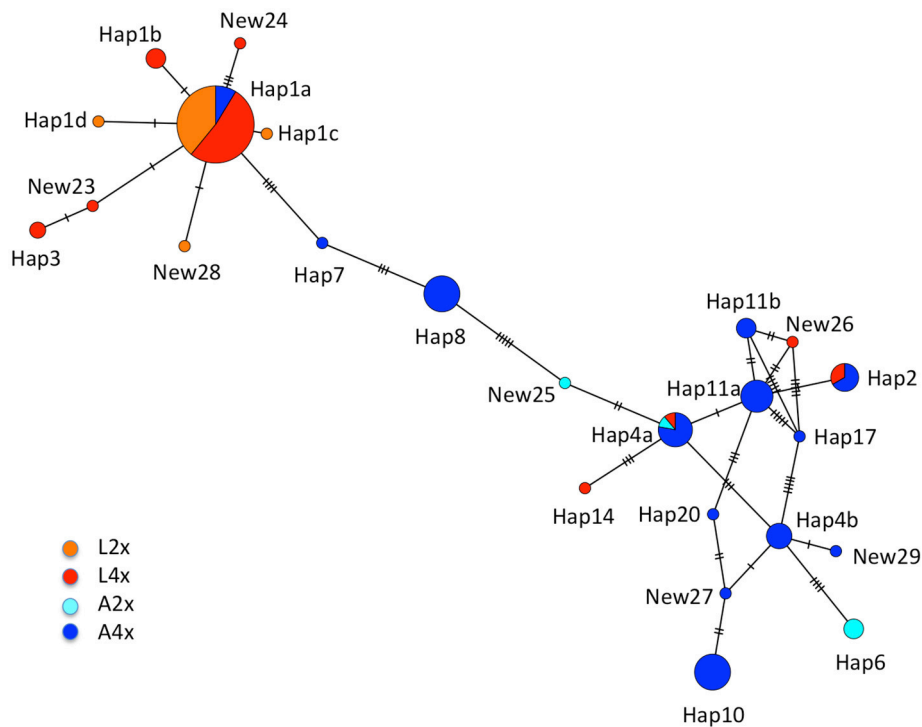


FIGURE 4 | Minimum spanning network for *SRK01* haplotypes resolved using direct Sanger sequencing. Circles are drawn proportionately to the frequency of the haplotype and colored by relative frequency in each population type: A2x = light blue; A4x = dark blue; L2x = orange; L4x = red. Vertical bars on the connecting branches indicate the number of nucleotide substitutions separating haplotypes. Haplotypes 7, 8, and 10 were predicted to be recombinants between “arenosa” and “lyrata” specific sequences; haps 7 and 8 appear intermediate between the two clusters whereas hap10 is on a tip in the “arenosa” part of the network. NEW25 also appears intermediate and so could be another introgressed haplotype but it was only found in a single individual. Note that extensive reticulation was found predominantly among sequences found in A4x populations and that there is less variation among haplotypes restricted to diploids than those found in tetraploids.

products using degenerate primers, and additional sequences available in Genbank) was used to uncover receptor-like kinases from published diploid and tetraploid genome sequences (Supplementary Table 10). This resulted in identification of 1-2 predicted *SRK* alleles in the diploid and 1-4 in the tetraploid accessions for both species among the 13 short read sets screened (Table 6; Supplementary Table 11). In total 29/177 contigs were assigned as *SRK*, but 12 of these would have been mis-assigned based only on BLAST (Supplementary Table 10). *Aly13-2*-like sequences were pulled out in seven accessions, but would have been classified as *SRK* based only on BLAST (Table 6; Supplementary Table 10). This locus is not present in all individuals, so copy number variation is expected (Mable et al., 2017). Other alleles predicted to be unlinked to the SI phenotype were also resolved by the clustering analysis but none of these would have been assigned as *SRK*-like based on BLAST (Table 6; Supplementary Table 10). One published allele (*AlySRK32*) whose phylogenetic position and dominance have not been resolved in previous studies was pulled out from six accessions; based on the length of its branch to other *SRK* sequences, it has been predicted to be unlinked to the SI phenotype (Tedder et al., 2011). *AlySRK47* (found in four of the accessions) is also predicted to be unlinked, based on its phylogenetic position relative to linked sequences. The other four paralogs tested were present in all accessions, except for

one diploid *A. lyrata* that lacked *ARK3* (*Aly8*). Since this latter locus is tightly linked to *SRK* in some specificities and shows high polymorphism (Kusaba et al., 2001; Charlesworth et al., 2003b; Guo et al., 2011; Vekemans et al., 2014), this could be due to divergence from the reference sequence. AL2G2623090 included sequences similar to both *Aly10.1* (*ARK1* in *A. thaliana*) and *Aly10.2* (*ARK2* in *A. thaliana*), which were detected in all individuals. *Aly10.2* is a suspected pseudogene in *A. lyrata* due to a large deletion and does not amplify in all individuals (Charlesworth et al., 2003b), whereas *Aly10.1* is predicted to be functional and amplifies in more individuals. Clustering suggested that only four individuals had both genes but not all contigs could be resolved due to missing parts of the sequence. AL6G484380 (*Aly3*) was found in all individuals. Fourteen of the contigs clustered into two distinct clades that did not show similarity to any known paralogs (contig-only clusters; Supplementary Table 10). One was found in 10/13 accessions while the other was only found in four; these could represent previously uncharacterized members of the S-receptor kinase gene family.

For the *SRK* sequences, five of the putatively new alleles found by 454 pyrosequencing were pulled out, all of which also were detected by cloning and sequencing using degenerate primers; multi-exon sequences were mined from the genomes for two of them (NEW2, NEW16; Table 6). Multi-exon sequences were also

TABLE 6 | Identity of *SRK*-like (AL7G32730) contigs pulled out by genome mining and confirmed by phylogenetic clustering.

Accession	Type	AL7G32730 (SRK)				Total linked	Unlinked	Total unlinked
SRR2040821	A4x	<i>AlySRK01</i>	<i>AlySRK15</i>	<i>AlySRK42</i>	NEW17	4		0
SRR2040822	A4x	<i>AlySRK01</i> x 2	<i>AHASRK08</i> x 2			4	<i>Aly13-2*</i> <i>AlySRK32</i>	2
SRR2040825	A4x	<i>AlySRK01</i>	<i>AlySRK12</i>			2	NEW2*	1
SRS945917	A2x	<i>AHASRK17</i>	NEW16*			2	<i>Aly13-2*</i> <i>AlySRK32</i> x 2 <i>AlySRK47</i>	4
SRS1256176	A2x	<i>AlySRK13</i>				1	<i>Aly13-2*</i> <i>AlySRK32</i> <i>AlySRK47</i> NEW9	4
SRS1256175	A2x	<i>AlySRK01*</i>	<i>AlySRK23</i>			2		0
SRR2040827	L4x	<i>AlySRK01</i>				1	<i>AlySRK32</i> <i>AlySRK47</i>	2
SRR2040828	L4x	<i>AlySRK01</i>	<i>AlySRK25</i>	<i>AlySRK33</i>		3	<i>Aly13-2</i> <i>AlySRK32</i>	2
SRR2040829 ^a	L4x	<i>AlySRK01</i>	<i>AlySRK12</i>	NEW17	NEW7	6 ^a	<i>AlySRK47</i>	1
SRR2040830	L4x	<i>AlySRK01*</i>	<i>AlySRK42</i>	<i>AaSRK50</i>		3		0
SRR3111440	L2x	<i>AlySRK15*</i>				1		0
SRR3111441	L2x	<i>AlySRK44</i>	<i>AlySRK17</i>			2	<i>Aly13-7*</i> <i>AlySRK32</i>	2
SRR2040791	L2x	<i>AlySRK01*</i>	<i>AlySRK42</i>			2	NEW2*	0

Contigs in red would have been mis-assigned based only on BLAST; those in blue were not resolved by BLAST. Cloned sequences were also obtained from contigs indicated in bold; asterisks indicate sequences where multi-exon sequences were pulled out using the genome mining. Alleles showing high similarity to *SRK* but not predicted to be linked to the *Sl* phenotype are also indicated (Unlinked). The total number of linked and unlinked alleles resolved per accession is also indicated.

^aAlso has *AlySRK10* and *AlySRK28*.

pulled out for *AlySRK01*, *AlySRK15*, *Aly13-2*, and *Aly13-7*. While the genome mining approach seems promising, the presence of homozygotes for *SRK* for three individuals suggests that not all *SRK* alleles were identified within individual genomes: one L2x and one A2x individual had a single dominant allele each (*AlySRK15*, in dominance Class A2 and *AlySRK13* in dominance Class A3, respectively). One L4x individual was homozygous for *AlySRK01*, which is plausible, as homozygotes for this recessive allele have been found in previous segregation-based analyses of tetraploid *A. lyrata* from Austria (Mable et al., 2004).

Challenge: Extracting Full-Length Sequences of Polymorphic Genes From Short Read Data

While the genome mining holds promise for investigating copy number variation and obtaining full-length sequences from new alleles, the approach that worked best required a detailed reference database of alleles in order to accurately assign sequences to loci. BLAST analyses alone resulted in mis-assignment of *SRK* alleles to other paralogs and other paralogs were sometimes assigned as *SRK* alleles. While part of this was because not all sequences were available in Genbank for BLAST analysis, the gene conversion with unlinked loci that makes similarity alone unreliable (Prigoda et al., 2005) remained problematic in these analyses. For example, *Aly13-2/13-7* sequences (which are not linked but are highly similar to Class B *SRK* alleles) were assigned as *SRK* in the initial analyses using only the five genes extracted from the MN47 genome. Manual alignments and phylogenetic clustering were required to determine allelic identities and to assign sequences to paralogous loci. However, there were clues in the BLAST analyses that suggested mis-assignment of *SRK*-like alleles; a signature of high E-value and low score in all cases predicted clustering to *SRK*-like sequences (although including unlinked loci such as *Aly13-2*). Nevertheless, the presence of only a single dominant allele in some accessions suggested that the genome mining did not pull out all *SRK* sequences that should have been

present (since homozygotes should only be possible for recessive alleles).

While we had hoped also to be able to use this approach to map the potentially new alleles found using 454 sequencing to genomic regions to predict linkage to the *S*-locus, the failure of the mapping approach meant that this was not possible. However, when amplifying longer sequences using degenerate primers, we were able to obtain full-length sequences for some of the potentially new specificities predicted from the 454 analyses that we could use to BLAST the *de novo* assemblies (Table 6).

CONCLUSIONS AND RECOMMENDATIONS

The results presented here suggest that the highly polymorphic *SRK* alleles could be useful for interpreting evolutionary patterns of gene flow among populations, species and ploidy levels. We have demonstrated that tetraploids show no apparent advantage in terms of allelic or haplotypic repertoire due to more relaxed selection than diploids but that there is increased evidence for introgression (at least based on the most recessive *SRK* allele) among tetraploids from suspected hybrid populations. We also demonstrated that following up high throughput genotyping with targeted PCR can help to increase accuracy and completeness. We also identified new alleles not previously characterized and predicted dominance based on phylogenetic clustering.

Nevertheless, there are some important caveats from the analyses, which highlight considerations for future studies based on more robust approaches to high throughput genotyping. We make the following recommendations for future investigations of gene family evolution, in diploids as well as polyploids: (1) applying a hierarchical strategy to filtering decisions for cluster analyses could improve assignment of sequence variants to allelic variants, similar to suggestions for hierarchical AMOVA or STRUCTURE analyses (Holsinger and Mason-Gamer, 1996;

Herdegen et al., 2014); (2) amplicon-based approaches for genotyping using deep sequencing should be avoided if there are other options available, as differential amplification and the difficulty of distinguishing PCR errors from real biological processes are difficult to overcome by any current sequencing technology; (3) due to the difficulty of assigning variants to gene copies, interpretation of gene family evolution should always be accompanied by co-segregation of sequence variants with the phenotype, whenever possible; (4) genome mining of resequenced genomes has the potential to investigate copy number variation and obtain full-length sequences that would be useful for population genetics analyses and tests for selection but lack of assembly of highly polymorphic genes to references means that this might only be practical for genes where there is already extensive knowledge about the components of the gene family.

While our results have demonstrated some useful insights into the dynamics of a complex gene family in polyploids and hybrids, we recommend that non-PCR-based sequence capture approaches hold the most promise for assessing patterns of selection on genes under balancing selection, where trans-specific polymorphism, reduced differentiation among alleles, and intermediate frequency alleles are predicted. Such approaches, for example, have been successfully applied to investigating R-gene variation in crop plants (Jupe et al., 2012, 2013; Andolfo et al., 2014; Giolai et al., 2016; Russell et al., 2016; Van Weymers et al., 2016). Whole genome resequencing approaches could be useful for setting the genomic context and fate of duplications, but there are still substantial challenges to resolve in distinguishing loss of copies from lack of coverage or lack of assembly to the reference due to high sequence divergence. A hierarchical approach to filtering or assembly to multiple references (e.g., multiple individuals or multiple alleles or gene family members) could help to overcome such difficulties but resolving fine-scale variation among variants from errors (e.g., haplotypes within specificities) and resolving complete heterozygous genotypes (particularly in polyploids) will require some creative bioinformatic solutions.

DATA AVAILABILITY STATEMENT

The 200 bp fragments generated by 454 sequencing are too short for submission to Genbank but a full alignment of the sequences

identified has been provided as **Supplementary Data Sheet 1** (including only the 454 sequences and references), **2** (including all unique alleles found across analyses), and **3** (SRK01 sequences). All new Sanger sequences have been deposited to Genbank (Accession numbers: MH507371–MH507400). Accession numbers and details for all unique sequences identified, along with those for reference sequences already available in Genbank are provided in **Supplementary Table 12**.

AUTHOR CONTRIBUTIONS

BM wrote the paper and performed the bulk of the sequence analyses described. MJ, AC, and PR-D generated sequence data for objectives 1, 2, and 3, respectively. MJ also performed the crosses for segregation analyses. KL and CK developed bespoke bioinformatics pipelines for objectives 1 and 4, respectively. AB contributed conceptually and financially to objectives 1 and 2. MK contributed conceptually to all aspects of the project and provided samples and advice on sampling locations and interpretation of the hybrid zones. AB, MK, and CK contributed to writing the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fevo.2018.00114/full#supplementary-material>

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