Identification and Mapping of Eastern Filbert Blight Resistance Quantitative Trait Loci in European Hazelnut Using Double Digestion Restriction Site Associated DNA Sequencing

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Abstract. European hazelnut (Corylus avellana L.) is an economically important edible nut producing species, which ranked sixth in world tree nut production in 2016. European hazelnut production in the United States is primarily limited to the Willamette Valley of Oregon, and currently nonexistent in the eastern United States because of the presence of a devastating endemic disease, eastern filbert blight (EFB) caused by Anisogramma anomala (Peck) E. Muller. The primary commercial means of control of EFB to date is through the development and planting of genetically resistant european hazelnut cultivars, with an R-gene introduced from the obsolete, late-shedding pollinizer ‘Gasaway’. Although the ‘Gasaway’ resistance source provides protection against EFB in the Pacific northwestern United States (PNW), recent reports have shown that it is not effective in parts of the eastern United States. This may be in part because the identification and selection of ‘Gasaway’ and ‘Gasaway’-derived cultivars occurred in an environment (PNW) with limited genetic diversity of A. anomala. The objectives of the current research were to develop a genetic linkage map using double digestion restriction site associated DNA sequencing (ddRADseq) and identify quantitative trait loci markers associated with EFB resistance from the resistant selection Rutgers H3R07P25 from southern Russia. A mapping population composed of 119 seedling trees was evaluated in a geographic location (New Jersey) where the EFB fungus is endemic, exhibits high disease pressure, and has a high level of genetic diversity. The completed genetic linkage map included a total of 2217 markers and spanned a total genetic distance of 1383.4 cM, with an average marker spacing of 0.65 cM. A single QTL region associated with EFB resistance from H3R07P25 was located on European hazelnut linkage group (LG) 2 and was responsible for 72.8% of the phenotypic variation observed in the study. Based on its LG placement, origin, and disease response in the field, this resistance source is different from the ‘Gasaway’ source located on LG6. The current results, in combination with results from previous research, indicate that the H3R07P25 source is likely exhibiting resistance to a broader range of naturally occurring A. anomala isolates. As such, H3R07P25 will be important for the development of new european hazelnut germplasm that combines EFB resistance from multiple sources in a gene pyramiding approach. Identification of EFB resistance in high disease pressure environments representing a diversity of A. anomala populations is likely a requirement for identifying plants expressing durable EFB resistance, which is a precursor to the development of a commercially viable european hazelnut industry in the eastern United States.

The genus Corylus L. (2n = 2x = 22), in the family Betulaceae, order Fagales, is composed of 11 edible nut producing species (Bassil et al., 2013; Chen et al., 1999; Erdogan and Mehlenbacher, 2000; Muehlbauer et al., 2014a; Yoo and Wen, 2002) that are geographically distributed across North America, Europe, and Asia (Mehlenbacher, 1991). The most commercially important species is the european hazelnut,
which ranked sixth in world tree nut production in 2016 (743,455 t) [Food and Agricultural Organization of the United Nations (FAO), 2016]. Approximately 10% of the nuts of this species are sold in-shell, whereas the remaining 90% are processed as dairy, bakery, chocolate, and confectionery products (Ciarmiello et al., 2014; Petriccione et al., 2010).

The top five leading european hazelnut producers in 2016 included Turkey (420,000 t), Italy (120,572 t), the United States (34,473 t), Azerbaijan (33,941 t), and the Republic of Georgia (29,500 t) (FAO, 2016). In the United States, commercial european hazelnut production occurs almost exclusively in the Willamette Valley of Oregon, accounting for 99% of total U.S. production (Chen et al., 2007; Lunde et al., 2000; U.S. Department of Agriculture, 2018). European hazelnut production has been attempted in the eastern United States as far back as the colonial time period (Thompson et al., 1996); however, the presence of a devastating endemic disease, EFB caused by the fungus *A. anomala*, harbored by the tolerant native american hazelnut (*Corylus americana* Marsh.), has prevented the establishment of a viable european hazelnut industry east of the Rocky Mountains (Capik and Molnar, 2012, 2014; Fuller, 1908; Halsted, 1892; Johnson and Pinkerton, 2002; Thompson et al., 1996).

European hazelnut production in the PNW United States remained EFB free from the time of the establishment of the industry in the late 1800s (Thompson et al., 1996) until an inadvertent introduction of *A. anomala* occurred in southwestern Washington in the 1960s (Davison and Davidson, 1973). Since that time, EFB has spread throughout the Willamette Valley of Oregon (Johnson and Pinkerton, 2002; Pinkerton et al., 1992). Disease management practices now include scouting for cankers, extensive pruning, and liberal fungicide applications; however, they are not always completely effective, which is a challenge when dealing with a perennial stem canker disease that has a 16-month latent period. This point, combined with the expense associated with these control measures, indicates that host genetic resistance is the most sustainable, long-term, economically viable solution for EFB management (Chen et al., 2007; Johnson et al., 1996; Julian et al., 2008; Lunde et al., 2006; Mehlenbacher, 1995; Molnar et al., 2013; Sathuvalli et al., 2010).

The first source of host genetic resistance identified in the PNW was the obsolete, late-shedding pollinator *C. avellana* ‘Gasaway’ (Cameron, 1976; Mehlenbacher et al., 1991). Although ‘Gasaway’ exhibits low yields and poor nut and kernel characteristics, when this cultivar is crossed with EFB-susceptible germplasm, it transmits resistance to its progeny in a 1:1 resistant:susceptible ratio, indicative of carrying a dominant EFB resistance allele (R-gene) in a heterozygous state at a single locus (Coyne et al., 1998; Mehlenbacher et al., 1991, 2006; Osterbauer et al., 1997; Sathuvalli et al., 2017). ‘Gasaway’ has subsequently been widely used in the Oregon State University (OSU) european hazelnut breeding program, resulting in the release of many advanced generation european hazelnut cultivars (Mehlenbacher et al., 2009, 2011, 2013, 2014, 2016) and pollinizers (Mehlenbacher and Smith, 2004; Mehlenbacher and Thompson, 1991; Mehlenbacher et al., 2012). Largely based on these new EFB-resistant cultivars, the Oregon european hazelnut industry expanded to 31,809.5 ha in 2018 (N. Wiman, personal communication).

Hazelnut breeders and plant pathologists have been concerned about the long-term durability of, and reliance on, a single source of EFB resistance in the european hazelnut industry (Coyne et al., 1998; Leadbetter et al., 2016; Lunde et al., 2006; Molnar et al., 2010a; Osterbauer, 1996; Pinkerton et al., 1998). Adding to this concern is that *A. anomala* found in the PNW is believed to have originated and spread from a single point introduction (Gottwald and Cameron, 1980; Johnson et al., 1996; Pinkerton et al., 1993), raising the possibility that the ‘Gasaway’ R-gene may have been initially discovered and screened against a limited number of fungal isolates. Support for this hypothesis can be found in recent research using microsatellite or simple sequence repeat (SSR) markers to study the diversity of *A. anomala* samples collected from multiple locations across the United States. Studies showed that *A. anomala* isolates collected from Oregon were genetically similar, whereas isolates collected from east of the Rocky Mountains were found to be much more genetically diverse (Cai et al., 2013; Muehlbauer et al., 2014b, 2019; Tobia et al., 2017). In addition, ‘Gasaway’ and its offspring, which have been shown to be resistant to EFB in Oregon and were initially resistant to EFB in New Jersey (T.J. Molnar, unpublished data), have recently been shown to be susceptible to EFB at Rutgers University in New Jersey (Capik and Molnar, 2012; Capik et al., 2013; Molnar et al., 2010a, 2010b; T.J. Molnar, unpublished data). This is a location where the fungus is endemic, exhibits greater genetic diversity (Cai et al., 2013; Muehlbauer et al., 2014b, 2019), and more important, has also been shown to exhibit pathogenic variation (Molnar et al., 2010a; T.J. Molnar, unpublished data).

In response to the concerns and observations listed previously, both the OSU and Rutgers University hazelnut breeding programs have outlined goals to collect, identify, and develop *C. avellana* germplasm sources with EFB resistance that differ from the ‘Gasaway’ R-gene source. This has included the goals of searching for R-genes exhibiting resistance to multiple isolates of *A. anomala*, and/or the identification or development of *C. avellana* germplasm with multiple R-genes in a gene pyramiding approach. Starting in 2002, a wide germplasm collection was made across the native range of *C. avellana* in eastern Europe and the Caucasus Republics. In total, more than 5000 seedling trees were grown from nearly 200 individual seed lots collected across 10 countries. After 16 years of evaluation in New Jersey, ~3% of this collected germplasm has remained resistant to highly tolerant of EFB (Capik et al., 2013; Leadbetter et al., 2016; Molnar et al., 2007, 2018), whereas the ‘Gasaway’ source has not offered the same level of protection (Capik and Molnar, 2012; Capik et al., 2013; Molnar et al., 2010a, 2010b; Muehlbauer et al., 2018; T.J. Molnar, unpublished data) and has shown differential pathogenic responses to *A. anomala* isolates in New Jersey and elsewhere across the native range of the fungus (Molnar et al., 2010a; T.J. Molnar, unpublished data). These observations and the wide origins of the plant materials (Muehlbauer et al., 2014a), when considered together, suggest that some of the newly identified EFB-resistant germplasm may not only differ from the ‘Gasaway’ source but also provide resistance to a broader range of *A. anomala* isolates.

The *C. avellana* reference genetic linkage map and trait association research at OSU have shown that the putative ‘Gasaway’ R-gene resides on european hazelnut LG6 (Colburn et al., 2015, 2017; Gürçan and Mehlenbacher, 2010; Mehlenbacher et al., 2006; Sathuvalli and Mehlenbacher, 2013; Sathuvalli et al., 2012, 2017). Interestingly, related work at
OSU showed that resistance from the Spanish cultivar Culplà, the Serbian cultivar Crvenje, and OSU 495.072 (a seed source from the N.I. Vavilov Research Institute of Plant Industry in St. Petersburg, Russia) also map to LG6 in a region similar to the ‘Gasaway’ source (Colburn et al., 2015). However, resistance from the Spanish cultivar Ratoli was shown to reside on LG7 (Sathuvalli et al., 2011a), as does resistance from C. americana ‘Rush’ (Bhattarai et al., 2017), whereas resistance from C. avellana OSU 759.010 (a selection from the Republic of Georgia) resides on LG2 (Sathuvalli et al., 2011b). All of these genetic mapping and QTL identification studies were conducted in the PNW (Oregon), where the genetic diversity of A. anomala appears to be limited. The objective of the current research was as follows: 1) conduct ddRADseq-based genetic linkage mapping and QTL experiments in a cross of European hazelnut selection Rutgers H3R07P25, resistant to EFB in the eastern United States, and EFB-susceptible selection OSU 1155.009; 2) expose full-sib progeny from that cross to naturally occurring high EFB disease pressure in New Jersey where A. anomala is endemic and genetically diverse; and 3) compare the LG locations of newly identified EFB resistance QTLs to those previously reported.

Materials and Methods

Plant Material

MAPPING POPULATION PARENTS. The full-sib mapping population used in the current study was developed from a controlled cross between two heterozygous individuals, with EFB-resistant H3R07P25 as the female parent and EFB-susceptible OSU 1155.009 as the male parent. H3R07P25 originated as a single EFB-resistant selection from a population of trees (seed lot RUS 12) grown from open-pollinated seed collected in Aug. 2002 at an outdoor market in Holmskij, Krasnodarskiy Kray, Russia (southern Russia, Black Sea region) (Capik et al., 2013; Molnar et al., 2007). The initial EFB resistance screening and identification of H3R07P25 was described in detail in Molnar et al. (2007). H3R07P25 was one of only 13 trees of 605 described in Molnar et al. (2007) that remained free of EFB. In addition, the original H3R07P25, as well as three grafted trees of the same clone, have been continually exposed to EFB in additional field trials at Rutgers, where all have remained disease free as of Dec. 2018 (Capik et al., 2013; Leadbetter et al., 2016; T.J. Molnar, unpublished data). H3R07P25 also remained free of EFB after greenhouse exposure in Oregon as well as in the field (S.A. Mehlenbacher, unpublished data). The male parent, OSU 1155.009, was identified as EFB-susceptible at OSU (data not shown) and selected as a parent for the cross based on its susceptibility, improved nut and kernel characteristics, and high yields. It originated from a cross of OSU 474.084 (‘Lewis’ × ‘Tonda di Giffoni’) × OSU 540.084 (a full sibling of ‘Sacajawea’) (Mehlenbacher et al., 2000, 2008).

MAPping population development and field evaluation of EFB response. The controlled hybridization (H3R07P25 × OSU 1155.009) was conducted following protocols described in Mehlenbacher (1994). H3R07P25 was located at Rutgers University Horticultural Research Farm #3 (East Brunswick, NJ) and pollen of OSU 1155.009 was provided by OSU. Pollen was collected in Jan. 2011 and stored at −28.9 °C until shipping on dry ice by overnight mail before its use in Feb. 2011 at Rutgers University. In Sept. 2011, the resulting hybrid seeds were harvested, provided a moist chilling period at 4 °C for 4 months, then germinated and grown in the greenhouse according to protocols described by Molnar and Capik (2012). Seedlings were removed from the greenhouse in June 2012 for acclimation outdoors under shade cloth (40% shade) until field planting in Nov. 2012 at Rutgers University Horticultural Research Farm #3. Tree spacing was 1.0 m in the row by 3.0 m between rows. Weed control using herbicides, irrigation, and fertilizer application were provided as needed, with no use of insecticides or fungicides. Trees were field inoculated by tying diseased twigs to the stems of the seedlings before budbreak each year (Molnar et al., 2007) to keep disease pressure high and reduce the possibility of susceptible plants escaping infection. In addition, EFB infected trees in nearby research plots contributed a steady influx of additional EFB inoculum. All of the trees were evaluated in Jan. 2016 and Jan. 2017 for response to EFB, which was recorded using a scale of 0 to 5, according to a modified index adapted from Pinkerton et al. (1992): 0 = no detectable EFB (includes presence of “sunken lesion” phenotypes where a few small sunken cankers that lack fungal stroma) = 0% of stems diseased; 1 = single canker (with fully formed stroma) = 1% of stems diseased; 2 = multiple cankers on a single branch = 5% of stems diseased; 3 = multiple branches with cankers = 25% of stems diseased; 4 = 50% branches with cankers = 50% of stems diseased; 5 = all branches contain cankers (except basal sprouts) = 100% of stems diseased. Plants scored 0 or 1 were considered resistant to infection by A. anomala. The 0 to 5 scale was converted to percent disease, as described previously, for the QTL analyses.

DNA extraction. Genomic DNA was isolated from both parents and 119 seedling samples using a DNeasy Plant Mini Kit (Qiagen, Germantown, MD), following the manufacturer’s instructions. Sample DNA quality and quantity were assessed using a spectrophotometer (NanoDrop; Thermo Fisher Scientific, Waltham, MA). Extracted DNA was used for SSR marker characterization and the preparation of ddRADseq libraries for all samples.

MICROSATELLITE (SSR) MARKERS. A total of 126 microsatellite or SSR markers from various sources (Akin et al., 2016; Bassil et al., 2005; Bhattarai and Mehlenbacher, 2018; Boccacci et al., 2005; Colburn et al., 2015, 2017; Gürcan and Mehlenbacher, 2010; Gürcan et al., 2010a, 2010b; Ives et al., 2014; Mehlenbacher et al., 2006; Sathuvalli and Mehlenbacher, 2013; Sathuvalli et al., 2012) were genotyped as potential anchor markers for LG comparison between the current genetic linkage map and previous maps (Beltramo et al., 2016; Bhattarai and Mehlenbacher, 2017; Colburn et al., 2015, 2017; Gürcan et al., 2010a; Ives et al., 2014; Mehlenbacher et al., 2006; Sathuvalli et al., 2011a, 2011b, 2012; Torrello Marinoni et al., 2018). A prefix indicated that LG was added to the SSR marker names. Polymerase chain reaction (PCR) amplification of SSR markers for genotyping was conducted in 13-μL reactions, using ≈5 ng genomic DNA per sample, 1 × PCR buffer, 2.0 mM MgCl₂, 0.25 mM dNTPs, 0.5 pmol forward primer with 5′-M13(-21) addition (Schuelke, 2000), 1 pmol reverse primer with 5′-‘PIG-tailing’ addition (Brownstein et al., 1996), 1 pmol forward M13(-21) primer with FAM, NED, PET, or VIC fluorescent labels (Schuelke, 2000), and 0.5 U IMMOLASE DNA polymerase (Bioline; Meridian Life Science, Memphis, TN). All SSR marker primers were synthesized by Integrated DNA Technologies (Coralville, IA). Thermal-cycling (9700 GeneAmp PCR system; Applied Biosystems,
libraries were then quantified using a fluorometer (Qubit 3.0; ThermoFisher Scientific) and Genemapper 5.0 software (Applied Biosystems), and sized using LIZ 600 size standard v2.0 (Applied Biosystems) and Genemapper 5.0 software (Applied Biosystems).

**ddRADseq library construction and Illumina sequencing.** ddRADseq libraries for H3R07P25, OSU 1155.009, and 119 mapping population seedlings were created using a protocol adapted from Poland et al. (2012). Briefly, 200 ng of genomic DNA from each individual sample was double-digested with the rare-cutting *PstI* (NEB, Ipswich, MA) and the common-cutting *MspI* restriction enzymes for 2 h at 37 °C. Uniquely barcoded [5–10 base pair (bp)] forward *PstI* adapters and reverse *MspI* Y-adapters were ligated to the digested european hazelnut DNA in a mastermix consisting of 200 U of T4 DNA ligase, 2 μL of 10 × NEBuffer 4, and 4 μL of ATP (10 mM) (NEB, Ipswich, MA) per sample. The ligation reaction was incubated for 2 h at 22 °C, with ligase inactivation occurring via an additional incubation of 20 min at 65 °C. Samples were then subjected to a “clean-up” with 0.5 v/v magnetic beads (Agencourt Ampure XP; Beckman Coulter, Brea, CA), followed by washing with 70% ethanol to remove DNA fragments smaller than 300 bp. Individual cleaned library samples were then PCR amplified, with primers that included sequences allowing for Illumina (San Diego, CA) next-generation sequencing (NGS) flow cell binding, under the following thermocycling conditions: initial denaturation of 95 °C for 30 s; followed by 16 cycles of 95 °C for 30 s, 62 °C for 20 s, 68 °C for 15 s; with a final extension of 68 °C for 5 min. All DNA libraries were then quantified using a fluorometer (Qubit 3.0; Thermo Fisher Scientific), normalized to 5 ng·μL⁻¹, and then pooled for Illumina short-read sequencing. Before sequencing, pooled libraries were subjected to a second clean-up with magnetic beads and wash step as previously described. Final pooled library quality was assessed via high-resolution automated electrophoresis (2100 Bioanalyzer System; Agilent Technologies, Santa Clara, CA).

Sequencing pools were constructed such that all progeny samples were divided equally across six MiSeq runs (Illumina) (2 × 300 paired-end) conducted at Rutgers University (New Brunswick, NJ), as well as an additional single pool of all samples on a single lane of a HiSeq 2500 [Illumina (2 × 125 paired-end high-output sequencing)] sequenced by Genewiz, Inc. (South Plainfield, NJ). Each mapping population parent sample was sequenced a total of 14 times across all sequencing runs. Each sequencing run was loaded with 10% and 30% PhiX for the MiSeq and HiSeq sequencing runs, respectively.

**ddRADseq single nucleotide polymorphism (SNP) marker calling.** Raw ddRADseq FASTQ data files were processed using the software pipeline Stacks 1.30 (Catchen et al., 2011, 2013). All data files were demultiplexed according to barcode, quality filtered, and trimmed to 100 bp using the *process_radtags* command (with barcode rescue, −r). The denovo_map.pl command wrapper of Stacks was then used for de novo SNP marker discovery (without reference genome), specifying the “CP” genetic map (−A). denovo_map.pl combines the *ustacks*, *cstacks*, and *stacks* programs. Briefly, the *ustacks* program was used to align matching reads of all samples into putative loci (stacks) and for making SNP marker calls, with a minimum of four matching reads (−m) required to create a stack and a maximum distance (nucleotide mismatch, −M) of three. A catalog of loci was then generated from the two parental genotypes using the *cstacks* program, followed by matching the generated stacks of the progeny samples back to the parental catalog using the *stacks* program. Finally, the genotypes program of Stacks was used to export map-specific genotypes in a format compatible with JoinMap 4.1 (van Ooijen, 2006) and OneMap/R (Margarido et al., 2007). Loci with more than 4.2% missing data (i.e., loci must be present in 115 progeny) were excluded from further analysis (−r of genotypes program).

**Linkage map construction.** Three approaches were used for the construction of genetic linkage maps using the software JoinMap 4.1 and the package OneMap/R. The first strategy used the two-way pseudo-testcross approach for cross-pollinated (CP) species (Grattapaglia and Sederoff, 1994) in JoinMap 4.1, with additional procedures as described by Mehlenbacher et al. (2006). Briefly, both SSR and SNP markers were initially combined into one dataset and formatted so that markers were coded as heterozygous in one parent and homozygous in the other parent (*lm xx ll* or *nn xx np*). All markers were then recoded as BC1 markers in JoinMap 4.1, and initial maps for each parent were generated. Markers assumed to be linked in repulsion, and not initially mapped, were recoded as “dummy variables” (a for present, and h for absent) to allow for mapping with markers previously assumed to be in coupling phase to create a single map for each parent. All individual markers were subjected to χ² analysis in JoinMap to test for goodness of fit to expected segregation ratios (1:1), and markers showing severe segregation distortion (*P ≤ 0.001*) were excluded from further analyses. Marker grouping was performed over a range of logarithm of odds (LOD) values from 2 to 30, with a step of one. After grouping, the order of loci was determined using the multipoint maximum likelihood mapping algorithm with a Gibbs sampling procedure (van Ooijen, 2006). This algorithm specifically deals with deterioration in computation time of the regression mapping algorithm in high-density linkage maps with greater than an average of 50 loci per LG (van Ooijen, 2006). Spatial sampling was conducted with five threshold values (0.1, 0.05, 0.03, 0.02, and 0.01), with three map optimization runs for each spatial sampling. Map optimization parameters included the following: chain length of 1000; cooling control parameter of 0.001; and chain termination after 10,000 chains without improvement. Map distances were converted from recombination frequency to centimorgans using the Kosambi mapping function.

The second strategy, also in JoinMap 4.1, used the extension of the multipoint maximum likelihood mapping algorithm to CP full-sib populations described in van Ooijen (2011a). In this extension, the JoinMap 4.1 software first constructs two parental maps simultaneously, estimating phasing, and then integrates the two parental maps into a single single map (van Ooijen, 2011b). For this strategy, SSR and SNP markers were combined into one dataset, with both common markers (*ab cd*, *ef eg*, and *hk hk*) and parental markers (*lm ll* and *nn np*) represented. Marker grouping was initially performed over a range of independence LOD values from 2 to 30, with a step of one; with a final LOD value ≥24 used to group loci. After grouping, the multipoint maximum likelihood mapping algorithm was used to determine the order of loci, with the same
parameters described previously. The Kosambi mapping function was used to determine map distances (centimorgans), also as described previously.

The third strategy was implemented in the package OneMap/R. The core functionality of this package is the construction of integrated linkage maps combining information from markers exhibiting mixed segregation patterns (common and parental markers), as well as simultaneous maximum likelihood estimation of linkage and linkage phases between markers in outcrossing (CP) species (Margarido et al., 2007). Briefly, locus genotypes used in the previous JoinMap analyses were imported into OneMap/R in outcross format using the read.outcross function. Next, the recombination fractions among all pairs of markers was calculated using the rf.2pts function. Markers were assigned to LGs using the group function, testing a range of LOD scores (5–30) and a range of maximum recombination frequencies (0.2–0.5). A final LOD score of LOD = 20 was used to create all LGs. The ordering of markers within each group was determined using the order.seq function, with an initial comparison of six informative markers, addition of markers sequentially, and an LOD threshold of three. All LG map distances (centimorgans) were calculated using the Kosambi mapping function. Alternative marker orders on all LGs were tested with the ripple.seq function. Maps from all three genetic mapping approaches described previously were viewed using MapChart 2.30 (Voorips, 2002).

**QTL analysis.** QTL analysis of single year (2016 and 2017) and mean EFB response was conducted using MapQTL 6.0 software (van Ooijen, 2009). Because of memory constraints with the CP mapping population, the two-way pseudo-testcross approach with markers recoded as DH population type as described by van Ooijen (2009) was used to construct two separate parental maps that were then used for all QTL mapping procedures (Herrmann et al., 2006; McAdam et al., 2013; Studer et al., 2006; van Heerden et al., 2014; Yun et al., 2014; Zyprian et al., 2016). Initial analysis was conducted using interval mapping (IM), followed by multiple rounds of multiple QTL mapping (MQM) to refine the location and magnitude of QTL. Briefly, putative QTL identified from IM were used as initial cofactors in the MQM mapping analysis. For MQM mapping, a backward elimination procedure was then used to select additional cofactors in subsequent rounds of MQM, until QTL positions were stabilized. LOD thresholds for QTL significance were determined by permutation test \((n = 10,000)\), with a genome-wide significance level of 0.05. QTL positions and magnitude were visualized using MapChart 2.30 (Voorips, 2002).

**Results**

**Segregation for EFB resistance.** The EFB response of the 119 seedlings from the cross between H3R07P25 and OSU 1155.009 was 55 resistant and 64 susceptible (Table 1). EFB ratings were similar across years (data not shown), and were therefore averaged for \(\chi^2\) analysis. The segregation pattern fits the expected ratio of one resistant to one susceptible, indicating that EFB resistance from H3R07P25 may be controlled by a dominant allele at a single locus in a heterozygous state.

**Microsatellite (SSR) genotyping.** Ninety SSR markers previously used for genotyping hazelnuts and linkage mapping (Beltramo et al., 2016; Bhattacharai and Mehlenbacher, 2017; Colburn et al., 2015, 2017; Gürçan et al., 2010a; Ives et al., 2014; Mehlenbacher et al., 2006; Sathuvalli et al., 2011a, 2011b, 2012; Torello Marinoni et al., 2018) were found to be polymorphic between the two parents of the mapping population cross and are listed along with their motif type and primer sequence information (Supplemental Table 1). Six of the 90 SSR markers were removed from the final genetic linkage map because of greater than 4.2% missing data in the progeny population or suspect linkages. Thirty-one of the remaining SSR markers segregated 1:1 \((ln \times ll \text{ or } nn \times np)\); 50 SSR markers segregated 1:1:1 \((ab \times cd \text{ or } ef \times eg);\) whereas three SSR markers segregated 1:2:1 \((hk \times hh)\) (Supplemental Table 1).

**SNP marker discovery and polymorphic loci development.** The summary data for the sequencing runs performed in this study are presented in Supplemental Table 2. Each ddRADseq sequencing library was sequenced individually in a single lane of either an MiSeq or HiSeq instrument platform. The mean Q scores for the MiSeq sequencing runs ranged from 32.4 to 34.3, with an average mean Q score of 33.7 across all six MiSeq sequencing runs. The mean Q score for the single HiSeq 2500 sequencing run was 29.0. The percentage of Q score \(\geq 30\) for the MiSeq sequencing runs ranged from 76.9% to 84.9%, with an average percentage of Q score \(\geq 30\) of 82.3% for all six MiSeq sequencing runs, whereas the single HiSeq 2500 sequencing read had a percentage of Q score \(\geq 30\) of 63.5%.

The combined MiSeq and HiSeq 2500 sequencing runs yielded a total of 101.11 Gb of DNA sequences (Supplemental Table 2). After demultiplexing and trimming to remove barcodes, adapters, and low-quality bases (at sequence fragment ends) (Stacks process_radtags), the number of sequencing reads per seedling sample ranged from a low of slightly more than 1.83 million reads to a high of slightly more than 4.06 million reads, with an average of 2.89 million reads across all 119 samples. The parental samples had slightly more than 14.03 million reads, with an average of 2.89 million reads across all 119 samples. The parental samples had slightly more than 14.03 million and 15.76 million reads for OSU 1155.009 and H3R07P25, respectively. This resulted in an average of \(5.2\times\) number of reads for the parental samples relative to the seedling samples, where a substantial parental read number excess is important for generating accurate SNP marker calls in the Stacks software pipeline. Raw reads for the two parental samples and all 119 seedling samples were deposited in the National Center for Biotechnology Information Sequence Read Archive database (PRJNA542160) (Leinonen et al., 2011).

The Stacks software retained read catalog for the mapping population parents and progeny (after the usstacks, cstacks, and stacks steps) contained an average of 110,233 stacks, 16,741 polymorphic loci, and 25,864 raw SNP marker calls. After

| Parents | Seedlings (no.) | Test ratio | Value | \(P\) |
|---------|----------------|------------|-------|------|
| H3R07P25 × OSU 1155.009 | Resistant 55 | Susceptible 64 | 1:1 | 0.54 | 0.47 |

\(^{a}\)Plants with an average rating of 0 to 1.5 for the combined 2016 and 2017 seasons were considered resistant to EFB. Plants with an average rating score of 3.5 to 5.0 were considered susceptible. There were no plants with an average rating score of 1.6 to 3.4. Rating scale adapted from Pinkerton et al. (1992).
further filtering for progeny individuals containing <4.2% missing data, 2894 SNP markers were retained. An additional 761 SNP markers were removed from the dataset before final linkage grouping were due to duplicated loci, suspect linkages, failure to group during linkage mapping, and severe segregation distortion ($P \leq 0.001$). A total of 1927 of the remaining 2133 SNP markers segregated 1:1 ($lm \times ll$ or $nn \times np$) in the progeny population; 18 SNP markers segregated 1:1:1 ($ab \times cd$ or $ef \times eg$); whereas 188 SNP markers segregated 1:2:1 ($hk \times hk$) (Supplemental Table 3).

**LINKAGE MAP CONSTRUCTION.** The three previously described approaches used for linkage mapping (two-way pseudo-testcross in JoinMap 4.1; integrated linkage map in JoinMap 4.1; and integrated map in OneMap/R) of the 2133 SNP markers and 84 SSR markers produced identical results for the grouping analyses, with only very minor differences in microcolinearity between marker positions. Therefore, only the integrated linkage map from JoinMap 4.1 is presented in Fig. 1 and Supplemental Figs. 1–11. At LOD ≥24, JoinMap 4.1 returned 11 integrated LGs, corresponding to the expected haploid chromosome number for European hazelnut ($n = x = 11$) (Fig. 1, Supplemental Figs. 1–11) (Mehlenbacher et al., 2006). The map included a total of 2217 markers and spanned a total genetic distance of 1383.4 cM with an average marker spacing of 0.65 cM. The genetic distance of individual LGs ranged from a low of 70.5 cM (98 markers) for LG3, to a high of 196.0 cM (269 markers) for LG2. LG assignments were made via syntenic comparisons of SSR anchor markers to previous European hazelnut linkage maps (Beltramo et al., 2016) (also LG4). The 80 remaining SSR markers were then used as anchor markers for LG assignment, with a high of 11 anchored SSR markers on LG9, a low of five anchored SSR markers on LG10, and an average of 7.3 anchored SSR markers per LG (Supplemental Table 1, Supplemental Figs. 1–11).

**QTL ANALYSIS.** The initial IM QTL analysis in MapQTL 6.0 identified three potential marker/trait (EFB response) association regions on LG2, LG4, and LG9, all associated with the EFB-resistant H3R07P25 parental map [using the DH two-way pseudo-testcross approach for QTL mapping (van Ooijen, 2009)]. Further refinement of QTL location using cofactor selection and MQM mapping resulted in identification of a single EFB resistance QTL region on LG2 associated with two ddRADseq SNP markers (RUCaS1_18603 and RUCaS1_14004) (Fig. 2). QTL LOD scores for RUCaS1_18603 and RUCaS1_14004 were 32.5 and 34.8, respectively (Fig. 2), whereas the average percent phenotypic variation (EFB disease response) explained by the QTL region spanning the two SNP markers was 72.8%, indicating a major QTL region associated with EFB disease resistance. The initial IM marker/trait association regions on LG4 and LG9 fell far below the permutation test–calculated LOD threshold of 3.0 during the cofactor selection and MQM mapping analyses steps. BLASTN results for the sequences containing the RUCaS1_18603 and RUCaS1_14004 SNP markers (Supplemental Table 3) returned no matches (data not shown).

**Discussion**

Before the advent of NGS technologies, genetic linkage mapping and QTL identification experiments were often conducted with DNA markers such as restriction fragment length polymorphism markers, random amplified polymorphic DNA (RAPD) markers, amplified fragment length polymorphism markers, microsatellite or SSR markers, or some combination thereof. Although many of these markers proved valuable (and are still valuable for anchoring between linkage maps), the high costs, low throughput, and laborious nature of working with these marker systems resulted in low coverage linkage maps requiring an extensive investment of time and resources to...
answer a broader range of research questions than have genetic linkage maps and identify marker trait associations to lower cost methodology can then be used to quickly generate reproducible, and economic means of generating thousands to systems, are a relatively straightforward, highly specific, highly techniques, such as genotyping-by-sequencing or ddRADseq, that construct. In contrast, reduced representation sequencing tech-

Fig. 2. The quantitative trait locus (QTL) on the genetic linkage map of the eastern filbert blight (EFB) disease resistance QTL and single nucleotide polymorphism (SNP) markers RUCA_S1_14004 and RUCA_S1_18603 are located on linkage group (LG) 2. Logarithm of odds (LOD) thresholds for QTL significance (dashed line) were determined by permutation test (n = 1000), with a genome-wide significance level of 0.05. QTL LOD scores (solid line) for RUCA_S1_18603 and RUCA_S1_14004 were 32.5 and 34.8, respectively. The average percent phenotypic variation (EFB disease response) explained by the QTL region spanning the two SNP markers was 72.8%.

The putative ‘Gasaway’ R-gene (as well as resistance from ‘Culplá’, ‘Crvenje’, ‘Uebov’, OSU 408.040, and OSU 495.072) was previously shown to reside on European hazelnut LG6 (Colburn et al., 2015, 2017; Gürcan and Mehlhenbacher, 2010; Mehlhenbacher et al., 2006; Sathuvalli and Mehlhenbacher, 2013; Sathuvalli et al., 2012). Interestingly, EFB resistance from H3R07P25 resides on LG2 (Fig. 1, Supplemental Fig. 2). An important finding from this work is that the resistance source from H3R07P25 is different from the ‘Gasaway’ resistance source (previously found to reside on LG6). This is based on the differences in LG locations of resistance QTL (and lack of a resistance signal on LG6 in the current work), as well as the phenotype of the plants protected by this source of resistance in a high disease pressure environment within the native range of the fungus. The position of the disease resistance locus on LG2 in H3R07P25 appears to be similar to the disease resistance source identified in OSU 759.010 (selection from the Republic of Georgia), as both QTL locations are distal to the SSR marker CAT_B501 on LG2 (Supplemental Fig. 2) (Sathuvalli et al., 2011b). Additional work will be required to determine if these resistance genes are identical or allelic at the same locus, if they differ and are part of a cluster of disease resistance genes in that region, or if they differ in some other means from one another. Interestingly, the phenotypic response of OSU 759.010 in New Jersey differs from H3R07P25. In Molnar et al. (2010a), replications of an identical clone (OSU 759.007) expressed EFB after inoculation with most isolates in the study. In the field, Capik and Molnar (2012) also showed that it developed EFB; however, only four of six trees of OSU 759.007 developed EFB and their average proportion of diseased wood after 10 years of exposure was less than 1% of the tree canopy, which would qualify as resistant in this current study. Additional work with OSU 759.007 is currently under way to evaluate the similarity of these two resistance sources.

The findings from the current research, in combination with those from previous studies, have numerous important implications for European hazelnut breeders and growers. First, previous studies have shown differential EFB disease responses in the field between the ‘Gasaway’ resistance source and H3R07P25, different LG locations of resistance sources (LG6 vs. LG2, respectively), various pathogenic responses from different A. anomala isolates (Molnar et al., 2010a), and a high degree of fungal genetic diversity in EFB high disease pressure environments (Cai et al., 2013; Muehlbauer et al., 2014b, 2019; Tobia et al., 2017). These observations indicate that different isolates of A. anomala in high disease pressure environments may be eliciting differential pathogenic responses from European hazelnut plant material carrying different disease resistance genes. Although this was hypothesized in previous research (Capik and Molnar, 2012; Leadbetter et al., 2016; Molnar et al., 2010a, 2010b; Muehlbauer et al., 2014a; Sathuvalli et al., 2011a, 2011b), the current work further emphasizes the need for breeders to develop European hazelnut germplasm exhibiting EFB resistance via R-gene pyramiding approaches or through the identification of R-genes conferring resistance to multiple isolates of A. anomala (as indicated in the current work). Second, screening European hazelnut germplasm and developing new cultivars exhibiting EFB resistance in a high disease pressure environment with greater fungal genetic diversity may allow for the identification of germplasm capable of maintaining resistance against a greater number of fungal isolates. Third, the resistance conferred by the ‘Gasaway’ source is likely different from that identified in H3R07P25, does not appear to confer complete resistance to EFB in all high disease pressure environments, and therefore may confer resistance to only a limited number of A. anomala isolates. Accordingly, continuing efforts are needed to prevent the introduction of additional isolates of A. anomala into the
PNW. In addition, should that effort fail, European hazelnut breeders need to continue to search for and identify new sources of EFB resistance that offer broader protection against genetically diverse EFB fungal populations. Fourth, identification of European hazelnut germplasm that maintains EFB resistance in high disease pressure environments is a requirement for the development of a commercially viable European hazelnut industry in the eastern United States. Identification of the LG location of that resistance, and transmission to progeny as a single locus with a dominant allele for resistance, will likely expedite that development. Last, the identification of genetic markers tightly linked to EFB resistance, as well as previously discovered linked genetic markers, will help facilitate resistance gene pyramiding approaches (e.g., ‘Gasaway’-based resistance from LG6, ‘Rotoli’-based resistance from LG7, and H3R07P25-based resistance from LG2 in the current work) via marker-aided breeding, and/or the identification of EFB resistance genes that confer resistance to multiple *A. anomala* isolates.

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Supplemental Fig. 1. Representation of European hazelnut linkage group (LG) 1. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 2. Representation of European hazelnut linkage group (LG) 2. The eastern filbert blight (EFB) disease resistance quantitative trait loci (QTL) region is denoted by “+++.” Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 3. Representation of European hazelnut linkage group (LG) 3. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 4. Representation of European hazelnut linkage group (LG) 4. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 5. Representation of European hazelnut linkage group (LG) 5. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 6. Representation of European hazelnut linkage group (LG) 6. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 7. Representation of European hazelnut linkage group (LG) 7. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 8. Representation of European hazelnut linkage group (LG) 8. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.

LG8 [1]

RLCaS1_18407
RLCaS1_1999 RUCaS1_91481
RLCaS1_23914
RLCaS1_71617
RLCaS1_45571 RUCaS1_73393
RLCaS1_18382
RLCaS1_8962
RLCaS1_77327
RLCaS1_9165 RUCaS1_24933
RLCaS1_5762 RUCaS1_89457
RLCaS1_89003

LG8 [2]

RLCaS1_12955
RLCaS1_91136
RLCaS1_33245
RLCaS1_6154
RLCaS1_13309
RLCaS1_94835
RLCaS1_13206 RUCaS1_9098
RLCaS1_58070
RLCaS1_87469
RLCaS1_5349 RUCaS1_91769

A616 LG8

RLCaS1_63525
RLCaS1_18293
RLCaS1_17548
RLCaS1_92270 B617_LG8
RLCaS1_128800 RUCaS1_18355
RLCaS1_65186
RLCaS1_8671 RUCaS1_74367
RLCaS1_9816 RUCaS1_17567
RLCaS1_49145 RUCaS1_17647
RLCaS1_39569 RUCaS1_34004
RLCaS1_62590 RUCaS1_66060
RLCaS1_91390
RLCaS1_4876 RUCaS1_75394
RLCaS1_61918 RUCaS1_36188

B788 LG8

RLCaS1_50634
RLCaS1_18327 RUCaS1_85041
RLCaS1_79974
RLCaS1_24479
RLCaS1_35281
RLCaS1_17652 RUCaS1_79233
RLCaS1_19801
RLCaS1_43086
RLCaS1_63680

B726 LG8

RLCaS1_10880

B654 LG8

RLCaS1_18733 RUCaS1_49000
RLCaS1_49609 RUCaS1_64657
RLCaS1_18144 RUCaS1_60270
RLCaS1_23600
RLCaS1_54270
RLCaS1_68401 RUCaS1_62576
RLCaS1_90635
RLCaS1_75767 RUCaS1_35721
Supplemental Fig. 9. Representation of European hazelnut linkage group (LG) 9. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 10. Representation of European hazelnut linkage group (LG) 10. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.
Supplemental Fig. 11. Representation of European hazelnut linkage group (LG) 11. Microsatellite or simple sequence repeat (SSR) anchor markers are highlighted in bold italics.