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<u>ARTICLE</u>

Discovery of a resistance gene cluster associated with smut resistance in peanut

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ABSTRACT

Peanut smut, caused by Thecaphora frezzii Carranza & J.C. Lindq., is an emerging threat for the global peanut industry. The disease's destructive potential can be exemplified by pod incidence values as high as 70% and yield losses reaching 30%. Because fungicides have shown moderate but highly variable levels of control levels, development and deployment of smut resistant cultivars are the best strategies for disease management. Screening for smut-resistant germplasm requires years of field trials and is currently the only option for breeders because genetic markers for resistance have not yet been developed. The objectives of this study were to perform whole genome sequencing (WGS) on populations developed for smut resistance mapping and subsequently fine map discovered QTL associated with smut resistance. An expedited strategy was employed by phenotyping in the F3 generation. We phenotyped 200 families with 3 plants per family for smut resistance in infested fields during the 2019/2020 season in General Deheza (32°45'23"S 63°47'20"W), Argentina. Each individual was sequenced using iGenomXRipTide library preparation and Illumina NovaSeq sequencing to yield approximately 1 times genome coverage. Analysis of the phenotype and genotype data using Khufu resulted in the identification of a single major smut resistance QTL on chromosome 12 (B02). Chromosome level genome sequences were assembled for the resistant parent (Ascasubi) and susceptible parent (Granoleico) using PacBio HIFI sequencing. A validation population was sequenced using Khufu to validate the QTL region and analysis confirmed the major locus on chromosome 12. The identified variation will be used to develop smut resistant varieties quickly using molecular assisted breeding strategies.

INTRODUCTION

Peanut smut, caused by Thecaphora frezzii Carranza & J.C. Lindq. (T. frezzii), is an emerging threat for the global peanut industry (BICON, 2017; Rago et al., 2017). First described in 1962, T. frezzii was observed on a collected sample of a wild peanut from Mato Grosso do Sul, Brazil, that was sent to INTA's Manfredi Exp. Stn. in Córdoba Province, Argentina (Carranza & Lindquist 1962). In 1995, peanut smut was first detected on cultivated peanuts on three farms in north-central Córdoba (Marinelli et al., 2008). Since then, it has spread to all peanut-producing provinces in Argentina (Cazón et al., 2018). The disease's destructive potential can be exemplified by pod incidence values as high as 70% (Bonessi et al. 2011, Chamberlin et al. 2022) and yield losses reaching 30% (Oddino et al. 2010). Disease management strategies using pesticides or biocontrol agents (Figueredo et al. 2017), liming (Bonessi et al. 2011), tillage (Cignetti et al. 2010; Cazón et al. 2014), and rotation (Marraro Acuña and Haro, 2011) have been attempted with modest results. Fungicides have shown moderate but highly variable control levels (Rago et al. 2017; Paredes et al. 2021). Smut resistant cultivars seem the best tool for disease management as they suffer less crop loss and limit the pathogen's multiplication (Rago et al. 2017). Sources of smut resistance have been reported in wild peanuts (de Blas et al. 2019), synthetic amphidiploids (de Blas et al. 2021) and in A. hypogaea core collections (Chamberlin et al. 2022; Wann et al. 2020). Smut genetic resistance has shown high broad sense heritability (Bressano et al. 2019; de Blas et al. 2021). Recently, major and a minor quantitative trait loci (QTL) associated with moderate resistance have been identified in a recombinant inbred line (RIL) population developed from a cross between a susceptible A. hypogaea genotype and a resistant synthetic amphidiploid [A. correntina × A. cardenasii] × A. batizocoi (de Blas et al. 2021). Peanut reaction to T. frezzii is assessed by opening each pod by hand and inspecting it visually for incidence (i.e., the presence or absence of *T. frezzii* in pods), which is time and labor intensive. Screening for smut resistant genetic sources requires years of in-field trials along with pod phenotyping and is currently the only option for breeders because markers for resistance have not yet been developed.

Marker assisted selection (MAS) is now widely used in plant breeding to increase the efficiency of developing cultivars with highly desired traits. Early marker work identified several for disease resistance in peanut. Random amplified polymorphic DNA (RAPD) was used to identify root-knot nematode (RKN) resistance (Burow et al., 1996), restriction fragment length polymorphisms (RFLPs) were identified for resistance to Meloidogyne arenaria (Church et al., 2000), amplified fragment length polymorphisms (AFLPs) were found linked to resistance to the vector of groundnut rosette virus (Herselman et al., 2004), and single sequence repeats (SSRs) associated with resistance to Sclerotinia minor (Jagger) (Chenault, et al., 2009) along with those for tomato spotted wilt virus (TSWV) and leaf spot (LS) (Wang et al., 2013; Tseng et al., 2016; Zhao et al, 2018), rust (Khedikar et al., 2010; Mondal and Badigannavar, 2018), and bud necrosis disease (Jadhav et al., 2019) were identified.

Sequencing of the diploid progenitors (Bertioli *et al*, 2016), *A duranensis* and *A. ipaensis*, provided the reference

genomes necessary to construct the high-density single polymorphism (SNP) genotyping nucleotide arrav Axiom_Arachis (Pandey et al., 2017) which led to the generation of SNP-based linkage maps enabling the identification SNPs associated with traits. SNPs are preferred over other marker types due to their wide distribution throughout the peanut genome (Liao and Lee, 2010). Whole genome resequencing (WGRS) of mapping populations has also been used to generate high-density SNP-based maps and identify QTL along with candidate genes for resistance to peanut diseases such as early LS (ELS), late LS (LLS), and spotted wilt (Agarwal et al., 2018). Such linkage maps have also been used to identify QTL for resistance in peanut to RKN (Leal-Bertioli et al., 2016), Aspergillus flavus (Khan et al., 2020), bacterial wilt (Luo et al, 2020a), stem rot (Luo et al. 2020b), Sclerotinia blight (Liang et al., 2021), and bud necrosis disease (Jasani et al., 2021). The recent release of the tetraploid (cultivated) peanut genome sequence (Bertioli et al., 2019) has further clarified questions regarding A. hypogaea evolution and the generation of a high-density SNP array for tetraploid peanut (Axiom_Arachis2) (Clevenger et al., 2017, Clevenger et al., 2018) provide further foundation for rapid marker-trait association. Using the Axiom Arachis SNP array, potential SNPs and QTL associated with peanut smut resistance were identified by association mapping using a limited number of genotypes (Massa et al., 2021), but no confirmed QTL or candidate genes have been reported to date. Therefore, the objectives of this study were to perform whole genome sequencing (WGS) on a population developed for smut resistance mapping and subsequently fine map discovered QTL associated with smut resistance.

MATERIALS AND METHODS

Plant Materials and Population Development.

The mapping population consisting of 200 lines was developed by crossing Mf10_2870, smut susceptible (J. Baldessari, unpublished data), and Ascasubi Hispano, smut resistant (Ibañez et al., 2018). An additional population used for validation purposes was also developed by crossing Granoleico, smut susceptible (Soave, 2002; Oddino et al., 2013) and Ascasubi Hispano. To rapidly identify QTL associated with smut resistance, a non-traditional approach was used where families within the population were phenotyped and genotyped at familial stages prior to the F7generation rather than developing a uniform recombinant inbred line (RIL) population. Briefly, three individuals from each F2:3 family (for the mapping population) and from each F_{3:4} family (for the validation population) were phenotyped under heavily smut infested field conditions. This was done during the 2019-2020 season for the mapping population and during the 2020-2021 season for the validation population, in General Deheza, Córdoba Province, Argentina under rainfed conditions.

Experimental Design and Field Testing.

Smut resistance tests were conducted as described in Chamberlin *et al.*, 2022, in fields adjacent to a farm (-32.759, -63.770) in the town of General Deheza (Córdoba Province, Argentina). Soil at the site was a General Deheza coarse-silty sandy loam type (0.3% slope, coarse-silty, mixed, thermic, Entic Haplustoll). The tests were planted one km downwind from a peanut processing plant where peanut smut is prevalent. The production field site was on a rotation schedule of corn-cornsoybean-peanut, where the preceding crop of the test site was always soybean.

Two control entries, one resistant (Ascasubi Hispano) and one susceptible (Colorado Irradiado INTA) were planted in both seasons. They were replicated twelve times each. Each family was represented by three individuals, each one constituting a plot. Plots were arranged in a square grid and randomized within it. Controls were placed, alternately, along a diagonal of the grid. Individual plots were spaced one m apart among rows and between plants.

The tests were planted on 11/7/19 and 12/23/20, using an augmented grid design with three replications, using single plants as experimental units. Plants were respectively dug at 147 and 140 days after planting (DAP), and all pods from each plant were harvested by hand and placed in mesh bags to air dry for two months. Yearly soil spore count was done as described by Chamberlin *et al.* (2022) to determine disease pressure. The test was conducted following the extension guidelines of the Instituto Nacional de Tecnología Agropecuaria (INTA).

Harvest and Evaluation of Smut Resistance.

Each pod was opened by hand and rated for disease incidence, i.e., presence or absence of *T. frezzii* sori or spore masses on the kernels. Disease incidence (DI) was calculated as DI = Infected pods / Total pods.

QTL mapping and validation.

DNA was taken from 572 individuals (approximately 3 individuals from each of the 200 lines) of the $F_{2:3}$ family (Mf10_2870 and Ascasubi Hispano. DNA was also taken from 96 individual plants of $F_{2:3}$ family of Ascasubi x Granoleico selected for phenotype. DNA extraction was performed from 20 mg of dried young leaves in silica gel. The samples were ground for two 10-sec cycles in bead mill at 15.000 rpm (Super FastPrep-2 Bead Beating System, MP BiomedicalsLLC, Irvine, CA, USA). A modified CTAB method was used with a sorbitol cleaning wash before the lysis step (Inglis *et al.* 2018). DNA quality was determined using 0.8% agarose gel and the DNA quantity was estimated with DS-11+ Nano UV-Vis Spectrophotometer (DeNovix Inc., DE, USA).

Parental genomic DNA extraction.

For whole-genome sequencing, seeds of Ascasubi INTA Cv, Granoleico Cv and Mf10_2870L were germinated in an incubation oven, under total darkness at constant temperature of 28 °C for 15 days. Fresh young etiolated leaves were collected for each genotype, immediately frozen with liquid nitrogen, and preserved at -80 C (Revco, Model ULT1386-5V41). The nucleii were extracted following the Nucleii Isolation – LN2 Plant Tissue Protocol (Ciculomics Inc./ PacBio) optimized from Workman *et al.* (2018). In the step 5 of protocol, filtration of the lysate with steriflip 20 μ m pore was replaced with two lysate filtrations with a cell strainer of 100 μ m pore and 40 μ m pore. High-molecular weight (HMW) DNA was extracted from nucleii following the specifications of Nanobind Plant Nuclei Big DNA Kit (Circulomics Inc./ PacBio). DNA quality was determined using 0.8% agarose gel and the DNA quantity was estimated by Qubit V1 (Invitrogen ThermoFisher Scientific Inc.) using Qubit[®] dsDNA BR Assay Kits.

Genotyping: Affymetrix Axiom SNP Array.

Seeds from the peanut mini core accessions were sowed in 4.7m pots with a mix of 50% Promix (Premier Tech Horticulture, Quaker, PA) and 50% steam-sterilized sandy soil from the Coastal Plain Experiment Station in Tifton, GA. Young leaf tissue was collected from 1-month-old plants for DNA extraction using DNeasy Plant mini kit (www.qiagen.com). Quantification of DNA was performed with Quant-iT dsDNA assay kit (www.thermofisher.com). DNA were submitted for genotyping with the Arachis version SNP array consisting of 47K SNPs features (www.thermofisher.com). SNP data were curated from the Axiom analysis suite (www.thermofisher.com). SNP markers were classified into six categories, i.e., PolyHighResolution, NoMinorHom, MonoHighResolution, CallRateBelowThreshold, OfftargetVariant, and Other according to the SNP QC matrix of the software (Clevenger et al., 2017). All data analysis in this study was performed with markers in the PolyHighResolution category due to its high quality in signal separation.

Genotyping: PacBio Sequencing

High molecular weight DNA was received from A. hypogea Granoleico and A. hypogea Ascasubi. DNA was sheared using the Diagenode Megaruptor 3 targeting 20kb fragments. Sheared DNA was prepared for PacBio sequencing using the SMRTbell Express Template Prep Kit 2.0. The library was size selected with AMpure PB beads to remove fragments less than 3kb. Sequencing was performed on a Sequel IIe System (Pacific Biosciences, Menlo Park, CA) using Binding Kit 2.2, Sequel II Sequencing Kit 2.0, and SMRTCell 8M. To target HiFi reads, the library was sequenced using a 30-hour movie time using Instrument Control Software Version 10. Raw subreads were converted to HiFi data by processing with CCS to call a single high quality consensus sequence for each molecule, using a 99.5% consensus accuracy cutoff. A. hypogea Granoleico was sequenced on 2 SMRTcells yielding 73.11Gb of HiFi data, and A. hypogea Ascasubi was sequenced on 3 SMRTcells yielding 61.37Gb of HiFi data.

Genotyping: Khufu whole-genome sequencing and analysis

DNA from populations were prepped into sequencing libraries using iGenomX Riptide library prep. For the F2:3 population, Six Riptide libraries, representing 576 samples were sequenced on 2 lanes of NovaSeq S4 chemistry (800 Gb raw base pairs per lane) to yield an average of 0.99X genome coverage per individual. The raw reads were demultiplexed and processed using Khufu (hudsonalpha.org/khufudata). To identify QTL for resistance, bulks were identified using one year of field data. Analysis of the bulks were done in silico, by bulking individual sequenced samples and calculating allele frequency differences between the putative resistant and susceptible bulks. The F3:4 samples were processed and analyzed the same way, except were sequenced to an average of 1.14X genome coverage. Haplotype analysis was done using HawkHAP, which is a component of Khufu.

RESULTS AND DISCUSSION

Generation of reference genomes

Long read sequencing and assembly led to the generation of highly contiguous genome assemblies for smut resistant parent Ascasubi, and smut susceptible parent Granoleico (Table 1).

Mapping of smut resistance

A total of 576 individuals to construct bulks for mapping. They represented 200 F2:3 sub-families where each family was represented by 3 individuals. The smut pressure was low due to drought stress during pod development, but there were enough 'susceptible' individuals to map construct bulks (Figure 1). The frequency of infection was heavily skewed to the low end of the distribution, indicating that there was a considerable amount of false negative disease scores. To construct bulks, we employed a conservative approach whereby we selected susceptible individuals when all three sub-family members had infection and the average infection was greater than 5%. It was more difficult to select resistant bulks, because more than 400 individuals had 0% infection. For the 'resistant' bulk we selected every family member if they each had 0% infection and at least one individual was located in the field close to a susceptible check that exhibited infection. These criteria were chosen to reduce false negatives in the bulk as much as possible.

We selected 45 individuals for the 'resistant' bulk and 49 individuals for the 'susceptible' bulk. The QTL-seq analysis was conducted by bulking bam alignment files *in silico* and assaying allele frequency differences genome-wide between the bulks (Figure 2A). Analysis of the bulks indicated one major

QTL located on the proximal end of chromosome 12. There were no other peaks identified, indicating that a single, major locus accounted for smut resistance from Ascasubi. In the following year (2019-2020), we phenotyped a related population of F3:4 progeny with Ascasubi as a parent and Granoleico as the susceptible parent. We selected a set of 48 resistant individuals and the 48 most susceptible individuals. We sequenced those 96 individuals and again analyzed them with QTL-seq analysis. The major locus on chromosome 12 was again indicated as a single, major resistance locus controlling resistance (Figure 2B).

We have identified and validated a major locus controlling smut resistance in peanut. The qualitative inheritance of this trait, its high level of heritability, low environmental variance, and strong phenotyping methodologies, we were able to map and validate this resistance within early generations (F2:3 and F3:4), by phenotyping single plants rather than replicated plots. Phenotyping and genotyping early generation families within a developing population saves time and allows the potential for selection for backcross breeding. This method greatly decreased the time needed for discovery. The resistance is indicated on an approximal 2 Mb region that contains the largest cluster of R genes in the peanut genome (Bertioli et al., 2016). We have generated highly contiguous genome assemblies of the resistant parent, Ascasubi, and of the most widely grown cultivar in Argentina, Granoleico. Granoleico is very susceptible to smut, and the differences between those two genomes within the 2 Mb region will contain the functional variation that confers strong resistance to smut. This resistance and the markers associated with it will be transformational in breeding smut resistant cultivars for Argentina and the world.

Table 1. Assembly stats for the Ascasubi and Granoleico genome assemblies. Contigs were scaffolded into 20 pseudomolecules comprising the 10 A chromosomes and 10 B chromosomes. N/L50 is the length of the contig after half of the assembly has been accounted for when contigs are sorted from largest to smallest. Briefly, for Ascasubi half of the genome is assembled into 16 contigs that are longer than 53 Mb. N/L90 is calculated the same, but for 90% of the assembly. Briefly, for Ascasubi, 90% of the assembly is in 39 contigs which are longer than 19 Mb.

| | Ascasubi (R) | Granoleico (S) |
|-------------------|--------------|----------------|
| Scaffold total | 20 | 20 |
| Contig total | 92 | 97 |
| Contig N/L50 | 16/53.095 МЬ | 18/53.926 Mb |
| Contig N/L90 | 39/19.502 Mb | 39/27.963 Mb |
| Max contig length | 136.375 МЬ | 159.756 МЬ |



Figure 1: Smut infection as measured by infected pods/total pods in individual F2:3 progeny plants (A) tested at Manfredi in 2018-2019 field season (B).



Figure 2: Mapping and validation of smut resistance. (A) A major QTL was identified on chromosome 12. Bulk analysis of 45 individuals with low infection and 49 individuals with infection >3% identifies a 2 Mb region conferring resistance. Y axis is the difference in allele frequency between the two bulks. A peak indicates a region where the allele frequencies significantly segregate between the bulks. The X axis is the physical location on the chromosome. (B) Validation of the QTL on chromosome 12 in a validation population of 48 resistant individuals and 48 susceptible individuals.

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