## The *rhg1-a* (*Rhg1* low-copy) nematode resistance source harbors a copiafamily retrotransposon within the *Rhg1*-encoded $\alpha$ -SNAP gene

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## **Abstract**

Soybean growers widely use the Resistance to Heterodera glycines 1 (Rhg1) locus to reduce yield losses caused by soybean cyst nematode (SCN). Rhg1 is a tandemly repeated four gene block. Two classes of SCN resistance-conferring Rhg1 haplotypes are recognized: rhg1-a ("Peking-type", low copy number, 3 or fewer Rhg1 repeats) and rhg1-b ("PI 88788-type", high copy number, 4 or more Rhg1 repeats). The rhg1-a and rhg1-b haplotypes encode α-SNAP (alpha-Soluble NSF Attachment Protein) variants  $\alpha$ -SNAP<sub>Rhg1</sub>LC and  $\alpha$ -SNAP<sub>Rhg1</sub>HC respectively, with differing atypical C-terminal domains, that contribute to SCN-resistance. Here we report that rhg1-a soybean accessions harbor a copia retrotransposon within their Rhg1 Glyma. 18G022500 (α-SNAP-encoding) gene. We termed this retrotransposon "RAC", for Rhg1 alpha-SNAP copia. Soybean carries multiple RAC-like retrotransposon sequences. The Rhg1 RAC insertion is in the Glyma. 18G022500 genes of all true rhg1-a haplotypes we tested and was not detected in any examined rhg1-b or  $Rhg1_{WT}$  (single-copy) soybeans. RAC is an intact element residing within intron 1, anti-sense to the rhg1-a  $\alpha$ -SNAP open reading frame. RAC has intrinsic promoter activities, but overt impacts of RAC on transgenic α-SNAP<sub>RhgI</sub>LC mRNA and protein abundance were not detected. From the native rhg1-a RAC<sup>+</sup> genomic context, elevated  $\alpha$ -SNAP<sub>Rhg/</sub>LC protein abundance was observed in syncytium cells, as was previously observed for α-SNAP<sub>RhgI</sub>HC (whose rhg1-b does not carry RAC). Using a SoySNP50K SNP corresponding with RAC presence, just ~42% of USDA accessions bearing previously identified rhg1-a SoySNP50K SNP signatures harbor the RAC insertion. Subsequent analysis of several of these putative rhg1-a accessions lacking RAC revealed that none encoded  $\alpha$ -SNAP<sub>Rhg1</sub>LC, and thus they are not rhg1-a. rhg1-a haplotypes are of rising interest, with Rhg4, for combating SCN populations that exhibit increased virulence against the widely used rhg1-b resistance. The present study reveals another unexpected structural feature of many Rhg1 loci, and a selectable feature that is predictive of *rhg1-a* haplotypes.

Keywords: Plant disease resistance, retrotransposon, soybean cyst nematode, Rhg1

## Introduction

To thrive in their natural environments, organisms must continually sense and respond to changing conditions, including biotic and abiotic stresses. Transposable elements can cause relatively stable variation in numerous plant phenotypes such as flowering time, trichome presence or fruit size (Lisch, 2013). Transposable elements (TEs) may insertionally disrupt genes, or if TE activity is repressed by epigenetic transcriptional silencing, small interfering RNAs and chromatin condensation, this can impact the expression of nearby genes (Sigman and Slotkin, 2016). TEs are also increasingly being identified as modulatory factors during periods of host stress (i.e., heat, pathogen) (Liu et al., 2004; Xiao et al., 2008; Ding et al., 2015). For instance, cis-regulatory motifs within certain TEs can recruit stress-responsive host transcriptional factors, thereby influencing nearby host gene expression and potentially conferring a host benefit (Slotkin and Martienssen, 2007; Woodrow et al., 2010; Matsunaga et al., 2012; McCue and Slotkin, 2012; Cavrak et al., 2014; Makarevitch et al., 2015; Matsunaga et al., 2015; Negi et al., 2016; Galindo-Gonzalez et al., 2017). Some TEs also beneficially modulate the expression of host plant defense or susceptibility genes (Tsuchiya and Eulgem, 2013; Berg et al., 2015). Additionally, numerous studies report TEs lying within or adjacent to putative plant immune genes, however, potential influences on host genes or positive effects are often not apparent (Bhattacharyya et al., 1997; Henk et al., 1999; Wawrzynski et al., 2008).

Glycine max (soybean) is an important food and industrial crop (Schmutz et al., 2010). A major pest afflicting global soybean production is the soybean cyst nematode (SCN, Heterodera glycines), which causes yearly U.S. soybean yield losses of over 1 billion USD (Niblack et al., 2006; Mitchum, 2016; T. W. Allen, 2017). SCN is an obligate parasite that invades host roots and induces individual host cells to form a complex syncytium that serves as the SCN feeding site (Niblack et al., 2006; Mitchum, 2016). SCN feeding depletes available host resources and a functional syncytium must be maintained for 2-4 weeks for the nematode to complete its lifecycle. Since the unhatched eggs within cysts can remain viable for many years in the field, SCN is difficult to manage and is primarily controlled by growing naturally resistant soybeans (Niblack et al., 2006). Among known soybean loci contributing to SCN-resistance, the Rhg1 (Resistance to Heterodera glycines 1) locus found on chromosome 18 provides the strongest protection (Concibido et al., 2004). Rhg1 causes the SCN-induced syncytium to fail a few days after induction, and the soybean PI 88788-type "rhg1-b" haplotype is the primary SCN resistance locus used in commercially grown soybeans (Concibido et al., 2004; Niblack et al., 2006; Mitchum, 2016).

Soybean *Rhg1* is an unusual disease resistance locus that consists of a ~31.2 kb unit that is tandemly repeated as many as 10 times (Cook et al., 2012). Within each 31.2 kb *Rhg1* repeat unit are four different *Rhg1*-encoded genes: *Glyma.18G022400*, *Glyma.18G022500*, *Glyma.18G022500*, and *Glyma.18G022700*, none of which have similarity to previously identified resistance genes (Cook et al., 2012; Cook et al., 2014; Lee et al., 2015). Of the three *Rhg1* genes that contribute to SCN-resistance, only *Glyma.18G022500*, an α-SNAP (alpha-

<u>Soluble NSF Attachment Protein</u>), has amino acid polymorphisms relative to the wild-type *Rhg1* gene alleles present in SCN-susceptible soybeans (Cook et al., 2012; Cook et al., 2014; Lee et al., 2015). The mRNA transcript abundance of all three resistance-associated *Rhg1* genes is significantly elevated in SCN-resistant multi-copy *Rhg1* soybeans, relative to SCN-susceptible single-copy *Rhg1* (WT *Rhg1*) soybeans (Cook et al., 2012; Cook et al., 2014).

At least two distinct Rhg1 genotype classes exist: "low-copy Rhg1" (rhg1-a, sometimes referred to as Rhg1<sub>LC</sub>, often derived from PI 548402 'Peking'), and "high-copy Rhg1" (rhg1-b, sometimes referred to as Rhg1<sub>HC</sub>, often derived from PI 88788) (Niblack et al., 2002; Brucker et al., 2005; Cook et al., 2012; Cook et al., 2014; Bayless et al., 2018). These Rhg1 genotype classes represent two distinct multi-copy Rhg1 haplotypes that vary most notably by: a) Rhg1 repeat number (a high or low number of Rhg1 repeats), and b) encoding distinctive resistancetype  $\alpha$ -SNAP proteins with C-terminal polymorphisms at a conserved functional site (Cook et al., 2014; Bayless et al., 2016). rhg1-a resistance is bolstered by an unlinked chromosome 8 locus, Rhg4, whose presence contributes to full-strength "Peking-type" SCN resistance (Meksem et al., 2001; Liu et al., 2012). Rhg4 encodes a polymorphic serine hydroxy methyl transferase with altered enzyme kinetics, but the molecular basis of resistance augmentation by Rhg4 is not yet understood (Liu et al., 2012; Mitchum, 2016). Several rhg1-b and rhg1-a accessions have been analyzed by whole genome sequencing (WGS) studies, and characteristic single nucleotide polymorphism (SNP) signatures predictive of rhg1-b or rhg1-a haplotype soybeans have been reported (Cook et al., 2014; Lee et al., 2015; Shi et al., 2015; Kadam et al., 2016; Patil et al., 2019). Additionally, studies by Arelli, Young and others have profiled SCN-resistance among thousands of USDA soybean accessions and noted substantial phenotypic variation (e.g., (Anand, 1984; Hussey et al., 1991; Young, 1995; Diers et al., 1997; Arelli et al., 2000; Vuong et al., 2015; Klepadlo et al., 2018)). However, the influence of all Rhg1 haplotype and/or allelic variation factors on SCN resistance expression or plant yield is not yet fully understood.

Several recent studies have deepened our understanding of RhgI molecular function and highlight a central role of the SNARE (Soluble NSF Attachment Protein REceptors)-recycling machinery in SCN-resistance (Matsye et al., 2012; Cook et al., 2014; Bayless et al., 2016; Lakhssassi et al., 2017; Bayless et al., 2018).  $\alpha$ -SNAP, and the ATPase NSF (N-ethylmaleimide Sensitive Factor), are conserved eukaryotic housekeeping proteins that form the core SNARE-recycling machinery. They sustain the pool of fusion-competent SNAREs necessary for new membrane fusion events (Sudhof and Rothman, 2009; Zhao et al., 2015). While most animals encode single NSF and  $\alpha$ -SNAP genes, soybean is a paleopolyploid that encodes two NSF, four or five  $\alpha$ -SNAP and two  $\gamma$ -SNAP (gamma-SNAP) genes, respectively. A C-terminal  $\alpha$ -SNAP domain conserved across all plants and animals recruits NSF to SNARE-bundles and stimulates the ATPase activity of NSF that powers SNARE-complex recycling. However, it is this otherwise conserved  $\alpha$ -SNAP C-terminal region that is atypical among both rhg1-b- and rhg1-a-encoded  $\alpha$ -SNAP proteins, and accordingly, both Rhg1 resistance type  $\alpha$ -SNAPs are impaired in promoting normal NSF function and instead mediate dosage-dependent cytotoxicity (Sudhof and Rothman, 2009; Cook et al., 2014; Zhao et al., 2015; Bayless et al., 2016). The abundance of the

atypical rhg1-b  $\alpha$ -SNAP<sub>Rhg1</sub>HC protein specifically increases in the SCN feeding site and contributes to Rhg1-mediated collapse of the SCN-induced syncytium (Bayless et al., 2016).

At least two additional loci associated with SCN-resistance are also components of the SNARE-recycling machinery (Lakhssassi et al., 2017; Bayless et al., 2018). Recently, a specialized allele of NSF, NSF<sub>RAN07</sub> (Rhg1-associated NSF on chromosome 07), was shown to be necessary for the viability of Rhg1-containing soybeans (Bayless et al., 2018). Compared to the WT NSF<sub>Ch07</sub> protein, the NSF<sub>RAN07</sub> protein more effectively binds to resistance type  $\alpha$ -SNAPs and confers better protection against resistance-type α-SNAP-induced cytotoxicity (Bayless et al., 2018). During the Rhg1-mediated resistance response, the ratio of Rhg1 resistance-type to WT α-SNAPs increases and is apparently an important factor underlying resistance (Bayless et al., 2016; Bayless et al., 2018). Two genetic events sharply reduce WT α-SNAP protein abundance in SCN-resistant rhg1-a soybeans (Bayless et al., 2018). First, the wild type α-SNAPencoding block at Rhg1 on chromosome 18 - a predominant source of total WT α-SNAP proteins in soybean – is absent from all examined rhg1-a accessions, thereby diminishing overall WT  $\alpha$ -SNAP protein abundance (Cook et al., 2014; Bayless et al., 2018). Secondly, rhg1-a lines often carry a null allele of the α-SNAP encoded on chromosome 11 (Glyma.11G234500) - the other major source of WT α-SNAP proteins – due to an intronic splice site mutation that causes premature translational termination and loss of protein stability (Matsye et al., 2012; Cook et al., 2014; Lakhssassi et al., 2017; Bayless et al., 2018). Together, the above studies support a paradigm whereby Rhg1 and associated SCN-resistance loci rewire major components of the soybean SNARE-recycling machinery. Importantly, soybean accessions that carry rhg1-a and Rhg4 can resist many of the virulent SCN populations that partially overcome rhg1-b resistance (Brucker et al., 2005; Niblack et al., 2008; Bayless et al., 2016). Therefore, there is considerable interest in understanding and using rhg1-a, the subject of the present study, as an alternative to *rhg1-b* in commercial soybean cultivars (Brucker et al., 2005; Liu et al., 2012; Yu et al., 2016).

Presence/absence variation of transposable elements at specific loci is common among different soybean accessions, and tens of thousands of non-reference genome TE insertions occur between cultivated and wild soybean (Tian et al., 2012). Moreover, high TE densities near genomic regions exhibiting structural polymorphisms such as copy number variation are also reported in soybean (McHale et al., 2012). While examining the *Rhg1* low-copy (*rhg1-a*) haplotype of soybean accession PI 89772, we uncovered an intact copia retrotransposon within all three copies of the *Rhg1*-encoded α-SNAP genes. We termed this retrotransposon "RAC", for *Rhg1* alpha-SNAP copia). The *RAC* element, which is entirely within the first intron of the *Glyma.18G022500* (α-SNAP) gene, appears to be intact and transcribes anti-sense to the α-SNAP ORF. BLAST searches revealed similar copia elements across the soybean genome, suggesting why assemblies of Illumina short-read whole-genome sequences failed to include this sequence within *rhg1-a* assemblies. This α-SNAP-RAC insertion was absent from all examined single-copy *Rhg1* (SCN-susceptible) and high-copy *rhg1-b* (*Rhg1<sub>HC</sub>*) accessions. More than half of the USDA accessions with SoySNP50K SNPs preliminarily indicative of a low-copy *rhg1-a* haplotype did not carry *RAC*, but sub-sampling among those accessions revealed that they do not

encode  $\alpha$ -SNAP<sub>Rhg1</sub>LC and thus are not *rhg1-a*. The increasingly important *rhg1-a* SCN-resistant soybean breeding lines do harbor this previously unreported retrotransposon within the  $\alpha$ -SNAP<sub>Rhg1</sub>LC-encoding gene.

#### **Materials & Methods**

## Transgenic soybean hairy root generation and root culturing

Transgenic soybean roots were produced using *Agrobacterium rhizogenes* strain "ARqua1" (Quandt et al., 1993) and the previously described binary vector pSM101, as in (Cook et al., 2012). Transgenic roots were sub-cultured in the dark at room temperature on hairy root medium as in (Cook et al., 2012).

#### **DNA** extraction

Soybean genomic DNAs were extracted from expanding trifoliates or root tissues of the respective soybean accessions using standard CTAB methods similar to (Cook et al., 2012).

#### Amplification and detection of RAC (Rhg1 alpha-SNAP copia)

For initial amplification and subcloning of native α-SNAP-RAC, approximately 100 ng of CTAB-extracted gDNA from PI 89772 (*rhg1-a*) was PCR amplified for 35 cycles using HiFi polymerase (KAPA Biosystems, Wilmington, MA). Primer annealing was at ~70°C for 30 seconds and extension was at 72°C for 5 minutes. The resulting α-SNAP-RAC amplicon from PI 89772 was separated by agarose gel electrophoresis, gel extracted using a Zymoclean Large Fragment DNA Recovery Kit (Zymo Research, Irvine CA) and TA overhang cloned into a pTopo xL vector using the Topo xL PCR Cloning Kit (Life Technologies Corp., Carlsbad CA), per manufacturer's recommendations. For PCR detection of α-SNAP-RAC junctions or WT exon distances, ~25 ng of CTAB-extracted genomic DNA from each respective accession was amplified using GoTAQ Green (New England Biolabs, Ipswich MA) for 32 cycles, separated on a 0.8% agarose gel and visualized.

#### Phylogenetic tree construction

For the *RAC*-like nucleotide tree, evolutionary analyses were conducted in MEGA7 (Kumar et al., 2016) and evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model (Tamura et al., 2004). The tree with the highest log likelihood (-47751.11) is shown. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. The analysis involved 24 nucleotide sequences. All positions containing gaps and missing data were eliminated. There were a total of 4157 positions in the final dataset.

For the *RAC*-polyprotein tree, evolutionary analyses were conducted in MEGA7 (Kumar et al., 2016) and the evolutionary history was inferred by using the Maximum Likelihood method based on the JTT matrix-based model (Jones et al., 1992). The tree with the highest log likelihood (-12400.87) is shown. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and then selecting the topology with superior log likelihood value. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 14 amino acid sequences. All positions containing gaps and missing data were eliminated. There were a total of 644 positions in the final dataset.

### Read Depth Analysis of RAC

Using previously published whole-genome-sequencing data (Cook et al., 2014), read depth was computed using the depth program of SAMtools (Li et al., 2009). Depth was averaged in 250 bp intervals on Chromosome 10 from bp 40,650,000 - 40,690,000 (includes flanking regions of the 99.7% identity *RAC*-like element). The copy number of the Chromosome 10 *RAC*-like element was then calculated as the ratio of the read coverage per 250 bp from bp 40,672,000- 40,675,750, divided by the average read coverage for the flanking regions between bp 40,650,000 - 40,690,000. Sequencing coverage was visualized using ggplot (Wickham, 2009) within RStudio (RStudio Team, 2015).

## Methylation analysis

McrBC methylation studies were performed similarly to (Cook et al., 2014). Control McrBC reactions contained equivalent amounts of gDNA in reaction buffer, but had no added McrBC enzyme. McrBC digestion was performed at 37°C for 90 minutes, followed by a 20 minute heat inactivation at 65°C. McrBC digested or mock treated samples were PCR amplified with primers flanking 5' or 3' α-SNAP-RAC junctions and visualized by agarose gel electrophoresis.

#### RNA isolation and cDNA synthesis

Total RNAs were extracted using Trizol (Life Technologies Corp., Carlsbad CA) or the Direct-Zol RNA Miniprep Kit (Zymo Research, Irvine CA), per manufacturer's instructions. All RNA samples were DNAase treated and quantified using a spectrophotometer. cDNA synthesis was performed using the iScript cDNA Synthesis Kit (Bio-Rad, Hercules CA) according to manufacturer's recommendations using 1.0 µg of purified total RNA.

#### qPCR analysis

qPCR was performed with a CFX96 real-time PCR detection system (BioRad Laboratories, Hercules CA) using SsoFast EvaGreen Supermix (Bio-Rad Laboratories, Hercules CA) as in (Cook et al., 2012). Following amplification, a standardized melting curve analysis program was performed. Overall cDNA abundances for each sample were normalized using the qPCR signal

for reference gene *Glyma.18G022300*. *RAC* transcript abundances are presented relative to the mean abundance of *RAC* transcript for Williams 82 leaf samples.

#### **RT-PCR** analysis

For RT-PCR, 31 cycles of amplifications were performed prior to loading PCR product samples for separation and visualization by agarose gel electrophoresis. The number of PCR cycles terminated prior to maximal amplification of product from the most abundant template pool. A primer set sitting on a conserved copia region detected both endogenous *RAC*-like transcripts as well as the uniquely tagged *RAC* transgene. Specific detection of the tagged *RAC* transgene was with a primer pair sitting atop the engineered region. Transcript from *Glyma.18G022300* or *Skp16* served as a control for both cDNA quality and relative transcript abundance.

#### **Vector construction**

Native α-SNAP-RAC was PCR amplified from pTopo XL subclones using Kapa HiFi polymerase with AvrII and SbfI restriction site overhangs. Following agarose gel (cut with XbaI/PstI) (New England Biolabs, Ipswich MA). Gel extractions were performed using the QIAquick Gel Extraction Kit (Qiagen, Hilden Germany) or the Zymoclean Large Fragment DNA Recovery Kit (Zymo Research, Irvine CA). Purified DNA fragments were ligated overnight at 4°C with T4 DNA ligase (New England Biolabs, Ipswich MA) per manufacturer's recommendations. To remove RAC from within the native  $\alpha$ -SNAP-RAC subclone in vector pSM101, the Polymerase Incomplete Primer Extension (PIPE) PCR method was used with Kapa HiFi polymerase (Klock and Lesley, 2009). Similarly, the synonymous tag added within the RAC ORF of native  $\alpha$ -SNAP-RAC was created using PIPE. Unique nucleotide tag was located ~160 bp downstream of the RAC ATG and maintains an intact ORF. For creating the RAC only vector which assessed inherent RAC transcription, the native RAC ORF with both LTRs (~4.77 kb) was amplified from the initial PI 89772 α-SNAP-RAC subclone in pTopoXL using Kapa HiFi. Restriction site overhangs for AvrII and SbfI were incorporated into the primer sequences, and following gel recovery, the PCR amplicon was restriction digested and ligated into a PstI/XbaI cut pSM101 binary vector using T4 DNA ligase (New England Biolabs, Ipswich MA). For the native α-SNAP-RAC with flanking Rhg1 sequence, an 11.1 kb native Rhg1 sequence containing Glyma. 18G022400 and Glyma. 18G022500 (and ~1 kb downstream of each stop codon), was PCR amplified from a previously published fosmid subclone "Fos-32", with AvrII and SbfI restriction ends using Kapa HiFi polymerase (Cook et al., 2012). After restriction digestion, this amplified native fragment was ligated into the binary vector pSM101 (digested with PstI and XbaI). This created a native RhgI two gene vector of the rhgI-b type. Then, to make the native rhg1-a construct, this native rhg1-b pSM101 vector was used as a scaffold for step-wise cloning of two different native rhg1-a fragments amplified from PI 89772 genomic DNA. The first was a 4 kb fragment with an SbfI primer overhang containing Glyma. 18G022400 up til an endogenous NruI site at exon 1; the second 11 kb fragment resumed at NruI until ~ 1.0 kb downstream of the

*Glyma.18G022500* (α-SNAP<sub>Rhg1</sub>LC) termination codon and contained an AvrII restriction overhang.

## Immunobloting & antibodies

Affinity-purified polyclonal rabbit antibodies raised against  $\alpha$ -SNAP<sub>Rhg1</sub>LC and wild-type  $\alpha$ -SNAP C-terminus were previously generated and validated in 1(Bayless et al., 2016). Protein lysates were prepared from ~100 mg of soybean roots that were immediately flash-frozen in liquid N<sub>2</sub>. Roots were homogenized in a PowerLyzer 24 (Qiagen) for three cycles of 15 seconds, with flash-freezing in-between each cycle. Protein extraction buffer [50 mM Tris·HCl (pH 7.5), 150 mM NaCl, 5 mM EDTA, 0.2% Triton X-100, 10% (vol/vol) glycerol, 1/100 Sigma protease inhibitor cocktail] was then added at a 3:1 volume to mass ratio. Lysates were then centrifuged at 10,000g for 10 minutes and supernatant was added to SDS-PAGE loading buffer. Immunoblots were performed essentially as in (Song et al., 2015; Bayless et al., 2016). Briefly, immunoblots for α-SNAP<sub>Rhg1</sub>LC or WT α-SNAPs were incubated overnight at 4°C in 5% (wt/vol) nonfat dry milk TBS-T (50 mM Tris, 150 mM NaCl, 0.05% Tween 20) at 1:1,000. NSF immunoblots were performed similarly. Secondary horseradish peroxidase-conjugated goat antirabbit IgG was added at 1:10,000 and incubated for 1 h at room temperature on a platform shaker, followed by four washes with TBS-T. Chemiluminescence signal detection was performed with SuperSignal West Pico or Dura chemiluminescent substrates (Thermo Scientific, Waltham WA) and developed using a ChemiDoc MP chemiluminescent imager (Bio-Rad, Hercules CA).

#### Immunolabeling and Electron Microscopy.

Immunolabeling was performed as in (Bayless et al., 2016). Transverse sections of ~2 mm long soybean (cv. Forrest) root areas containing syncytia were harvested by hand-sectioning at 4 dpi. Root sections were fixed in 0.1% glutaraldehyde and 4% (vol/vol) paraformaldehyde in 0.1M sodium phosphate buffer (PB) (pH 7.4) overnight after vacuum infiltration for about 1 hour. After dehydration in ethanol, samples were then embedded in LR White Resin. Ultrathin sections (~90-nm) were taken longitudinally along the embedded root pieces using an ultramicrotome (UC-6; Leica) and mounted on nickel slot grids. For the immunogold labeling procedure, grids were first incubated on drops of 50 mM glycine/PBS for 15 min, and then blocked in drops of blocking solutions for goat gold conjugates (Aurion; Wageningen, NL) for 30 min and then equilibrated in 0.1% BSA-C/PBS (incubation buffer). Grids were then incubated overnight at  $4^{\circ}$ C with custom α-SNAP<sub>RhgI</sub>LC polyclonal antibody (diluted 1:1000 in incubation buffer), washed five times in incubation buffer, and incubated for 2 h with goat anti-rabbit antibody conjugated to 15-nm gold (Aurion) diluted 1:50 in incubation buffer. After six washes in incubation buffer and two 5-min washes in PBS, the grids were fixed for 5 min in 2.0% (vol/vol) glutaraldehyde in 0.1 M phosphate buffer, followed by two 5-min washes in 0.1 M phosphate buffer and five 2-min washes in water. Images were collected with a MegaView III digital camera on a Philips CM120 transmission electron microscope.

#### **Oligonucleotide Primers**

RAC qRT For: GGGTTCGAAATGAATACCTG
RAC qRT Rev: CACGTTCTTCTCATGGATCCTA

RAC Delete PIPE For: CTT CAT CCA CAA TTC TAA TTT ATA TGC TAG RAC Delete PIPE Rev: GAA TTG TGG ATG AAG TAC GAC AAT CAA C

5' RAC Junction For: TGGCTCCAAGTATGAAGATGCC

5' RAC Junction Rev: AACTACAGTGGCTGACCTTCT

3' RAC Junction For: ACTGTTCATTCAGACCGCGT

3' RAC Junction Rev: GCAATGTGCAGCATCGACATGGG

WT Junction For: GAGTTTTGAGGTGTCCGATTTCCC

WT Junction Rev: GTGAGCGCAGTCACAAACAAC

5' Methylation For: TGGCTCCAAGTATGAAGATGCC

5' Methylation Rev: AACTACAGTGGCTGACCTTCT

3' Methylation For: ACTGTTCATTCAGACCGCGT

3' Methylation Rev: GCAATGTGCAGCATCGACATGGG

Skp16 qRT For: GAG CCC AAG ACA TTG CGA GAG

Skp16 qRT Rev: CGG AAG CGG AAG AAC TGA ACC

2570 qRT For: TGA GAT GGG TGG AGC TCA AGA AC

2570 qRT Rev: AGC TTC ATC TGA TTG TGA CAG TGC

For RAC Tag Mut: GCTCTGCTCCTGAGCCCTTGAAAACGGACAGAATGCACGGAG Rev RAC Tag Mut: GCTCAGGAGCAGAGCCATCTATGAACTCCACTTTATTCTTGGC

RAC Tag Detect For: CAGTCCTAGACTCAACCAATTACC

RAC Tag Detect Rev: CCTTGGCTATACCTGCTCTTTAAATC

For RAC initial TopoXL subclone: GAGATTACATTGGATGATACGGTCGACC

Rev RAC initial TopoXL subclone: AGATAAGATCAGACTCCAGCAACCTC

For RAC Alone subclone AvrII: cctaggGGTGTCCGATTTCCCGATTAATTGAAG

Rev RAC Alone subclone SbfI: cctgcaggCCAACATCAATTTCAAAGTTCGTCACTTTC

LC-Splice Reverse: AGTAATAACCTCATACTCCTCAAGTT

LC-Splice Full For: GAGGAGGTTGTTGCTATAACCAATGC

LC-Splice Isoform For: GAGGAGGAACTGGATCCAACATTTTC

SbfI Native Glyma.18G022400 For: cctgcaggGAGCAGTAGGCTTCTTTGGAACTTG

AvrII Native Glyma. 18g022500 Rev:cctaggGTTCCTAAAGTGGCAAACCCTAAGAACAAAG

BglII Native For: AGATCTCCCTGAGAGTATCTTGATTTCAGATCG

BgIII Native Rev: AGATCTTTTACGCATATCCGACCTTCAAC

#### **Accession Numbers**

NCBI Accessions will be established for the RAC sequences shown in Supplemental Figs S2 and S3.

#### **Supplemental Data files**

See: Bayless 2019-SI-combined.pdf

- **Fig. S1.** Native genomic rhg1-a α-SNAP amplicons from PI 89772 are unexpectedly large and contain an inserted DNA element.
- **Fig. S2.** Complete nucleotide sequence of the PI 89772 (*rhg1-a*, *Rhg1* low-copy) *RAC* (element and flanking exonic α-SNAP<sub>Rhg1</sub>LC regions.
- **Fig. S3.** Translation of the *RAC*-encoded polyprotein (1438 residues) from accession PI 89772.
- Fig. S4. Depth of short genomic reads that align to the RAC nucleotide sequence.
- **Fig. S5.** Sequence of the RAC SNP (ss715606985), nucleotide SNP signatures of consensus rhg1-a (Rhg1 low-copy) and rhg1-b (Rhg1 high-copy) haplotypes, and detected Rhg1  $\alpha$ -SNAP transcripts from RAC<sup>+</sup> or RAC<sup>-</sup> accessions with consensus rhg1-a SNP-signatures.
- **Fig. S6.** Flow chart summarizing findings regarding RAC<sup>+</sup> vs. RAC<sup>-</sup> accessions, which otherwise have consensus SNP signatures for *rhg1-a*.
- Fig. S7. α-SNAP<sub>Rhg1</sub>LC immunolabeling in 'Forrest' roots is highly specific.
- **Table S1.** RAC-like elements identified from NBLAST searches of RAC against the Williams 82 reference genome at Phytozome.org.

#### Results

## Multiple rhg1-a haplotypes harbor an intronic copia retrotransposon (RAC) within the Rhg1-encoded $\alpha$ -SNAP

The  $\alpha$ -SNAPs encoded by the *rhg1-a* and *rhg1-b* loci play a key role in SCN-resistance (Cook et al., 2012; Cook et al., 2014; Bayless et al., 2016; Liu et al., 2017; Bayless et al., 2018). While entire 31.2 kb Rhg1 repeats of rhg1-b and Rhg1wT-like haplotypes have been sub-cloned and characterized (Cook et al., 2012), we sought to study the native genomic rhg1- $a\alpha$ -SNAPencoding region to investigate potential regulatory differences between rhg1-a and rhg1-b. Fig 1A provides a schematic of the 31.2 kb Rhg1 repeat unit with the four Rhg1-encoded genes, while Fig 1B presents a schematic of the previously published Rhg1 haplotypes (single-, low-, and high-copy) and the respective C-terminal amino acid polymorphisms of their Rhg1-encoded α-SNAP proteins (Cook et al., 2012; Cook et al., 2014). Working from previously generated WGS data, we PCR-amplified the native genomic rhg1-a α-SNAP locus from PI 89772, and unexpectedly, obtained PCR amplicons ~5 kb larger than WGS-based estimates (Fig S1A) (Cook et al., 2014; Liu et al., 2017). Overly large amplicons were also obtained using different PI 89772 genomic DNA templates and/or other PCR cycling conditions, and no rhg1-a α-SNAP amplicons of expected size were observed (Fig S1A). Sanger DNA sequencing of these unusually sized rhg1-a α-SNAP amplicons matched WGS predictions until part way into α-SNAP intron 1, where a 4.77 kb element was inserted. Immediately following this 4.77 kb insertion, the amplicon sequence again matched WGS predictions (Fig 1C).

An NCBI nucleotide BLAST of the unknown 4.77 kb region returned hits for conserved features of the Ty-1 copia retrotransposon superfamily. Notably, the multi-cistronic open reading

frame (ORF) of this copia element was fully intact, and both 5' and 3' LTRs were present (Long Terminal Repeats; LTRs function as transcriptional promoters and terminators, respectively) (Havecker et al., 2004). Subsequently, we named this insert 'RAC' for Rhg1 alpha-SNAP copia. Fig 1D shows the rhg1-a α-SNAP-RAC structure and Fig S2 provides the complete RAC nucleotide sequence and highlights the  $\alpha$ -SNAP<sub>Rhg1</sub>LC intron 1 sequences directly flanking the RAC-integration. The RAC insertion effectively doubles the pre-spliced  $\alpha$ -SNAP<sub>Rhg1</sub>LC mRNA transcript from 4.70 kb to 9.47 kb, yet RAC apparently splices out effectively, as all reported cDNAs of mature α-SNAP<sub>Rhg/</sub>LC transcripts do not contain any RAC sequences (Cook et al., 2014; Lee et al., 2015; Liu et al., 2017). The RAC ORF is uninterrupted and encodes a 1438 residue polyprotein with conserved copia retrotransposon motifs for GAG (group specific antigen) protease, integrase and reverse transcriptase (Peterson-Burch and Voytas, 2002; Havecker et al., 2004; Kanazawa et al., 2009). These conserved RAC polyprotein motifs are highlighted in Fig S3. Intriguingly, RAC integrated just 396 bp downstream of the  $\alpha$ -SNAP<sub>Rhg1</sub>LC start codon, and in an anti-sense orientation (Fig 1D). The intact LTRs and uninterrupted ORF suggest that RAC integration could have been a relatively recent event, and that RAC may remain functional.

PI 89772 (used above) is one of seven rhg1-a and rhg1-b soybean accessions used to determine the HG type of SCN populations (Niblack et al., 2002). We subsequently tested for RAC insertions within the Rhg1-encoded  $\alpha$ -SNAP(s) genes in the other six soybean accessions used in HG type tests. The RAC integration within the PI 89772-encoded  $\alpha$ - $SNAP_{Rhg1}LC$  creates unique 5' and 3' sequence junctions within the  $\alpha$ - $SNAP_{Rhg1}LC$  intron 1, and substantially increases the distance from  $\alpha$ -SNAP exon 1 to exon 2 from ~400 bp to ~5,000 bp (Fig 1B and C). Therefore, to screen for  $\alpha$ -SNAP-RAC insertions, we devised PCR assays specific for RAC- $\alpha$ -SNAP junctions, or for wild-type (uninterrupted) "WT junctions" separated by the genomic distances from exon 1 to exon 2 that are annotated in the soybean reference genome (Schmutz et al., 2010), as depicted in Fig 1C and 1D. Among all rhg1-a haplotype HG type test accessions (PI 90763, PI 89772, PI 437654 and PI 548402(Peking)-derived 'Forrest'), both 5' and 3'  $\alpha$ -SNAP-RAC junctions were detected (Fig 1D).

### RAC is absent from rhg1-b and single-copy Rhg1<sub>WT</sub> accessions

The  $\alpha$ -SNAP-RAC junctions were absent from the rhg1-b accessions tested (PI 88788, PI 548316, PI 209332), which instead gave WT junction PCR products (Fig 1D). Because no SNPs exist at the WT junction primer sites across any of the rhg1-b repeats (Cook et al., 2014), absence of 5' RAC junction and 3' RAC junction PCR products for the rhg1-b accessions suggests that those accessions do not carry the RAC copia element in the first intron of their  $\alpha$ -SNAP gene. Accession Williams 82 (Wm82, SCN-susceptible, Rhg1 single-copy), the source of the soybean reference genome, also gave a product for the WT junction reaction, and no PCR product for a RAC integration within the  $Rhg1_{WT}$  (single-copy)  $\alpha$ -SNAP gene, consistent with the reference genome annotation (Schmutz et al., 2010) (Fig S1B). RAC absence from rhg1-b and WT Rhg1 repeats is also consistent with previous studies that sub-cloned and Sanger-sequenced large-

insert genomic fragments spanning entire rhg1-b and  $Rhg1_{WT}$ -like repeats and noted no unusual insertions (Cook et al., 2012). Although all seven HG type test soybean lines were previously analyzed via WGS, the RAC insertion was evidently omitted from the four rhg1-a accession assemblies during Illumina short sequence read filtering that excludes repetitive genome elements, and hence from subsequent read mapping and assembly to the Williams 82 reference genome, which lacks the RAC insertion (Cook et al., 2014; Lee et al., 2015). The RAC insertion was apparently missed in other studies due to sequencing of post-splicing Rhg1  $\alpha$ -SNAP cDNAs (Bayless et al., 2016; Liu et al., 2017).

The WT junction PCR experiment also interrogated if RAC is present within each encoded α-SNAP gene of all three rhg1-a repeats. Among all rhg1-a accessions, no WT exon 1 to exon 2 junctions were detected, while uninterrupted WT junction product distances were present in all rhg1-b accessions and Wm82 as noted above (Fig 1E, F, Fig S1B). Hence RAC is apparently present within the  $\alpha$ -SNAP of each rhg1-a repeat unit. To independently investigate the same question, previously available Illumina whole genome sequence data were queried (Cook et al., 2014). The read-depth for RAC was found to be 3-4 fold greater in rhg1-a accessions relative to the read-depth of the flanking DNA regions, or when compared to RAC read depth in rhg1-b accessions (Fig S4, SI Spreadsheet). Table 1 provides a summary of the Rhg1 haplotype composition, copy number, resistance-type  $\alpha$ -SNAP allele, and presence of normal vs. RAC-interrupted  $\alpha$ -SNAP among the HG type test accessions and the Wm82 reference genome. Together, these findings indicate that the HG type test rhg1-a accessions contain the RAC- $\alpha$ -SNAP introgression, and that their rhg1-a repeats are ~36.0 kb, as opposed to ~31.2 kb for rhg1-b repeats and Rhg1<sub>WT</sub>. Although RAC is integrated in anti-sense orientation and close to  $\alpha$ -SNAP<sub>Rhg1</sub>LC exon 1, RAC does not eliminate rhg1-a function, because the HG type test accessions are selected owing to their strong SCN-resistance, and, all have previously been shown to express α-SNAP<sub>Rhg1</sub>LC mRNA and protein (Cook et al., 2014; Bayless et al., 2016; Liu et al., 2017; Bayless et al., 2018). RAC presence within these rhg1-a lines may even be beneficial.

## RAC is not identical to other copia elements but the RAC-like copia subfamily is common in soybean

Copia retrotransposons frequently attain high copy numbers in plant and animal genomes, therefore, we assessed the abundance of *RAC* and *RAC*-like copia elements in the soybean genome (Du et al., 2010; Du et al., 2010; Zhao and Ma, 2013). SoyTE, the soybean transposon database, has recorded over 32,000 transposable elements, including nearly 5,000 intact retrotransposons (Du et al., 2010). We queried the SoyTE database via Soybase.org using a *RAC* nucleotide sequence BLASTN search, but no intact or high identity hits were returned. However, a similar BLASTN search against the Wm 82 soybean reference genome at Phytozome.org (Goodstein et al., 2012) returned 146 sequences. These 146 hits spanned all 20 *G. max* chromosomes and included several intact elements of high nucleotide identity with *RAC*, as well as numerous short length matches which likely represent fragments of inactive elements (Table

S1). We then constructed a nucleotide-based phylogenetic tree of the soybean RAC family using just one RAC-family element from each soybean chromosome (Chr) (Fig 2A). This analysis used the element from each chromosome that was most similar to RAC, as well as a previously reported copia retrotransposon (TGMR) residing near the soybean Rps1-k resistance gene (Bhattacharyya et al., 1997) and the highest RAC-identity element match from the common bean (Phaseolus vulgaris) genome (Fig 2A). The two RAC-like elements, from Chr10 and Chr18, had 99.7% and 97.6% nucleotide identity with RAC, respectively (Fig 2A, Table S1). Moreover, the Chr10 element retained an intact ORF and both LTRs. The near-perfect nucleotide identity with RAC (99.7%; 4464/4477 positions) suggests the Chr10 element as a possible source for the retrotransposition event that created the RAC introgression within  $\alpha$ -SNAP<sub>Rhg1</sub>LC. The abovenoted WGS read-depth analysis of soybean accessions (Fig S4, SI Spreadsheet) also found that 3 of 10 examined rhg1-b accessions gave a read-depth of zero or close to zero for RAC (with one mismatch allowed), indicating absence of the RAC or a close homolog at the Chr 10 locus in some soybean accessions. Like RAC, the Chr18 element was also integrated anti-sense within a host gene Glyma. 18G268000 (a putative leucine-rich repeat receptor kinase). We further noted that the Chr20 RAC-like element (82% identical) was intronically positioned within Glyma. 20G250200 (BAR-domain containing protein) (Table S1). That multiple intact and highly similar RAC-like elements are in soybean suggests that this retrotransposon family was recently active.

Subsequent work examined if *RAC*-family elements are present among other plant species. We performed a TBLASTN search of the *RAC*-encoded polyprotein at NCBI and obtained numerous hits against multiple species, including *Arabidopsis*, *Cajanus cajans* (pigeon pea), *Vigna angularis* (adzuki bean), *Phaseolus vulgaris* (common bean), *Lupinus angustifolius* (blue lupin), *Medicago truncatula*, and clover (*Trifolium subterraneum*). Similar to Fig 2A, a phylogenetic tree using MEGA (muscle alignment) and the *RAC*-family polyprotein sequences of these different plant species was constructed (Kumar et al., 2016) (Fig 2B). Together, these findings demonstrate that the *RAC*-like copia members are not only common in *Glycine max*, but also in other legumes.

# *RAC* is present within only a subclass of the soybean accessions that have the previous SoySNP50K-predicted *rhg1-a* signature

Although we detected RAC in all four rhg1-a accessions that are used for HG type determination, we sought to determine if the RAC insertion is universal among all rhg1-a-containing accessions. Recently, the USDA soybean collection ( $\sim$ 20,000 accessions) was genotyped using a 50,000 SNP DNA microarray chip - the SoySNP50K iSelect BeadChip (Song et al., 2015). We searched for and found a SNP on the SoySNP50K chip that detects RAC. Using the SoySNP50K browser at Soybase.org (Soybase.org/snps/), we found a SNP (ss715606985, G to A) that in the Wm82 soybean reference genome was assigned to the Chr10 RAC-family element (99.7% nucleotide identity with RAC). However, we noted that this ss715606985 SNP is a perfect match to the sequence of RAC within  $\alpha$ - $SNAP_{Rhg1}LC$  (Fig S5A). Using this SNP marker

for RAC, we then calculated the ss715606985 SNP prevalence among all USDA accessions, and found that the SNP is rare - only 390 of 19,645 accessions (~2.0%) were putative  $RAC^+$  lines homozygous for the SNP (Fig 3A). The RAC-SNP (ss715606985) was then directly tested as a marker for the  $\alpha$ -SNAP-RAC event using the PCR assays described in Fig 1, which test for  $\alpha$ -SNAP-RAC junctions and normal  $\alpha$ -SNAP exon 1-2 distances. We randomly selected several accessions with SNP-signatures of rhg1-a that were positive or negative for the RAC-SNP, and found that SNP presence correlated perfectly with RAC- $\alpha$ -SNAP junction detection, while accessions lacking the RAC-SNP had normal  $\alpha$ -SNAP exon 1-2 distances indicative of no inserted DNA (Fig 3B).

We next examined RAC presence among all  $Glycine\ max$  USDA accessions with the SoySNP50K SNP-signatures of rhg1-a or rhg1-b haplotypes, as reported by Lee  $et\ al$ . (Lee et al., 2015). The multi-SNP SoySNP50K signatures for rhg1-a and rhg1-b (Lee et al., 2015) are present in 705 and 150  $Glycine\ max$  accessions respectively, out of 19,645 USDA accessions; these SoySNP50K signatures are provided in Fig S5B. We found that 42% (299 of 705) of accessions with rhg1-a SNP-signatures and 0% (0 of 150) of accessions with rhg1-b SNP-signatures carry the RAC ss715606985 SNP (Fig 3C). That the RAC-SNP was absent from all rhg1-b signature accessions is consistent with the PCR screens of Fig 1E and F, which indicated that no rhg1-b HG type test accession contained RAC-a-SNAP junctions. A flow-chart is available as Supplementary Fig S6 that summarizes the above findings and additional work presented below.

Because only 299 of the 390 accessions with the RAC-SNP had a perfect-match rhg1-a SoySNP50K SNP signature (Fig 3C), we investigated the Rhg1 SNP signature of the remainder. To avoid false positives, the rhg1-a SNP signature of (Lee et al., 2015) uses 14 SNPs that extend to 11 kb and 54 kb beyond the edges of the ~30 kb Rhg1 repeat. Relaxing the stringency that required 14 perfect matches, we found that 83 of the remaining 91 RAC accessions carry a perfect match with the four rhg1-a SNP markers that map within the Rhg1 repeat or within <5 kb of the edge of the Rhg1 repeat. 86 of 91 have only a single reliably called SNP that varies from the rhg1-a consensus (SI Spreadsheet). Together, the combined findings indicate that the  $\alpha$ -SNAP-RAC integration is only present within rhg1-a haplotypes, and that RAC retrotransposition may have occurred within a subset of the Rhg1 population after Rhg1 divergence into the distinctive high- and low-copy haplotype classes.

#### The RAC-SNP allows more accurate prediction of rhg1-a presence

The above finding that a few hundred of the 705 USDA accessions with the previously identified rhg1-a SNP signature apparently do not contain RAC was surprising, given that all four of the rhg1-a HG type test accessions do contain RAC-a-SNAP junctions. However, it was possible that these non-RAC containing accessions, despite a consensus SNP-signature predicting an rhg1-a-haplotype, might not truly carry an rhg1-a resistance haplotype. rhg1-a and rhg1-b repeats encode distinct Rhg1 a-SNAP alleles, thus, we cloned and sequenced the genomic Rhg1

 $\alpha$ -SNAP regions from several non-RAC rhg1-a SNP-signature accessions and detected coding sequences for either rhg1-b ( $\alpha$ -SNAP<sub>Rhg1</sub>HC) or Rhg1<sub>WT</sub> ( $\alpha$ -SNAP<sub>Rhg1</sub>WT) alleles (Fig S5C). None of these accessions encoded  $\alpha$ -SNAP<sub>Rhg1</sub>LC and thus, they were not rhg1-a (Fig S5C). These findings indicate that the consensus rhg1-a SNP-signature, while useful, is not a perfect predictor of accessions carrying actual rhg1-a resistance haplotypes. Rather, combined presence of the ss715606985 SNP for RAC and a near-consensus rhg1-a signature is a more accurate predictor of accessions that truly carry rhg1-a resistance. Additionally, these data suggest that accessions carrying rhg1-b resistance haplotypes can share the SNP signatures of rhg1-a accessions. We again refer readers to the flow-chart (Supplementary Fig S6) that summarizes these and other findings.

# RAC presence correlates with a stronger SCN-resistance profile and the co-presence of other loci that augment rhg1-a resistance

rhg1-a (Rhg1 low-copy) loci encode unique  $\alpha$ -SNAP<sub>Rhg1</sub>LC alleles, however, robust rhg1a resistance requires the co-presence of Rhg4, and the  $\alpha$ -SNAP Ch11-IR allele bolsters rhg1-a resistance further (Liu et al., 2012; Lakhssassi et al., 2017; Bayless et al., 2018; Patil et al., 2019). We sought to compare the SCN resistance profiles of the rhgl-a signature accessions with RAC to those without RAC (which are not true rhg1-a), to assess how the two groups match what is known about rhg1-a resistance. Previously, Arelli, Young, and others obtained SCN resistance phenotype data, across multiple trials and with various SCN populations, for at least 573 different USDA accessions that are now known to carry the SoySNP50K signatures suggestive of rhg1-a (Anand, 1984; Hussey et al., 1991; Young, 1995; Diers et al., 1997; Arelli et al., 2000; Lee et al., 2015). We used these available SCN resistance data from the USDA GRIN database to compare the resistance profiles of rhg1-a-signature accessions which did or did not have the ss715606985 (RAC) SNP signature. Any accession that scored as "R" (resistant) in any single SCN trial was placed into the broad category "R". Likewise, any accession that scored "MR" (moderately resistant) or "MS" (moderately susceptible) in any trial, with no higher resistance scores in other trials, was placed into those respective categories. Only accessions that scored susceptible ("S") across all trials were placed into the "S" category. Consistent with previous reports that rhg1-a accessions possess broad and robust resistance(Concibido et al., 2004; Vuong et al., 2015; Kadam et al., 2016), 91% (51/56) of the accessions in the "R" group were positive for the ss715606985 + RAC SNP (Fig 4A). The frequency of RAC presence was substantially lower among the more susceptible phenotypic classes (Fig 4A) while the majority of the ss715606985 (no RAC) accessions scored either "S" or "MS". As was noted above, none of the non-RAC accessions that we examined had rhgl-a type resistance (Fig 3B, 3C, Fig S5C). The phenotype scores and relevant SNP markers for all 573 of these SCN-phenotyped rhg1-a SNP signature accessions are provided as a spreadsheet in the SI Data.

In the above analysis (Fig 4A), some of the  $RAC^+$  (ss715606985<sup>+</sup>) accessions, which are rhg1-a, had scored as "S" or "MS". This seemed likely to be because they lack a resistance-conferring allele at Rhg4 and/or the resistance-enhancing allele of the Chr 11-encoded  $\alpha$ -SNAP

 $(\alpha$ -SNAP<sub>Ch11</sub>-IR) (Liu et al., 2012; Lakhssassi et al., 2017; Bayless et al., 2018). Accordingly, we investigated if the SCN resistance phenotype scores also correlated with co-presence of those loci. None of the SoySNP50K markers resides within the Rhg4 gene but we noted that the two SoySNP50K SNPs that most closely flank the Rhg4 locus are rare among USDA accessions (Fig. 4B; ss715602757, ss715602764), and one or both of these SNPs is present in the Rhg4containing HG type test lines. We also used a SNP, ss715610416, previously associated with the Chr11  $\alpha$ -SNAP intron-retention allele ( $\alpha$ -SNAP<sub>Ch11</sub>-IR) (Bayless et al., 2018). Among the 51 RAC  $^+$  accessions with an SCN resistance score of "R", we found that SNPs associated with both  $\alpha$ -SNAP<sub>Ch11</sub>-IR and Rhg4 were enriched ~10-fold, as compared to the entire USDA collection (Fig. 4 B,C). Additionally, among the 705 USDA accessions with SoySNP50K signatures predictive of rhg1-a, we found that the Rhg4 and  $\alpha$ -SNAP<sub>Ch11</sub>-IR SNPs were enriched among the RAC<sup>+</sup> accessions relative to the RAC<sup>-</sup> accessions (Fig 4D). Thus, the heightened SCN resistance of the RAC-positive (ss715606985 +) rhg1-a-signature accessions is consistent with previous reports, and as expected, the resistance is associated with the co-presence of additional loci like Rhg4 and α-SNAP<sub>Ch11</sub>-IR (Vuong et al., 2015; Kadam et al., 2016; Lakhssassi et al., 2017; Bayless et al., 2018; Patil et al., 2019).

#### The rhg1-a RAC element has intrinsic transcriptional activity

While the RAC-SNP apparently identifies true rhg1-a resistance sources, possible impacts of the RAC element itself on  $\alpha$ - $SNAP_{Rhg1}LC$  expression remained to be explored. Typically, eukaryotic cells silence transposable elements using small RNA-directed DNA methylation pathways, and this can also silence adjacent genes (McCue et al., 2012; Kim and Zilberman, 2014). Since the  $\alpha$ -SNAP<sub>Rhg1</sub>LC mRNA transcript and protein are readily detected in RAC-containing soybeans, RAC does not, at least constitutively, eliminate  $\alpha$ - $SNAP_{Rhg1}LC$  expression (Cook et al., 2014; Bayless et al., 2016; Liu et al., 2017; Bayless et al., 2018). Nonetheless, we examined DNA methylation at the rhg1-a  $\alpha$ -SNAP-RAC junction, as well as transcriptional activity of the rhg1-a RAC element. The restriction enzyme McrBC cleaves only methylated DNA, so potentially methylated DNA regions may be assessed via McrBC digestion and subsequent attempted PCR across areas of interest. After genomic DNAs from 'Forrest' and 'Peking' (PI 548402) were treated with McrBC, both the 5' and 3' borders of the  $\alpha$ -SNAP-RAC did not PCR amplify relative to the mock treated controls, indicating the presence of methylated cytosines at the  $\alpha$ -SNAP-RAC junctions (Fig 5A).

Because RAC has both LTRs and an intact ORF, we tested for transcription of RAC in the rhg1-a soybean genotype 'Forrest' as compared to 'Fayette' (rhg1-b) and Wm82 ( $Rhg1_{WT}$ ). Fayette and Wm82 do not carry the Rhg1  $\alpha$ -SNAP-RAC but do carry other RAC-like copia elements that match the qPCR primers used (diagrammed in Fig 5C). qPCR analysis of cDNAs from root or leaf tissues indicated that mRNA transcripts from RAC or RAC-like sequences were ~200-fold higher in 'Forrest' than in Wm82 or 'Fayette' (rhg1-b) (Fig 5B). This suggested but did not firmly demonstrate that the Rhg1-embedded RAC is the primary source of the detected

transcript, because *RAC* has high nucleotide identity with other *RAC*-like elements (Fig 2A) whose activity may also vary between accessions.

We conducted additional tests for transcription of  $\alpha$ -SNAP-RAC by transforming Wm82 roots with a  $\sim$ 15 kb cloned segment of native  $\alpha$ -SNAP-RAC genomic DNA (including the upstream Glyma. 18G024400 Rhg1 gene, which shares the same bidirectional promoter; depicted in Fig 5C). Importantly, we engineered this otherwise native  $\alpha$ -SNAP-RAC cassette with a unique nucleotide tag to distinguish between transgene derived transcripts vs. transcripts from other RAC-family elements in the genome (Fig 5C). Low abundance of RAC transcripts in Wm82 roots relative to 'Forrest' roots had been documented (Fig 5B), so RT-PCR of Wm82 readily visualized RAC-specific transcript expression from the transgenically introduced construct. In control roots, sharp contrasts in RAC expression were again observed between 'Forrest' roots and Wm82 roots transformed with empty vector (Fig 5D). But Wm82 roots transformed with the uniquely tagged α-SNAP-RAC transgene had substantially elevated RAC transcripts compared to isogenic Wm82 controls, as indicated by a primer pair that amplifies all RAC sequences (native or tagged), and by a primer pair that amplifies only the uniquely tagged  $\alpha$ -SNAP-RAC transcript (Fig 5D). Controls using template samples prepared without reverse transcriptase verified successful DNAase treatment of cDNA preparations (Fig 5D). We further tested the activity of the RAC promoter itself by constructing a native 4.77 kb RAC element cassette divorced from the flanking Rhg1 DNA (Fig 5E), which we then transformed into Wm82. Similar to Fig 5D, the native 4.77 kb RAC transgene substantially increased RAC transcript abundance in Wm82 roots, relative to empty vector controls (Fig 5F). Together, these findings demonstrate that presence of the rhg1-a locus RAC can substantially elevate RAC mRNA transcripts, and that RAC itself possesses intrinsic promoter activity. The findings suggest that the high RAC transcript abundance observed in 'Forrest' (rhg1-a), but not 'Fayette' (rhg1-b) or Wm 82 (single-copy Rhg1), is likely to be derived from the rhg1-a locus RAC insertion. These findings also support the possibility that RAC may retain the potential to promote transposition.

### α-SNAP<sub>RhgI</sub>LC protein is expressed despite RAC presence

TEs can influence the expression of host genes. Because RAC is present in rhg1-a accessions previously chosen for use in agricultural breeding due to their strong SCN-resistance, RAC presence may benefit rhg1-a-containing soybeans. In light of RAC's anti-sense orientation and close proximity to the Rhg1  $\alpha$ - $SNAP_{Rhg1}LC$  promoter, we sought to examine if RAC influences  $\alpha$ - $SNAP_{Rhg1}LC$  protein expression. We were not able to compare expression of  $\alpha$ - $SNAP_{Rhg1}LC$  between native rhg1-a loci that do or do not contain RAC, because no rhg1-a accessions without RAC have been identified and deleting RAC from all Rhg1 repeats of a rhg1-a accession would not be trivial. Therefore, we left intact or removed the 4.77 kb RAC insertion from the native  $\alpha$ -SNAP-RAC construct used for Fig 5 (Fig 6A) and then examined  $\alpha$ - $SNAP_{Rhg1}LC$  protein abundance in transgenic Wm82 roots carrying the respective constructs. Immunoblotting was conducted using previously described  $\alpha$ - $SNAP_{Rhg1}LC$ -specific and WT  $\alpha$ -SNAP-specific antibodies (Bayless et al., 2016). The results of a representative experiment are

shown in Fig 6B. Across multiple experiments containing independently transformed roots, the constitutive expression of  $\alpha$ -SNAP<sub>RhgI</sub>LC protein was highly variable, regardless of *RAC* presence/absence. However, with respect to constitutive expression of  $\alpha$ -SNAP<sub>RhgI</sub>LC protein we observed no requirement for *RAC* nor any obvious detrimental impact of *RAC* (Fig 6B).

Previously, we reported that native  $\alpha$ -SNAP<sub>RhgI</sub>LC mRNA transcripts include an alternative splice product (Cook et al., 2014; Bayless et al., 2016). Because TEs can influence host mRNA splicing (Krom et al., 2008), we also examined how *RAC* influenced splicing of the known  $\alpha$ -SNAP<sub>RhgI</sub>LC alternative transcript. As above, we generated transgenic roots of Wm82 containing either a  $RAC^+$   $\alpha$ -SNAP<sub>RhgI</sub>LC native genomic segment or a version with *RAC* precisely deleted (Fig 6A). We then generated cDNAs and performed RT-PCR with primer sets specific for either the full length or shorter splice isoform. As shown by agarose gel electrophoresis, *RAC* presence was not required for alternate splicing of this  $\alpha$ -SNAP<sub>RhgI</sub>LC isoform (Fig 6B).

## α-SNAP<sub>Rhg1</sub>LC hyperaccumulates at SCN infection sites

We also examined infection-associated α-SNAP<sub>RhgI</sub>LC protein expression in non-transgenic soybean roots that carry the native rhgI-a locus. We previously reported that during rhgI-b-mediated SCN-resistance, α-SNAP<sub>RhgI</sub>HC abundance is elevated ~12-fold within syncytial cells (SCN feeding sites) relative to adjacent non-syncytial cells (Bayless et al., 2016). To test whether the RAC-containing rhgI-a follows a similar expression pattern during the resistance response, the present study examined α-SNAP<sub>RhgI</sub>LC abundance at SCN infection sites in soybean variety 'Forrest' using SDS-PAGE and immunoblots. Using the aforementioned α-SNAP<sub>RhgI</sub>LC-specific antibody, we detected increased α-SNAP<sub>RhgI</sub>LC accumulation within tissues enriched for SCN feeding sites, while expression was barely detectable in mock-treated roots (Fig 7A). As previously reported, NSF proteins were also increased in SCN-infested roots, albeit less prominently (Fig 7A) (Bayless et al., 2016). Thus, even in RAC presence, rhgI-a haplotypes drive an expression pattern of α-SNAP<sub>RhgI</sub>LC similar to that observed for rhgI-b and α-SNAP<sub>RhgI</sub>HC.

To more precisely locate the α-SNAP<sub>RhgI</sub>LC increases, SCN-infested root sections were imaged using transmission electron microscopy and immunogold labelling of bound α-SNAP<sub>RhgI</sub>LC-specific antibody. Syncytium-specific accumulation of α-SNAP<sub>RhgI</sub>LC protein was observed and quantified in root sections taken 7 days after SCN inoculation (Fig 7B, C). The average increase of immunogold particles per equal area of adjacent non-syncytial root cells (cells still carrying a large central vacuole) was ~25-fold (Fig 7C). In control experiments, EM sections from mock-inoculated roots (no SCN) exhibited no immunogold signal above background (Fig S7A). Similarly, no immunogold signal above background was observed when secondary antibody and all other reagents were used but the primary antibody was omitted (Fig S7B). The specificity of the antibody for α-SNAP<sub>RhgI</sub>LC protein was previously demonstrated (signal for recombinant α-SNAP<sub>RhgI</sub>LC protein or total protein from roots with *rhg1-a*, no signal for α-SNAP<sub>RhgI</sub>HC protein or total protein from roots with *rhg1-b* or *Rhg1<sub>WT</sub>*) [30]. The above

results, similar to the previously observed ~12-fold increase reported for the  $\alpha$ -SNAP<sub>Rhg1</sub>HC in syncytia from rhg1-b roots, indicate that  $\alpha$ -SNAP<sub>Rhg1</sub>LC protein abundance is also elevated within syncytia upon SCN infection [30]. Collectively, these findings demonstrate that while a potentially active retrotransposon (RAC) has integrated within the important rhg1-a  $\alpha$ -SNAP<sub>Rhg1</sub>LC resistance gene, and its presence correlates with rhg1-a haplotypes preferred for SCN resistance breeding, no negative impacts of RAC on  $\alpha$ -SNAP<sub>Rhg1</sub>LC mRNA or protein expression were detected.

#### **Discussion**

Rhg1 is the principal SCN resistance locus in commercially grown soybeans. The increasing occurrence of SCN populations that at least partially overcome the overwhelmingly utilized "PI 88788-type" rhg1-b resistance source is an important concern for soybean breeders and growers (McCarville et al., 2017), (www.thescncoalition.com). Alternating use of different Rhg1 haplotypes should help bolster and preserve resistance against these virulent SCN populations (Brucker et al., 2005; Niblack et al., 2008)(www.thescncoalition.com). In this study we report that the other Rhg1 haplotype available for SCN control, rhg1-a (also known as "Peking-type" Rhg1), carries a distinct genetic structure. rhg1-a unexpectedly contains an intact and transcriptionally active retrotransposon within an intron of the key Rhg1 α-SNAP resistance gene in each repeat. The "Hartwig-type" SCN resistance from PI 437654 also carries the rhg1-a haplotype, and also carries the RAC retrotransposon within the Rhg1 α-SNAP genes.

Transposons have been coopted for the service of defense responses in both plants and animals (Tsuchiya and Eulgem, 2013; Huang et al., 2016). V(D)J recombination, which underlies the remarkable diversity of vertebrate adaptive immunity, apparently derives from a domesticated RAG-family transposase (Huang et al., 2016). RAC has inherent transcriptional activity and is positioned anti-sense within the first intron of the rhg1-a  $\alpha$ -SNAP gene. It is unclear if RAC impacts rhg1-a function (discussed below). However, this study revealed the utility of RAC and the RAC-associated ss715606985 (G to A) SNP in more accurately identifying SCN resistance-conferring rhg1-a germplasm. Among the 19,645 USDA soybean accessions genotyped using the SoySNP50K iSelect BeadChip (Song et al., 2015), a few hundred accessions with a rhg1-a-type SNP signature apparently do not actually carry a rhg1-a locus. All of the rhg1-a-signature soybeans we examined that do not carry RAC encoded rhg1-b or  $Rhg1_{WT}$   $\alpha$ -SNAP alleles. Conversely, all examined rhg1-a-signature soybeans with RAC carried the rhg1-a-a-SNAP allele.

Active retrotransposon families are abundant in soybean (Wawrzynski et al., 2008). RAC has similarities to a copia element near a *Phytophthora sojae* resistance locus identified by Bhattacharyya *et al.*, however, SoyTE database searches returned no highly similar RAC-family TEs (Bhattacharyya et al., 1997; Du et al., 2010). Although whole-genome sequencing studies previously examined the HG type test soybean accessions, the RAC insertion within  $\alpha$ -

*SNAP<sub>Rhg1</sub>LC* was apparently omitted during the filtering steps of DNA sequence read mapping and assembly (Cook et al., 2014; Liu et al., 2017). Our findings revealed multiple *RAC*-family elements in soybean, and this abundance of *RAC*-family elements likely led to *RAC* omission from previous *rhg1-a* sequence assemblies. It is intriguing that *RAC*, at least from PI 89772, has inherent transcriptional activity and an intact ORF encoding conserved functional motifs – features of an autonomous element. Additionally, *RAC*'s near perfect identity with the Chr10 element supports that *RAC* family retrotransposons were recently active in soybean.

Host silencing of transposable elements can establish *cis*-regulatory networks where the expression of nearby host genes may also be impacted (Lisch and Bennetzen, 2011; McCue et al., 2012; McCue and Slotkin, 2012). In some cases, biotic stresses can influence transposon methylation, and subsequently, alter the expression of host genes near transposons (Dowen et al., 2012). We noted DNA methylation at the  $\alpha$ -SNAP-RAC junctions. However, we also found evidence that RAC is transcriptionally active. In addition, the Rhg1  $\alpha$ -SNAP gene that contains RAC successfully expresses the  $\alpha$ -SNAP<sub>Rhg1</sub>LC protein during SCN-resistance similarly to that observed for  $\alpha$ -SNAP<sub>Rhg I</sub>HC. Future analyses of small RNAs may provide evidence of differential silencing of RAC or  $\alpha$ -SNAP<sub>Rhg1</sub>LC. Cyst nematode infection of Arabidopsis has been reported to trigger the hypomethyation and activation of certain transposable elements, and moreover, many of these transposable elements reside near host genes whose expression is altered during syncytium establishment (Hewezi et al., 2017; Piya et al., 2017). Thus, it remains an intriguing hypothesis that RAC may influence the epigenetic landscape of  $\alpha$ -SNAP<sub>Rhg1</sub>LC or the overall ~36 kb rhg1-a repeat during infection, particular stresses, developmental stages or in specific tissues. Moreover, small RNAs deriving from the other RAC-like elements in the soybean genome could modulate  $\alpha$ -SNAP<sub>Rhg1</sub>LC expression in trans (Slotkin and Martienssen, 2007; McCue et al., 2012; McCue et al., 2013). Our BLAST searches revealed at least two other RAC-like elements positioned intronically or adjacent to putative host defense and/or developmental genes. Future studies may interrogate if RAC, and/or other endogenous retrotransposons, impact the regulation of host defense gene networks in soybean.

The currently available picture of *Rhg1* haplotype evolution is incomplete and has been dominated by study of lines that are the product of ongoing selection for the most effective SCN-resistance (*e.g.* modern 10-copy *rhg1-b* and 3-copy *rhg1-a* haplotypes). The finding of *RAC* in all copies of the *rhg1-a* repeat, and in all confirmed *rhg1-a* haplotypes that were tested to date, suggests but does not confirm that *RAC* plays an adaptive role in those haplotypes. The sequence of the *Rhg1* repeat junction is identical between *rhg1-b* and *rhg1-a* haplotypes, as are many SNPs not present in the Williams 82 soybean reference genome, providing evidence of the shared evolutionary origin of *rhg1-b* and *rhg1-a* (Cook et al., 2012). The finding to date of *RAC* only in *rhg1-a* haplotypes suggests that this retroelement probably inserted in *Rhg1* after the divergence of *rhg1-b* and *rhg1-a*. However, it is also possible that the *RAC* retroelement was ancestrally present but then purged from the progenitors of current *rhg1-b* accessions. A correlation has been demonstrated between *Rhg1* copy number and SCN-resistance, and, *rhg1-a* in the absence of *Rhg4* confers only partial SCN resistance (Liu et al., 2012; Cook et al., 2014;

Lee et al., 2016; Yu et al., 2016; Kandoth et al., 2017). Yet there are no known instances of rhg1-a accessions with an Rhg1 copy number above three. The RAC may allow increased and/or more tightly regulated expression of rhg1-a. Alternatively, it is possible that absence of RAC is advantageous in allowing increased copy number of Rhg1, but that too is only a hypothesis, raised by the present work and in need of future testing. Additional questions about Rhg1 locus evolution remain that have functional implications for the efficacy of SCN resistance. For example, we know of no rhg1-a haplotypes that carry an  $\alpha$ -SNAP $_{Rhg1}$ WT-encoding Rhg1 repeat, which all examined rhg1-b haplotypes do contain. Might RAC acquisition have influenced this absence of the WT Rhg1 repeat? Do any  $\alpha$ -SNAP $_{Rhg1}$ LC-expressing accessions exist that do not carry the RAC integration? The USDA soybean collection contains numerous accessions that are positive for  $NSF_{RAN07}$  but which carry Rhg1 copy numbers below 3 or 10, or have slight deviations from consensus rhg1-a or rhg1-b SNP signatures. Intensive study of these accessions may shed further light on Rhg1 haplotype evolution, and moreover, may facilitate the discovery of new and agriculturally useful Rhg1 alleles.

The finding that popular rhg1-a breeding sources contain an intact retrotransposon within  $\alpha$ -SNAP<sub>Rhg1</sub>LC was surprising, given that these accessions have previously been sequenced multiple times by different groups (Cook et al., 2014; Liu et al., 2017; Patil et al., 2019). The RAC-SNP is rare among all USDA accessions, and approximately 700 of 19,645 USDA accessions carry a SoySNP50K signature predictive of an rhg1-a haplotype. However, ~400 of these putative rhg1-a accessions do not carry the RAC-SNP and all of the non-RAC putative rhg1-a accessions that we sampled did not encode  $\alpha$ -SNAP<sub>Rhg1</sub>LC, indicating that they are not true rhg1-a. Accordingly, most of these non-RAC accessions scored phenotypically as SCN-susceptible. Taken together with our PCR assays showing perfect correlation of the RAC-SNP with RAC presence, our findings indicate that the RAC-SNP successfully identifies true rhg1-a loci (i.e., those that encode the  $\alpha$ -SNAP<sub>Rhg1</sub>LC protein) which, in combination with Rhg4 and other loci, confers strong SCN resistance.

Correlation of SNP data with previously published SCN resistance phenotype data indicated that the vast majority of "R" scoring rhg1-a-signature accessions were  $RAC^+$ . Many of the 705 accessions postulated (using the earlier SoySNP50K SNP signature) to be lines that carry rhg1-a turned out to carry rhg1-b, which would explain their resistance to SCN. The large majority of the subset that are not positive for RAC were scored as SCN-susceptible or moderately susceptible. Some  $RAC^+$  lines also were scored as SCN-susceptible or moderately susceptible, but most of these are apparently due to the absence of a resistance-associated Rhg4 and/or the Chr 11  $\alpha$ -SNAP intron-retention allele ( $\alpha$ - $SNAP_{Ch11}$ -IR), consistent with the established contributions of those loci to SCN resistance (Meksem et al., 2001; Yu et al., 2016; Kandoth et al., 2017). Among the "R" scoring  $RAC^+$  accessions, SNPs genetically linked to Rhg4 and the Chr 11  $\alpha$ -SNAP intron-retention allele were substantially elevated.

Potential modulation of  $\alpha$ -SNAP<sub>Rhg1</sub>LC expression by RAC, either during the SCN-resistance response or in certain developmental or stress situations, could benefit *rhg1-a*-

containing soybeans that have significantly depleted WT α-SNAP proteins. α-SNAPs, together with NSF, carry out essential eukaryotic housekeeping functions by maintaining SNARE proteins for vesicle trafficking. Notably, among true rhg1-a accessions, the abundance of wildtype (WT) α-SNAP proteins is sharply diminished, as compared to rhg1-b- or SCN-susceptible soybeans (Bayless et al., 2018). Moreover, α-SNAP<sub>RhgI</sub>LC protein was shown to be cytotoxic in Nicotiana benthamiana while WT α-SNAP co-expression alleviated this toxicity (Bayless et al., 2016). The more recent discovery of the  $NSF_{RAN07}$  allele as a requisite for the viability of Rhg1 soybeans further underscores the necessity of the SNARE-recycling machinery for overall plant health (Bayless et al., 2018). The intronic copia element within the Arabidopsis RPP7 (Recognition of Peronospora Parasitica 7) gene serves as an example of a retroelement with immunomodulatory function, having been shown to modulate RPP7 transcript splicing and expression (Tsuchiya and Eulgem, 2013). However, we did not detect any influence of RAC on constitutive α-SNAP<sub>Rhg1</sub>LC protein expression from transgenes delivered to roots. We also did not detect a functional influence of RAC in its native rhg1-a haplotype context, insofar as that  $\alpha$ - $SNAP_{RhgI}LC$  protein abundance is successfully elevated in syncytia similarly to the reported syncytium elevation of the  $\alpha$ -SNAP<sub>RhgI</sub>HC protein of rhgI-b haplotypes, which do not carry RAC (Bayless et al., 2016). Hence although we did not find any true rhg1-a soybeans without RAC integrations, it remains possible that RAC integration was a neutral event that confers no host advantage or disadvantage.

SCN causes the most yield loss of any disease for U.S. soybean farmers and *rhg1-a* offers a potential solution to SCN populations that overcome commonly used *rhg1-b* resistance sources (Brucker et al., 2005; Niblack et al., 2008; T. W. Allen, 2017). Findings continue to emerge that further characterize different sources of SCN resistance, including exciting new findings regarding copy number variation at *Rhg4* (Patil et al., 2019). An attractive overall hypothesis for future study of *RAC* is that, in the presence of SCN or other stresses, *RAC* provides an additional regulatory layer to optimize the SCN resistance response mediated by *rhg1-a* and *Rhg4*, and/or promotes plant health in the absence of SCN. By revealing the existence of *RAC* within the important *rhg1-a* haplotype, the present study provides a marker for finding such soybeans, and expands our knowledge regarding the genetic structure and divergence of the agriculturally valuable *Rhg1* source of SCN resistance.

## **Literature Cited**

- **Anand SC, Gallo, K.M.** (1984) Identification of additional soybean germplasm with resistance to race 3 of the soybean cyst nematode. Plant Diseases (USA)
- **Arelli PR, Sleper DA, Yue P, Wilcox JA** (2000) Soybean Reaction to Races 1 and 2 of *Heterodera glycines*. Contribution of the Missouri Agric. Exp. Stn. Journal Series No. 12,751. Crop Science **40**: 824-826
- Bayless AM, Smith JM, Song J, McMinn PH, Teillet A, August BK, Bent AF (2016) Disease resistance through impairment of alpha-SNAP-NSF interaction and vesicular trafficking by soybean *Rhg1*. Proc Natl Acad Sci U S A **113**: E7375-E7382
- Bayless AM, Zapotocny RW, Grunwald DJ, Amundson KK, Diers BW, Bent AF (2018) An atypical N-ethylmaleimide sensitive factor enables the viability of nematode-resistant *Rhg1* soybeans. Proc Natl Acad Sci U S A **115**: E4512-E4521
- Berg JA, Appiano M, Santillan Martinez M, Hermans FW, Vriezen WH, Visser RG, Bai Y, Schouten HJ (2015) A transposable element insertion in the susceptibility gene *CsaMLO8* results in hypocotyl resistance to powdery mildew in cucumber. BMC Plant Biol 15: 243
- **Bhattacharyya MK, Gonzales RA, Kraft M, Buzzell RI** (1997) A copia-like retrotransposon Tgmr closely linked to the *Rps1-k* allele that confers race-specific resistance of soybean to *Phytophthora sojae*. Plant Mol Biol **34:** 255-264
- Brucker E, Carlson S, Wright E, Niblack T, Diers B (2005) *Rhg1* alleles from soybean PI 437654 and PI 88788 respond differentially to isolates of *Heterodera glycines* in the greenhouse. Theor Appl Genet 111: 44-49
- Cavrak VV, Lettner N, Jamge S, Kosarewicz A, Bayer LM, Mittelsten Scheid O (2014)

  How a retrotransposon exploits the plant's heat stress response for its activation. PLoS Genet 10: e1004115
- Concibido VC, Diers BW, Arelli PR (2004) A decade of QTL mapping for cyst nematode resistance in soybean. Crop Sci. 44: 1121-1131
- Cook DE, Bayless AM, Wang K, Guo X, Song Q, Jiang J, Bent AF (2014) Distinct copy number, coding sequence, and locus methylation patterns underlie *Rhg1*-mediated soybean resistance tosoybean cyst nematode. Plant Physiol **165**: 630-647
- Cook DE, Lee TG, Guo X, Melito S, Wang K, Bayless AM, Wang J, Hughes TJ, Willis DK, Clemente TE, Diers BW, Jiang J, Hudson ME, Bent AF (2012) Copy number variation of multiple genes at *Rhg1* mediates nematode resistance in soybean. Science 338: 1206-1209
- **Diers BW, Skorupska HT, Rao-Arelli AP, Cianzio SR** (1997) Genetic relationships among soybean plant introductions with resistance to soybean cyst nematodes. Crop Science **37**: 1966-1972
- Ding M, Ye W, Lin L, He S, Du X, Chen A, Cao Y, Qin Y, Yang F, Jiang Y, Zhang H, Wang X, Paterson AH, Rong J (2015) The hairless stem phenotype of cotton (*Gossypium barbadense*) is linked to a copia-like retrotransposon insertion in a homeodomain-leucine zipper gene (*HD1*). Genetics 201: 143-154
- Dowen RH, Pelizzola M, Schmitz RJ, Lister R, Dowen JM, Nery JR, Dixon JE, Ecker JR (2012) Widespread dynamic DNA methylation in response to biotic stress. Proc Natl Acad Sci U S A 109: E2183-2191

- **Du J, Grant D, Tian Z, Nelson RT, Zhu L, Shoemaker RC, Ma J** (2010) SoyTEdb: a comprehensive database of transposable elements in the soybean genome. BMC Genomics **11**: 113
- Du J, Tian Z, Hans CS, Laten HM, Cannon SB, Jackson SA, Shoemaker RC, Ma J (2010) Evolutionary conservation, diversity and specificity of LTR-retrotransposons in flowering plants: insights from genome-wide analysis and multi-specific comparison. Plant J 63: 584-598
- Galindo-Gonzalez L, Mhiri C, Deyholos MK, Grandbastien MA (2017) LTR-retrotransposons in plants: Engines of evolution. Gene **626**: 14-25
- Goodstein DM, Shu S, Howson R, Neupane R, Hayes RD, Fazo J, Mitros T, Dirks W, Hellsten U, Putnam N, Rokhsar DS (2012) Phytozome: a comparative platform for green plant genomics. Nucleic Acids Res 40: D1178-1186
- **Havecker ER, Gao X, Voytas DF** (2004) The diversity of LTR retrotransposons. Genome Biol **5:** 225
- **Henk AD, Warren RF, Innes RW** (1999) A new Ac-like transposon of Arabidopsis is associated with a deletion of the *RPS5* disease resistance gene. Genetics **151**: 1581-1589
- Hewezi T, Lane T, Piya S, Rambani A, Rice JH, Staton M (2017) Cyst nematode parasitism induces dynamic changes in the root epigenome. Plant Physiol 174: 405-420
- Huang S, Tao X, Yuan S, Zhang Y, Li P, Beilinson HA, Zhang Y, Yu W, Pontarotti P, Escriva H, Le Petillon Y, Liu X, Chen S, Schatz DG, Xu A (2016) Discovery of an active RAG transposon illuminates the origins of V(D)J recombination. Cell 166: 102-114
- **Hussey RS, Boerma HR, Raymer PL, Luzzi BM** (1991) Resistance in soybean cultivars from maturity groups V-VIII to soybean cyst and root-knot nematodes. J Nematol **23:** 576-583
- **Jones DT, Taylor WR, Thornton JM** (1992) The rapid generation of mutation data matrices from protein sequences. Comput Appl Biosci **8:** 275-282
- Kadam S, Vuong TD, Qiu D, Meinhardt CG, Song L, Deshmukh R, Patil G, Wan J, Valliyodan B, Scaboo AM, Shannon JG, Nguyen HT (2016) Genomic-assisted phylogenetic analysis and marker development for next generation soybean cyst nematode resistance breeding. Plant Sci 242: 342-350
- Kanazawa A, Liu B, Kong F, Arase S, Abe J (2009) Adaptive evolution involving gene duplication and insertion of a novel Ty1/copia-like retrotransposon in soybean. J Mol Evol 69: 164-175
- Kandoth PK, Liu S, Prenger E, Ludwig A, Lakhssassi N, Heinz R, Zhou Z, Howland A, Gunther J, Eidson S, Dhroso A, LaFayette P, Tucker D, Johnson S, Anderson J, Alaswad A, Cianzio SR, Parrott WA, Korkin D, Meksem K, Mitchum MG (2017) Systematic mutagenesis of serine hydroxymethyltransferase reveals an essential role in nematode resistance. Plant Physiol 175: 1370-1380
- **Kim MY, Zilberman D** (2014) DNA methylation as a system of plant genomic immunity. Trends Plant Sci **19:** 320-326
- Klepadlo M, Meinhardt CG, Vuong TD, Patil G, Bachleda N, Ye H, Robbins RT, Li Z, Shannon JG, Chen P, Meksem K, Nguyen HT (2018) Evaluation of soybean germplasm for resistance to multiple nematode species: *Heterodera glycines*, *Meloidogyne incognita*, and *Rotylenchulus reniformis*. Crop Science **58**: 2511-2522

- **Klock HE, Lesley SA** (2009) The Polymerase Incomplete Primer Extension (PIPE) method applied to high-throughput cloning and site-directed mutagenesis. Methods Mol Biol **498:** 91-103
- **Krom N, Recla J, Ramakrishna W** (2008) Analysis of genes associated with retrotransposons in the rice genome. Genetica **134:** 297-310
- **Kumar S, Stecher G, Tamura K** (2016) MEGA7: Molecular Evolutionary Genetics Analysis Version 7.0 for Bigger Datasets. Mol Biol Evol **33:** 1870-1874
- Lakhssassi N, Liu S, Bekal S, Zhou Z, Colantonio V, Lambert K, Barakat A, Meksem K (2017) Characterization of the Soluble NSF Attachment Protein gene family identifies two members involved in additive resistance to a plant pathogen. Sci Rep 7: 45226
- **Lee TG, Diers BW, Hudson ME** (2016) An efficient method for measuring copy number variation applied to improvement of nematode resistance in soybean. Plant J **88:** 143-153
- Lee TG, Kumar I, Diers BW, Hudson ME (2015) Evolution and selection of *Rhg1*, a copynumber variant nematode-resistance locus. Mol Ecol **24:** 1774-1791
- Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R, Genome Project Data Processing S (2009) The Sequence Alignment/Map format and SAMtools. Bioinformatics 25: 2078-2079
- Lisch D (2013) How important are transposons for plant evolution? Nat Rev Genet 14: 49-61
- **Lisch D, Bennetzen JL** (2011) Transposable element origins of epigenetic gene regulation. Curr Opin Plant Biol **14:** 156-161
- Liu J, He Y, Amasino R, Chen X (2004) siRNAs targeting an intronic transposon in the regulation of natural flowering behavior in Arabidopsis. Genes Dev 18: 2873-2878
- Liu S, Kandoth PK, Lakhssassi N, Kang J, Colantonio V, Heinz R, Yeckel G, Zhou Z, Bekal S, Dapprich J, Rotter B, Cianzio S, Mitchum MG, Meksem K (2017) The soybean *GmSNAP18* gene underlies two types of resistance to soybean cyst nematode. Nat Commun 8: 14822
- Liu S, Kandoth PK, Warren SD, Yeckel G, Heinz R, Alden J, Yang C, Jamai A, El-Mellouki T, Juvale PS, Hill J, Baum TJ, Cianzio S, Whitham SA, Korkin D, Mitchum MG, Meksem K (2012) A soybean cyst nematode resistance gene points to a new mechanism of plant resistance to pathogens. Nature 492: 256-260
- Makarevitch I, Waters AJ, West PT, Stitzer M, Hirsch CN, Ross-Ibarra J, Springer NM (2015) Transposable elements contribute to activation of maize genes in response to abiotic stress. PLoS Genet 11: e1004915
- **Matsunaga W, Kobayashi A, Kato A, Ito H** (2012) The effects of heat induction and the siRNA biogenesis pathway on the transgenerational transposition of ONSEN, a copialike retrotransposon in *Arabidopsis thaliana*. Plant Cell Physiol **53**: 824-833
- Matsunaga W, Ohama N, Tanabe N, Masuta Y, Masuda S, Mitani N, Yamaguchi-Shinozaki K, Ma JF, Kato A, Ito H (2015) A small RNA mediated regulation of a stress-activated retrotransposon and the tissue specific transposition during the reproductive period in Arabidopsis. Front Plant Sci 6: 48
- Matsye PD, Lawrence GW, Youssef RM, Kim KH, Lawrence KS, Matthews BF, Klink VP (2012) The expression of a naturally occurring, truncated allele of an alpha-SNAP gene suppresses plant parasitic nematode infection. Plant Mol Biol 80: 131-155
- McCarville MT, Marett CC, Mullaney MP, Gebhart GD, Tylka GL (2017) Increase in Soybean cyst nematode virulence and reproduction on resistant soybean varieties in Iowa from 2001 to 2015 and the effects on soybean yields. Plant Health Progress 18: 146-155

- McCue AD, Nuthikattu S, Reeder SH, Slotkin RK (2012) Gene expression and stress response mediated by the epigenetic regulation of a transposable element small RNA. PLoS Genet 8: e1002474
- McCue AD, Nuthikattu S, Slotkin RK (2013) Genome-wide identification of genes regulated in trans by transposable element small interfering RNAs. RNA Biol 10: 1379-1395
- McCue AD, Slotkin RK (2012) Transposable element small RNAs as regulators of gene expression. Trends Genet 28: 616-623
- McHale LK, Haun WJ, Xu WW, Bhaskar PB, Anderson JE, Hyten DL, Gerhardt DJ, Jeddeloh JA, Stupar RM (2012) Structural variants in the soybean genome localize to clusters of biotic stress-response genes. Plant Physiol 159: 1295-1308
- Meksem K, Pantazopoulos P, Njiti V, Hyten L, Arelli P, Lightfoot D (2001) 'Forrest'resistance to the soybean cyst nematode is bigenic: saturation mapping of the *Rhg1* and *Rhg4* loci. Theoretical and Applied Genetics 103: 710-717
- **Mitchum MG** (2016) Soybean Resistance to the Soybean Cyst Nematode Heterodera glycines: An Update. Phytopathology **106**: 1444-1450
- **Negi P, Rai AN, Suprasanna P** (2016) Moving through the stressed genome: Emerging regulatory roles for transposons in plant stress response. Front Plant Sci 7: 1448
- Niblack T, Colgrove A, Colgrove K, Bond J (2008) Shift in virulence of soybean cyst nematode is associated with use of resistance from PI 88788. Plant Health Prog 10
- Niblack TL, Arelli PR, Noel GR, Opperman CH, Orf JH, Schmitt DP, Shannon JG, Tylka GL (2002) A revised classification scheme for genetically diverse populations of *Heterodera glycines*. J Nematol **34:** 279-288
- **Niblack TL, Lambert KN, Tylka GL** (2006) A model plant pathogen from the kingdom Animalia: *Heterodera glycines*, the soybean cyst nematode. Annu Rev Phytopathol **44:** 283-303
- Patil GB, Lakhssassi N, Wan J, Song L, Zhou Z, Klepadlo M, Vuong TD, Stec AO, Kahil SS, Colantonio V, Valliyodan B, Rice HJ, Piya S, Hewezi T, Stupar RM, Meksem K, Nguyen HT (2019) Whole genome re-sequencing reveals the impact of the interaction of copy number variants of the *rhg-1* and *Rhg4* genes on broad-based resistance to soybean cyst nematode. Plant Biotechnol J
- **Peterson-Burch BD, Voytas DF** (2002) Genes of the Pseudoviridae (Ty1/copia retrotransposons). Mol Biol Evol **19:** 1832-1845
- **Piya S, Bennett M, Rambani A, Hewezi T** (2017) Transcriptional activity of transposable elements may contribute to gene expression changes in the syncytium formed by cyst nematode in arabidopsis roots. Plant Signal Behav **12**: e1362521
- **Quandt HJ, Pühler A, Broer I** (1993) Transgenic root nodules of *Vicia hirsuta*: a fast and efficient system for the study of gene expression in indeterminate-type nodules. MPMI-Molecular Plant Microbe Interactions **6:** 699-706
- Schmutz J, Cannon SB, Schlueter J, Ma J, Mitros T, Nelson W, Hyten DL, Song Q, Thelen JJ, Cheng J, Xu D, Hellsten U, May GD, Yu Y, Sakurai T, Umezawa T, Bhattacharyya MK, Sandhu D, Valliyodan B, Lindquist E, Peto M, Grant D, Shu S, Goodstein D, Barry K, Futrell-Griggs M, Abernathy B, Du J, Tian Z, Zhu L, Gill N, Joshi T, Libault M, Sethuraman A, Zhang XC, Shinozaki K, Nguyen HT, Wing RA, Cregan P, Specht J, Grimwood J, Rokhsar D, Stacey G, Shoemaker RC, Jackson SA (2010) Genome sequence of the palaeopolyploid soybean. Nature 463: 178-183

- Shi Z, Liu S, Noe J, Arelli P, Meksem K, Li Z (2015) SNP identification and marker assay development for high-throughput selection of soybean cyst nematode resistance. BMC Genomics 16: 314
- **Sigman MJ, Slotkin RK** (2016) The first rule of plant transposable element silencing: location, location, location. Plant Cell **28**: 304-313
- **Slotkin RK, Martienssen R** (2007) Transposable elements and the epigenetic regulation of the genome. Nat Rev Genet **8:** 272-285
- **Song J, Keppler BD, Wise RR, Bent AF** (2015) PARP2 Is the predominant poly(ADP-ribose) polymerase in Arabidopsis DNA damage and immune responses. PLoS Genet **11**: e1005200
- Song Q, Hyten DL, Jia G, Quigley CV, Fickus EW, Nelson RL, Cregan PB (2015)
  Fingerprinting soybean germplasm and its utility in genomic research. G3 (Bethesda) 5: 1999-2006
- **Sudhof TC, Rothman JE** (2009) Membrane fusion: grappling with SNARE and SM proteins. Science **323**: 474-477
- T. W. Allen CAB, A. J. Sisson, E. Byamukama, M. I. Chilvers, C. M. Coker, A. A. Collins, J. P. Damicone, A. E. Dorrance, N. S. Dufault, P. D. Esker, T. R. Faske, L. J. Giesler, A. P. Grybauskas, D. E. Hershman, C. A. Hollier, T. Isakeit, D. J. Jardine, H. M. Kelly, R. C. Kemerait, N. M. Kleczewski, S. R. Koenning, J. E. Kurle, D. K. Malvick, S. G. Markell, H. L. Mehl, D. S. Mueller, J. D. Mueller, R. P. Mulrooney, B. D. Nelson, M. A. Newman, L. Osborne, C. Overstreet, G. B. Padgett, P. M. Phipps, P. P. Price, E. J. Sikora, D. L. Smith, T. N. Spurlock, C. A. Tande, A. U. Tenuta, K. A. Wise, and J. A. Wrather. (2017) Soybean yield loss estimates due to diseases in the United States and Ontario, Canada, from 2010 to 2014. Plant Health Research
- **Tamura K, Nei M, Kumar S** (2004) Prospects for inferring very large phylogenies by using the neighbor-joining method. Proc Natl Acad Sci U S A **101**: 11030-11035
- Tian Z, Zhao M, She M, Du J, Cannon SB, Liu X, Xu X, Qi X, Li MW, Lam HM, Ma J (2012) Genome-wide characterization of nonreference transposons reveals evolutionary propensities of transposons in soybean. Plant Cell 24: 4422-4436
- **Tsuchiya T, Eulgem T** (2013) An alternative polyadenylation mechanism coopted to the Arabidopsis RPP7 gene through intronic retrotransposon domestication. Proc Natl Acad Sci U S A **110**: E3535-3543
- Vuong TD, Sonah H, Meinhardt CG, Deshmukh R, Kadam S, Nelson RL, Shannon JG, Nguyen HT (2015) Genetic architecture of cyst nematode resistance revealed by genome-wide association study in soybean. BMC Genomics 16: 593
- Wawrzynski A, Ashfield T, Chen NW, Mammadov J, Nguyen A, Podicheti R, Cannon SB, Thareau V, Ameline-Torregrosa C, Cannon E, Chacko B, Couloux A, Dalwani A, Denny R, Deshpande S, Egan AN, Glover N, Howell S, Ilut D, Lai H, Del Campo SM, Metcalf M, O'Bleness M, Pfeil BE, Ratnaparkhe MB, Samain S, Sanders I, Segurens B, Sevignac M, Sherman-Broyles S, Tucker DM, Yi J, Doyle JJ, Geffroy V, Roe BA, Maroof MA, Young ND, Innes RW (2008) Replication of nonautonomous retroelements in soybean appears to be both recent and common. Plant Physiol 148: 1760-1771
- Wickham H (2009) Ggplot2: elegant graphics for data analysis. Springer, New York

- Woodrow P, Pontecorvo G, Fantaccione S, Fuggi A, Kafantaris I, Parisi D, Carillo P (2010) Polymorphism of a new Ty1-copia retrotransposon in durum wheat under salt and light stresses. Theor Appl Genet 121: 311-322
- Xiao H, Jiang N, Schaffner E, Stockinger EJ, van der Knaap E (2008) A retrotransposonmediated gene duplication underlies morphological variation of tomato fruit. Science 319: 1527-1530
- **Young LD** (1995) Soybean germplasm resistant to Races 3, 5, or 14 of soybean cyst nematode. Crop Science **35**: 895-896
- Yu N, Lee TG, Rosa DP, Hudson M, Diers BW (2016) Impact of *Rhg1* copy number, type, and interaction with *Rhg4* on resistance to *Heterodera glycines* in soybean. Theor Appl Genet **129:** 2403-2412
- **Zhao M, Ma J** (2013) Co-evolution of plant LTR-retrotransposons and their host genomes. Protein Cell **4:** 493-501
- Zhao M, Wu S, Zhou Q, Vivona S, Cipriano DJ, Cheng Y, Brunger AT (2015) Mechanistic insights into the recycling machine of the SNARE complex. Nature **518**: 61-67

## **Figure Legends**

Fig 1. Multiple rhg1-a haplotypes harbor an intronic copia retrotransposon (RAC) within the Rhg1-encoded a-SNAP (Glyma.18g022500). (A) Diagram of a single 31.2 kb Rhg1 block and the four Rhg1-encoded genes: Glyma. 18G022400 (amino acid permease, AAP), Glyma. 18G022500 (α-SNAP), Glyma. 18G022600 (PLAC-domain protein), and Glyma. 18G022700 (wound-inducible protein, WIP). Glyma. 18G022300 and Glyma. 18G022800 flank Rhg1, but each repeat also includes a truncated 3' fragment of Glyma.18G022300. (B) Schematic of the three known Rhg1 haplotypes: Rhg1 wild-type (single-copy, shown blue), rhg1a (low-copy, shown red) and rhg1-b (high-copy, shown orange). Rhg1 α-SNAP C-terminal amino acid polymorphisms colored to match Rhg1 block type. (C, D) Model from DNA sequencing of *Rhg1* alpha-SNAP copia (*RAC*) integration site within the PI 89772 (*rhg1-a*) encoded  $\alpha$ -SNAP. The 4.77 kb RAC element (shown grey) is anti-sense to  $\alpha$ -SNAP<sub>Rhg1</sub>LC and increases overall repeat size to ~36 kb. RAC ORF is intact and encodes a 1438 residue polyprotein. RAC LTRs shown dark grey,  $\alpha$ -SNAP<sub>Rhg1</sub>LC promoter shown red. "Ex." and "int." are  $\alpha$ -SNAP<sub>Rhg1</sub>LC exons and introns, respectively. LTR: long terminal repeat; GAG: groupspecific antigen, RT: reverse-transcriptase. (E) Agarose gel showing 5' and 3' α-SNAP-RAC junction products from the rhg1-a (low-copy, red dots) accessions: 'Forrest', PI 90763, PI 437654, PI 89772. No α-SNAP-RAC junctions detected from rhg1-b (high-copy; orange dots) accessions: PI 88788, PI 209332, or PI 548316. (F) Similar to E and using same template DNA samples, but PCR amplification of a WT (wild type)  $\alpha$ -SNAP exon 1 – 2 distance, as in the Williams 82 reference genome. WT  $\alpha$ -SNAP exon 1 to exon 2 distances outlined in C by an arrow and \*.

**Fig 2.** The *RAC*-like subfamily of copia retrotransposons is common in soybean and other **legumes.** (A) Maximum likelihood phylogenetic tree of *RAC*-like element nucleotide sequences from soybean. The top hit from each soybean chromosome was included, as was the known soybean retrotransposon "TGMR" and the top *RAC*-like match from *Phaseolus* (common bean). (B) Similar to *A*; a maximum-likelihood tree, but using the *RAC*-encoded polyprotein sequences from the four most similar soybean *RAC*-like elements, and the most similar element matches from the indicated plant species.

Fig 3. RAC is present within a subclass of rhg1-a-signature soybean accessions. (A) Frequency of RAC-associated SNP, ss715606985, among 19,645 SoySNP50K-genotyped USDA soybean accessions. (B) Agarose gel showing PCR detection of  $\alpha$ -SNAP-RAC junctions or WT  $\alpha$ -SNAP exon 1-2 distances among rhg1-a-signature accessions positive or negative for the RAC-SNP, ss71560698. Williams 82 (Rhg1wT), 'Forrest' (rhg1-a) and 'Fayette' (rhg1-b) included as controls; Rhg1 haplotypes color coded with dots as in Fig 1. An \* denotes an rhg1-a-signature

accession lacking the *RAC*-SNP. (C) Frequency of *RAC*-associated SNP among all USDA *Glycine max* accessions with consensus SNP signatures of *rhg1-a* or *rhg1-b* haplotypes.

Fig 4. *RAC* presence correlates with a stronger SCN-resistance profile and the co-presence of other loci that augment *rhg1-a* resistance. (A) Proportion of *RAC*<sup>+</sup> (ss715606985 A SNP) vs. *RAC*<sup>-</sup> (G SNP) accessions among 573 SCN-phenotyped soybeans with consensus SoySNP50K SNP signatures predictive of *rhg1-a*. \*Note that none of the sampled *RAC*<sup>-</sup> (G SNP) accessions had *rhg1-a* (none encoded *α-SNAP<sub>Rhg1</sub>LC*). "S": susceptible in all trials, "MS": moderately susceptible in at least one trial, "MR": moderately resistant in at least one trial, "R": resistant in at least one trial. Fisher's Exact Test pairwise comparisons: "R-MR" (p = 2.6E-4), "R-MS" (p = 2.3E-11), "R-S" (p = 2.2E-16), "MR-MS" (p = 1.0), "MR-S" (p = 0.25), "MS-S" (p = 2.4E-3). (B) Frequency of SNPs associated with *Rhg4* (ss715602757, ss715602764) or the Chromosome 11-encoded α-SNAP-Intron Retention (α-SNAP<sub>Ch11</sub>IR) allele, ss71559743 among 19,645 USDA accessions. (C) Frequency of the *Rhg4* and *α-SNAP<sub>Ch11</sub>IR* associated SNPs among the 51 "Resistant" scored *RAC*<sup>+</sup> *rhg1-a*-signature accessions. (D) Frequency of the *Rhg4* and *α-SNAP<sub>Ch11</sub>-IR* associated SNPs among all *RAC*<sup>+</sup> (300) or *RAC*<sup>-</sup> (403) USDA *G. max* accessions with consensus SNP signatures predictive of *rhg1-a* (705 total; 2 accessions undefined for ss715606985 SNP).

Fig 5. The rhg1-a RAC element is methylated but has intrinsic transcriptional activity. (A) Agarose gel showing PCR amplicons for  $\alpha$ -SNAP-RAC regions from McrBC treated (+) or mock treated (-) genomic DNAs from 'Forrest' (Forr) or 'Peking' (Pek, PI 548402) roots. (B) qPCR analysis of mRNA transcript abundance for RAC and similar RAC-like elements, in leaf or root tissues of Williams 82 (Wm; Rhg1<sub>WT</sub>), 'Forrest' (Forr; rhg1-a) or 'Fayette' (Fay; rhg1-b). Colored dots indicate Rhg1 haplotype as in Fig 1. Normalized RAC transcript abundances are presented relative to the mean abundance of RAC transcript for Williams 82 leaf samples. Y-axis uses  $\log_2$  scale. (C) Schematic showing unique nucleotide tag addition to an otherwise native  $\alpha$ -SNAP-RAC cassette. This construct contains native flanking Rhg1 sequence including Glyma. 18G022400 (transcribes from the bidirectional  $\alpha$ -SNAP promoter) and 1.8 kb upstream, as well as 4.7 kb of downstream RAC flanking sequence ( $\sim$ 1.0 kb after the  $\alpha$ -SNAP<sub>Rhg1</sub>LC termination codon). The RAC region detected and amplified via qPCR or RT-PCR is colored ivory and flanked by half-arrows. (D) Agarose gel of RT-PCR cDNAs of 'Forrest' or Wm 82 transgenic roots transformed with an empty vector (EV) or the native tagged  $\alpha$ -SNAP-RAC construct. Tag primers amplify only the modified  $\alpha$ -SNAP-RAC while the normal RAC primer set amplifies both endogenous RAC-like transcripts as well as the tagged  $\alpha$ -SNAP-RAC transgene. Glyma. 18G022300 mRNA transcript used as a cDNA quality and loading control; no RT (reverse transcriptase) ctrl verifies absence of amplifiable genomic DNA. (E) Schematic showing the subcloned 4.77 kb RAC expression cassette tested in F. (F) Like D, but 'Forrest' or Wm 82

roots transformed with empty vector or the 4.77 kb *RAC* element (all flanking *Rhg1* sequence context removed).

Fig 6. α-SNAP<sub>Rhg1</sub>LC protein is expressed despite *RAC* presence. (A) Schematic showing PIPE-mediated removal of *RAC* from the native α-SNAP-RAC construct, pSM101. (B) Immunoblots of independent 'Forrest' or Wm82 transgenic root lysates using previously described antibodies for α-SNAP<sub>Rhg1</sub>LC or WT α-SNAP proteins. "+" denotes α-SNAP-RAC transformation, "-" indicates α-SNAP<sub>Rhg1</sub>LC (RAC removed) transformed, and EV is empty vector transformed. Ponceau S staining serves as a loading control. (C) Agarose gel showing RT-PCR amplification of mature α-SNAP<sub>Rhg1</sub>LC transcript isoforms from roots of Wm 82 or 'Forrest' transformed with α-SNAP-RAC (+), or a native α-SNAP<sub>Rhg1</sub>LC cassette with RAC removed (-), or an empty vector control. WT refers to primers specific for WT α-SNAP transcripts, LC detects full length α-SNAP<sub>Rhg1</sub>LC transcripts, while "Iso" amplifies a previously described α-SNAP<sub>Rhg1</sub>LC alternative transcript isoform that splices out 36 bp (Cook et al., 2014).

Fig 7. α-SNAP<sub>Rhg1</sub>LC hyperaccumulates at SCN infection sites in low-copy *rhg1-a* soybean accession 'Forrest'. (A) Immunoblot of non-transgenic 'Forrest' root samples from SCN-infested root regions (SCN +) harvested 4 d after SCN infection, or similar regions from mockinoculated controls (SCN -). Blot was probed simultaneously with anti-α-SNAP<sub>Rhg1</sub>LC and anti-NSF polyclonal antibodies. Ponceau S staining before blotting served as a loading control. (B) Representative electron microscope image (7 dpi) showing anti-α-SNAP<sub>Rhg1</sub>LC immunogold signal in SCN-associated syncytium cells from 'Forrest' roots. Arrows highlight only some of the 15 nm immunogold particle dots. Frequent α-SNAP<sub>Rhg1</sub>LC signal was observed in syncytium cells (upper left, "Syn") but rare in adjacent cells (upper right and bottom, "Adj."). CW, cell wall; M, mitochondrion; Vac, vacuole. Bar = 1 μm. (C) Mean and SEM of α-SNAP<sub>Rhg1</sub>LC gold particle abundance in syncytia, normalized to the count from adjacent cells in the same image. Anti-α-SNAP<sub>Rhg1</sub>LC immunogold particles were counted for one 9 μm² area within cells having syncytium morphology and in a region with the highest observable signal in directly adjacent cells with normal root cell morphology (large central vacuole). Data are for 23 images (11 and 12 root sections respectively, from two experiments), for root sections 7 days after inoculation.

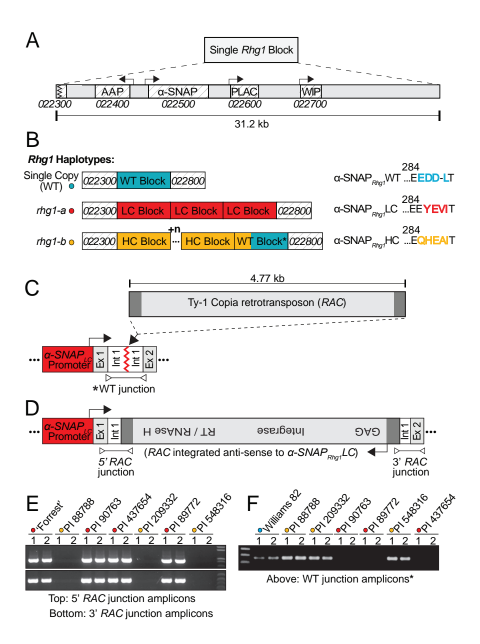
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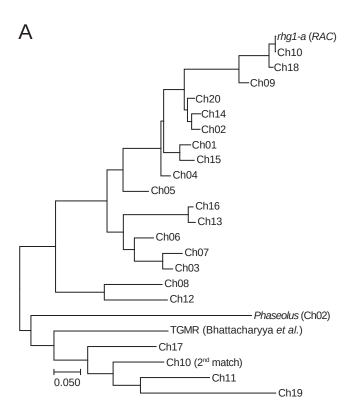
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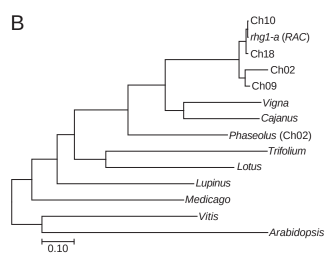
## **Author Contributions**

A.M.B., R.W.Z., S.H. and A.F.B designed the research; A.M.B., R.W.Z., S.H., D.J.G. and K.K.A. performed the research; all authors analyzed data and contributed to writing the paper that was drafted primarily by A.M.B.

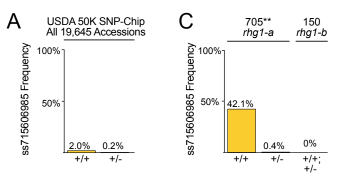




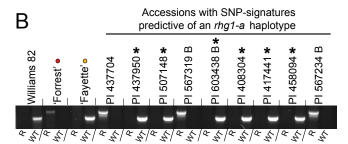
RAC-nucleotide sequence tree from BLASTN matches



RAC-encoded polyprotein tree from TBLASTN matches

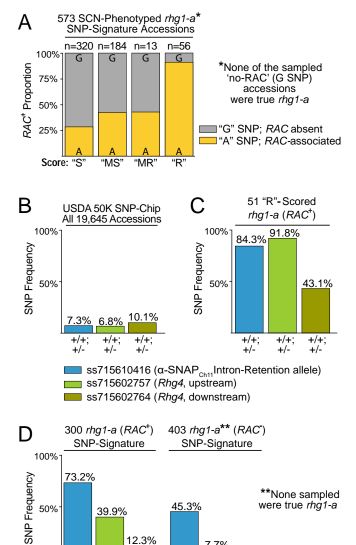


Homo (+/+) or heterozygous (+/-) for ss715606985 (RAC SNP) \*\*SNP undefined in 2 accessions



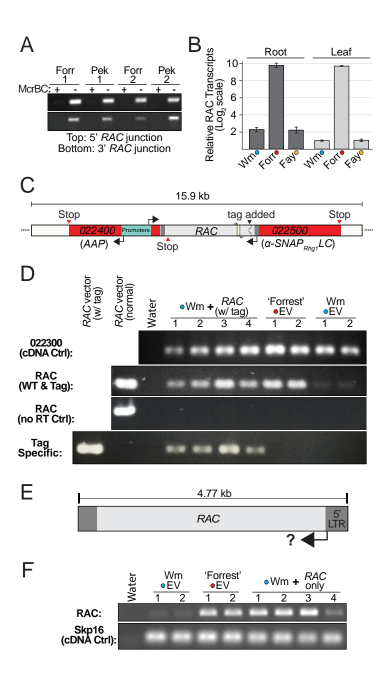
R: RAC- $\alpha$ -SNAP junction WT: Normal  $\alpha$ -SNAP intron 1 (no RAC integration)

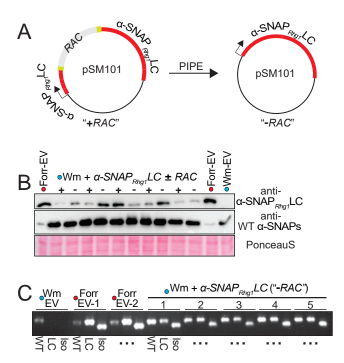
<sup>\*</sup>Lack ss715606985 SNP (RAC-associated)

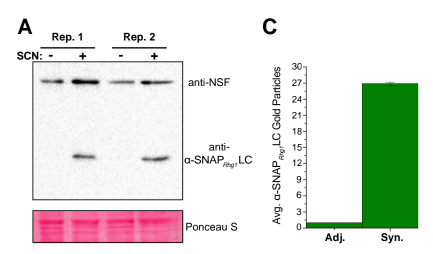


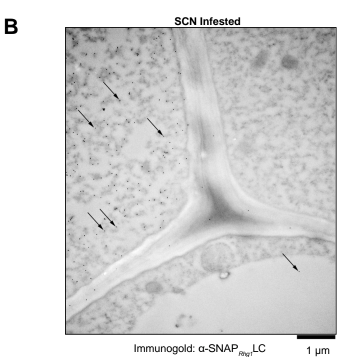
+/+;

+/-









Accession	Rhg1 Copies	SCN Resistant	<i>Rhg1</i> WT α-SNAP	Rhg1 HC α-SNAP	Rhg1 LC α-SNAP	WT Junction	RAC Junctions
Williams 82	1	_	<b>/</b>			<b>✓</b>	
'Forrest'	3	<b>/</b>			<b>✓</b>		<b>/</b>
PI 90763	3	<b>*</b>			<b>✓</b>		<b>~</b>
PI 437654	3	<b>/</b>			<b>\</b>		<b>/</b>
PI 89772	3	<b>✓</b>			<b>✓</b>		<b>/</b>
PI 548316	7	<b>/</b>	<b>/</b>	<b>/</b>		<b>✓</b>	
PI 88788	9	<b>\</b>	<b>/</b>	<b>/</b>		<b>✓</b>	
PI 209332	10	<b>/</b>	<b>/</b>	<b>/</b>		<b>/</b>	