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(54) SOYBEAN GENE FOR RESISTANCE TO ALPHIS GLYCINES

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5.5.e. 15 1(b) by 125 days.

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Related U.S. Application Data

- (62) Division of application No. 11/869,500, filed on Oct.9, 2007, now Pat. No. 7,928,286.
- (60) Provisional application No. 60/829,123, filed on Oct. 11, 2006.

(51)	Int. Cl.	
	A01H 1/00	(2006.01)
	A01H 1/02	(2006.01)
	A01H 5/00	(2006.01)

(52) U.S. Cl.

USPC **800/266**; 800/265; 800/298; 800/312; 800/302; 435/415; 435/6.11

(58) Field of Classification Search

None

See application file for complete search history.

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(57) ABSTRACT

An *Aphis glycines* resistance Rag2 gene is provided herein, along with methods for identifying its presence using marker-assisted selection. A cultivar of *G. max* having resistance to *Aphis glycines* conferred by the Rag2 gene has been identified. The Rag2 gene, as well as the methods, aphid-resistant varieties, and markers disclosed herein may be used to breed new elite lines expressing soybean aphid resistance.

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Kaloshian, I., et al. (1997), "The impace of Meu-1-mediated resistance in tomato on lengevity, fecundity and behavior of the potato aphid," *Macrosiphum euphorbiae, Entomol. Exp. Appl.* 83:181-187. Klinger, J. et al. (2001), "Mapping of cotton-melon aphid resistance in melon," *J. Am. Soc. Hortic. Sci.* 136:56-63.

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FIGURE 1A

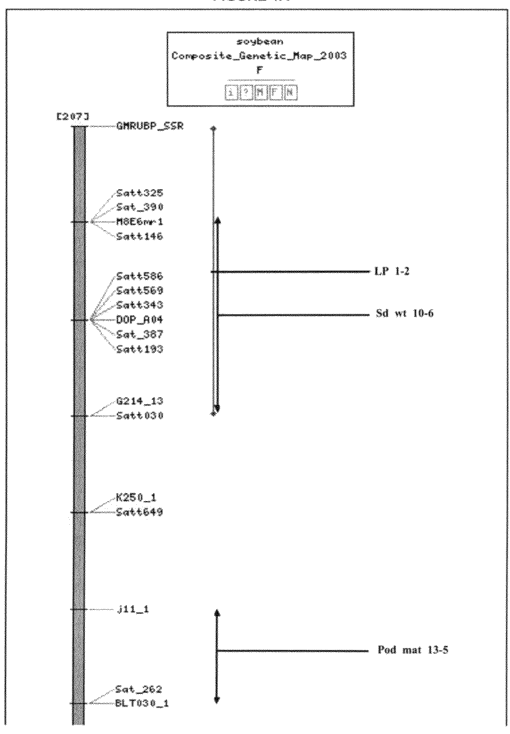
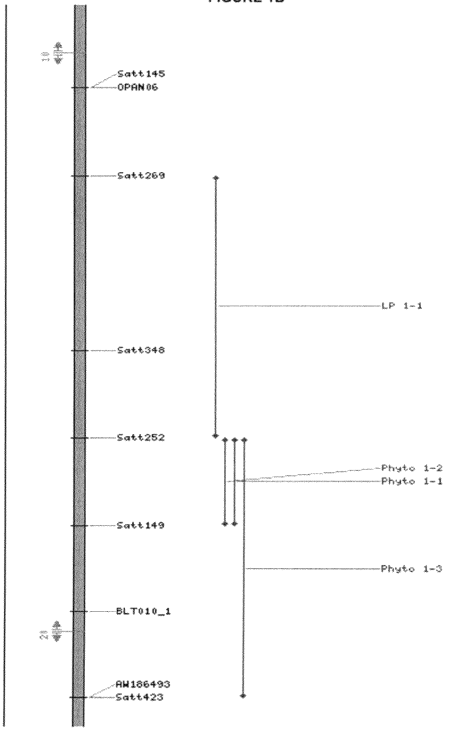


FIGURE 1B



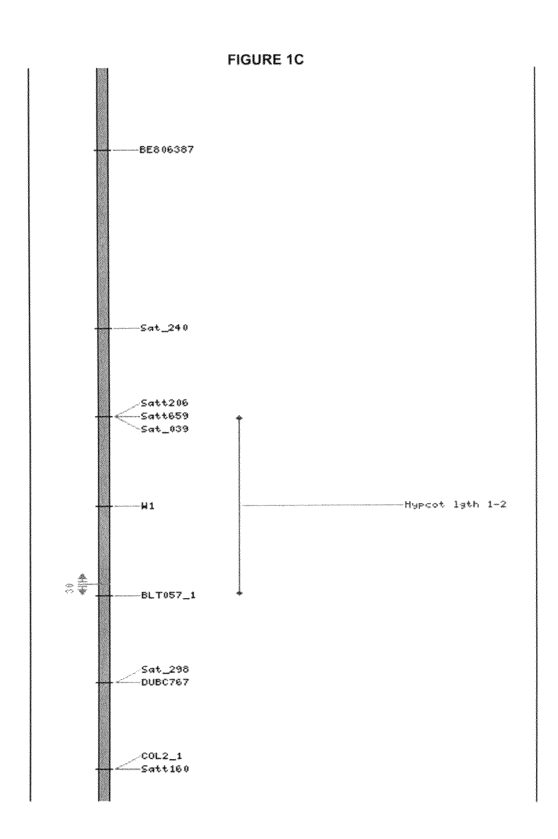
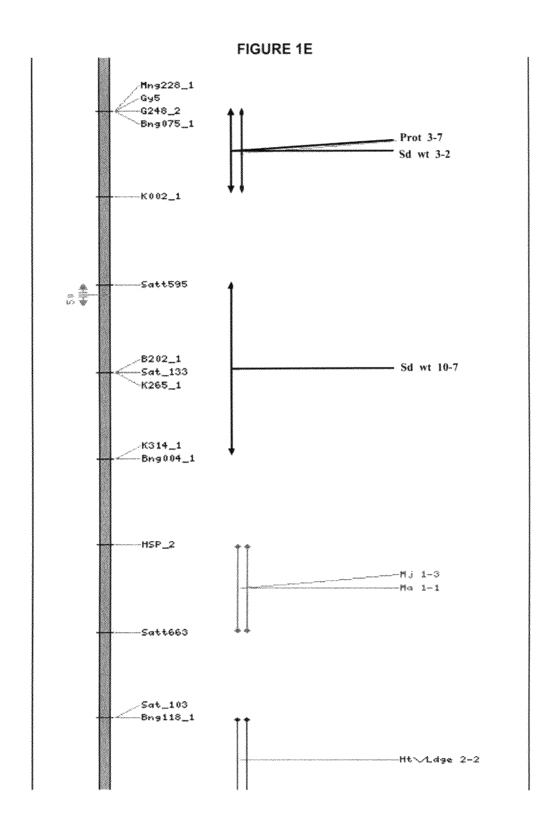
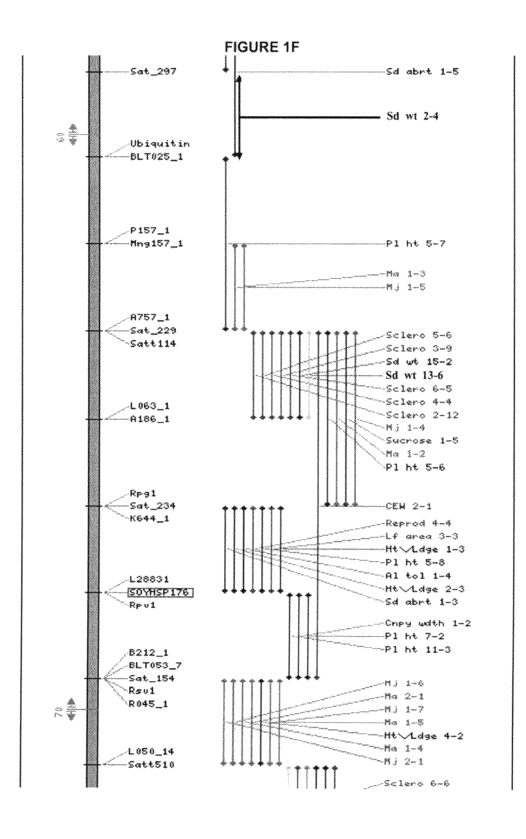
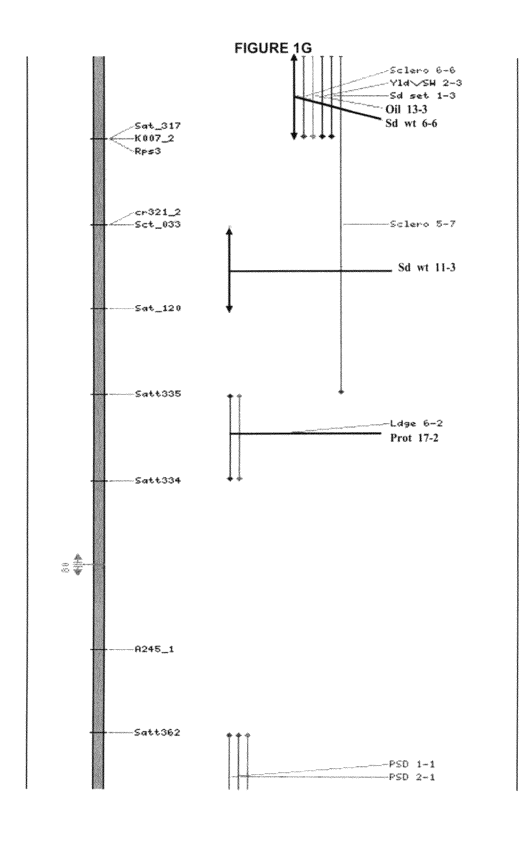
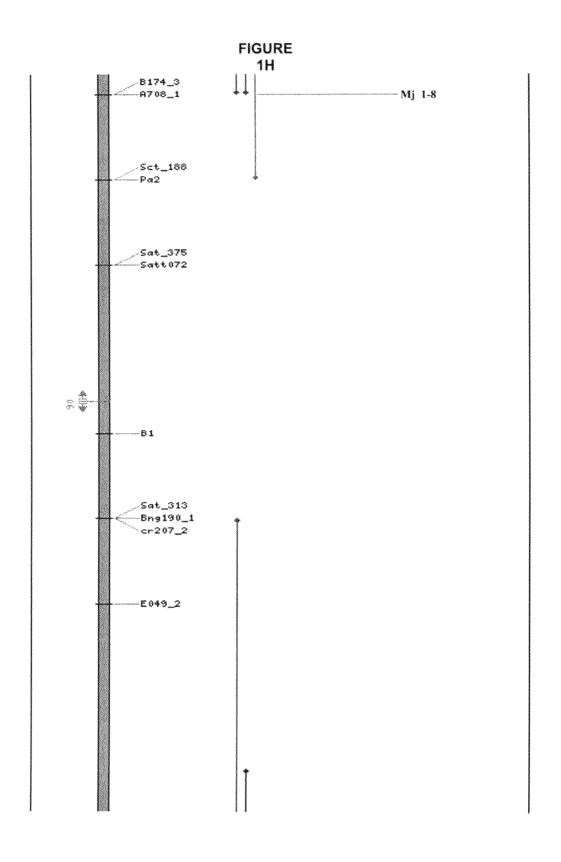


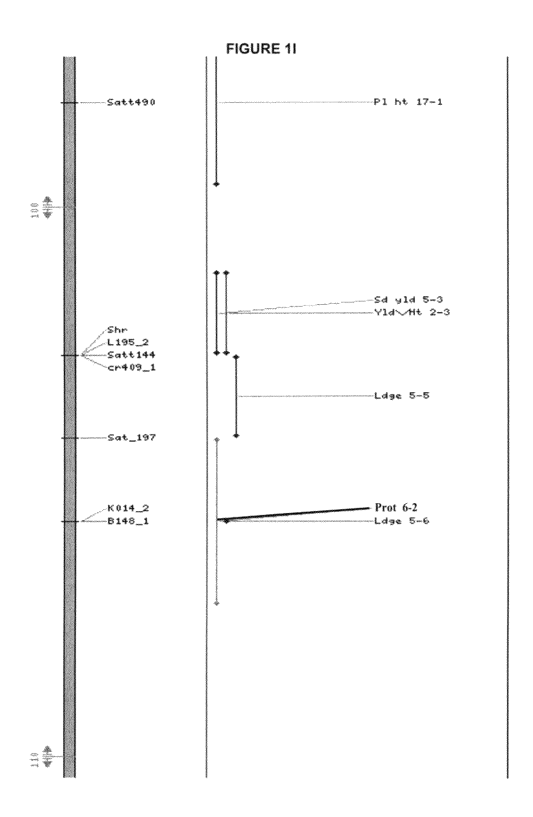
FIGURE 1D - Sd set 1-2 Sd abrt 1-7 A401_1 Lf wdth 1-2 -Cnpy ht 1-2 -Mj 1-2 A806_1 -K390<u>1</u> -Sat_309 -Satt374 Satt516 -Satt425

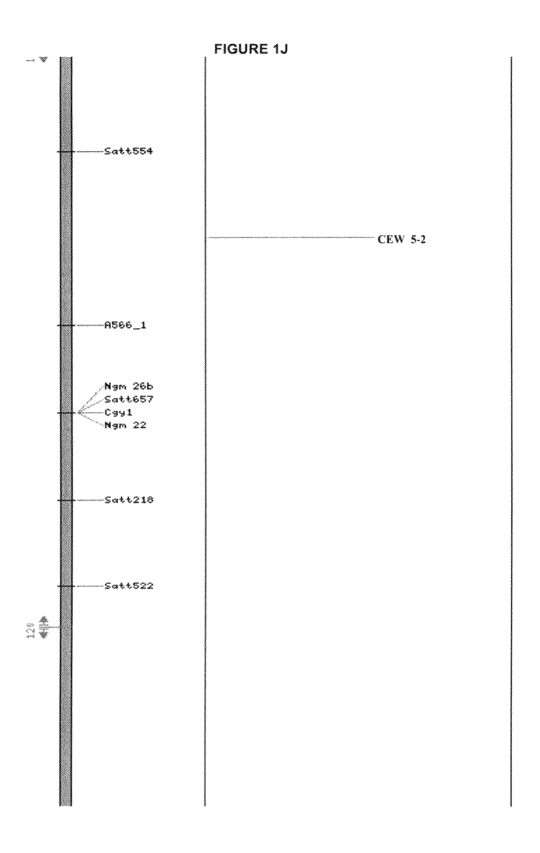


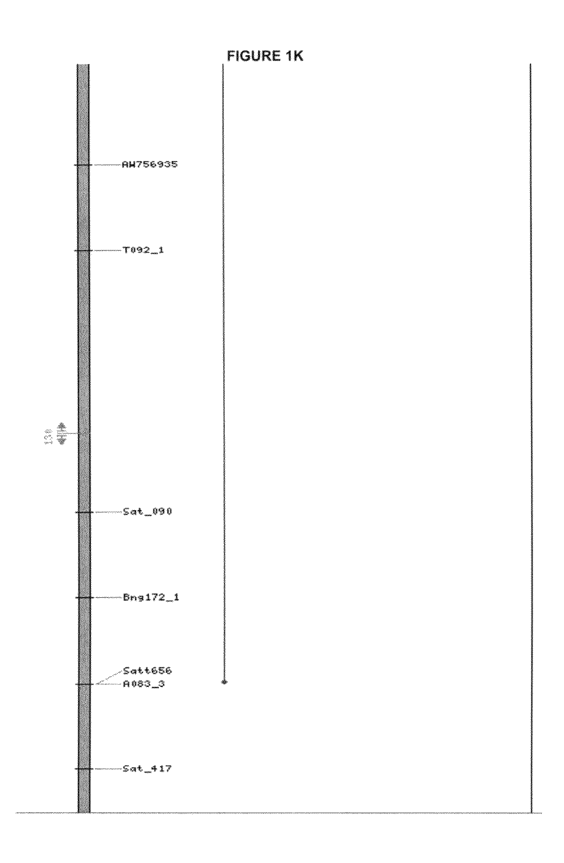












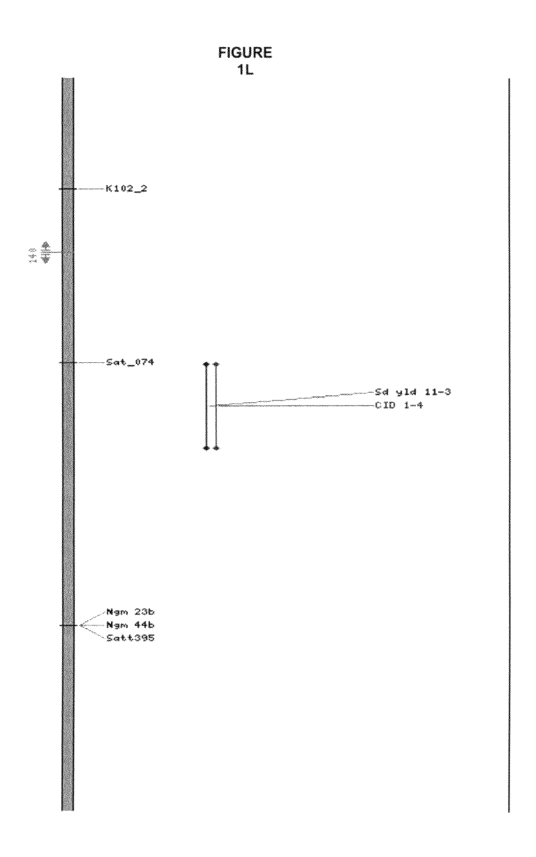
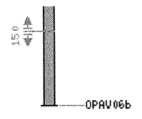


FIGURE 1M



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- Mag + 0-150.97 cM

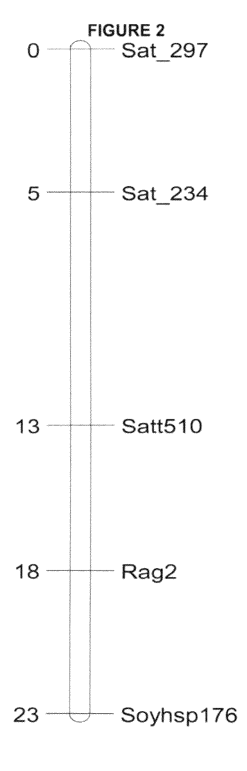
Feature Types:

- Gene PCR
- \$ QTL_fungal
- \$ QTL_inorganic
- \$ QTL_insect
- \$ QTL_leaf-stem
- \$ QTL_misc
- \$ QTL_nematode
- \$ QTL_oil
- Representation Repres
- Report QTL_pod
- \$ QTL_protein
- # QTL_reprod-period
- \$ QTL_whole-plant
- \$ QTL_yield RAPD RFLP SSR

Menu Symbols:

- i Map Set Info
- Map Details
- M Matrix View
- X Delete Map
- F Flip Map
- N New Map View

Chap #0.13



SOYBEAN GENE FOR RESISTANCE TO ALPHIS GLYCINES

CROSS-REFERENCE TO RELATED APPLICATION

This application is a divisional application of U.S. Ser. No. 11/869,500 filed Oct. 9, 2007, now U.S. Pat. No. 7,928,286 issued Apr. 19, 2011, which claims priority to U.S. Provisional Patent Application No. 60/829,123, filed Oct. 11, 2006, 10 both of which prior applications are incorporated herein by reference to the extent not inconsistent herewith.

BACKGROUND

Described herein are a soybean gene for resistance to *Aphis glycines*, soybean plants possessing this gene, which maps to a novel chromosomal locus, and methods for identifying and breeding these plants, the methods involving marker-assisted selection.

Soybeans (*Glycine max* L. Merr.) are a major cash crop and investment commodity in North America and elsewhere. Soybean oil is one of the most widely used edible oils, and soybeans are used worldwide both in animal feed and in human food production.

A native of Asia, the soybean aphid was first found in the Midwest in 2000 (Hartman, G. L. et al., "Occurrence and distribution of Aphis glycines on soybeans in Illinois in 2000 and its potential control," (1 Feb. 2001 available at the "plantmanagementnetwork" org website). It rapidly spread 30 throughout the region and into other parts of North America (Patterson, J. and Ragsdale, D., "Assessing and managing risk from soybean aphids in the North Central States," (11 Apr. 2002) available at the planthealth.info website in subdirectory soyaphid and further subdirectory aphid02. High aphid 35 populations can reduce crop production directly when their feeding causes severe damage such as stunting, leaf distortion, and reduced pod set (Sun, Z. et al., "Study on the uses of aphid-resistant character in wild soybean. I. Aphid-resistance performance of F₂ generation from crosses between culti- 40 vated and wild soybeans," (1990) Soybean Genet. News. 17:43-48). Yield losses attributed to the aphid in some fields in Minnesota during 2001, where several thousand aphids occurred on individual soybean plants, were >50% (Ostlie, K., "Managing soybean aphid," 2 Oct. 2002) available at the 45 soybeans University of Minnesota website under successive subdirectories crop, insects, aphid, aphid_publication_managingsba with an average loss of 101 to 202 kg ha⁻¹ in those fields (Patterson and Ragsdale, supra). In earlier reports from China, soybean yields were reduced up to 52% when there 50 was an average of about 220 aphids per plant (Wang, X. B. et al., "A study on the damage and economic threshold of the soybean aphid at the seedling stage," (1994) Plant Prot. (China) 20:12-13) and plant height was decreased by about 210 mm after severe aphid infestation (Wang, X. B. et al., 55 "Study on the effects of the population dynamics of soybean aphid (Aphis glycines) on both growth and yield of soybean," (1996) Soybean Sci. 15:243-247). An additional threat posed by the aphid is its ability to transmit certain plant viruses to soybean such as Alfalfa mosaic virus, Soybean dwarf virus, 60 and Soybean mosaic virus (Sama, S. et al., "Varietal screening for resistance to the aphid, Aphis glycines, in soybean, (1974) Research Reports 1968-1974, pp. 171-172; Iwaki, M. et al., "A persistent aphid borne virus of soybean, Indonesian Soybean dwarf virus transmitted by Aphis glycines," (1980) 65 Plant Dis. 64:1027-1030; Hartman, G. L. et al., supra; Hill, J. H. et al., "First report of transmission of Soybean mosaic

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virus and Alfalfa mosaic virus by *Aphis glycines* in the New World," (2001) Plant Dis. 561; Clark, A. J. and Perry, K. L., "Transmissibility of field isolates of soybean viruses by *Aphis glycines*," (2002) *Plant Dis.* 86:1219-1222).

Because *A. glycines* is a recent pest in the USA, a comprehensive integrated management approach to control the aphid has yet to be developed. Research to evaluate the efficacy of currently-available insecticides and other control measures has just begun.

An integral component of an integrated pest management (IPM) program to control aphids is plant resistance (Auclair, J. L., "Host plant resistance," pp. 225-265 In P. Harrewijn (ed.) Aphids: Their biology, natural enemies, and control, Vol. C., Elsevier, New York (1989); Harrewijn, P. and Minks, A. K., "Integrated aphid management: General aspects," pp. 267-272, In A. K. Minks and P. Harrewijn (ed.) Aphids: Their biology, natural enemies, and control, Vol. C., Elsevier, New York (1989). Insect resistance can significantly reduce input costs for producers (Luginbill, J. P., "Developing resistant 20 plants—The ideal method of controlling insects," (1969) USDA, ARS. Prod. Res. Rep. 111, USGPO, Washington, D.C. Resistance was reported in G. soja (Sun, Z. et al., "Study on the uses of aphid-resistant character in wild soybean. I. Aphid-resistance performance of F₂ generation from crosses 25 between cultivated and wild soybeans," (1990) Soybean Genet. News 17:43-48), a close relative of G. max (Hymowitz, T., "On the domestication of the soybean," (1970) Econ. Bot. 24:408-421), and other wild relatives (Zhuang, B. et al., "A study on resistance to soybean mosaic virus and Aphis glycines of perennial wild soybean," (1996) Soybean Genet. Newsl. 23:66-69). Prior to 2004, there were no reports of resistance in G. max. A report from Indonesia had indicated that there was no resistance in a test of 201 soybean cultivars and breeding lines (Sama, S. et al. (1974) Research Reports 1968-1974, p. 171-172. In Varietal screening for resistance to the aphid, Aphis glycines, in soybean. Agricultural Cooperation, Indonesia, the Netherlands).

There are numerous examples of the discovery and use of resistance genes to control aphids in crops other than soybean. Examples include Russian wheat aphid (Du Toit, F. (1987), "Resistance in wheat (*Triticum aestivum*) to *Diuraphis noxia* (Homoptera:Aphididae)," *Cereal Res. Commun.* 15:175-179; wheat greenbug (Tyler, J. M., et al. (1985), "Biotype E greenbug resistance in wheat streak mosaic virus-resistant wheat germplasm lines," Crop Science 25:686-688), potato aphid on tomato (Kaloshian, I., et al. (1997), "The impact of Meu-1-mediated resistance in tomato on longevity, fecundity and behavior of the potato aphid," *Macrosiphum euphorbiae,*" *Entomol. Exp. Appl.* 83:181-187), and cotton-melon aphid on melon (Klinger, J. et al. (2001), "Mapping of cotton-melon aphid resistance in melon," *J. Am. Soc. Hortic. Ci.* 136:56-63).

A number of soybean markers have been mapped and linkage groups created, as described in Cregan, P. B., et al., "An Integrated Genetic Linkage Map of the Soybean Genome" (1999) Crop Science 39:1464-1490.

U.S. Patent Publication 2006/0014964, Hill, C. B., et al. (2006), "Soybean aphid resistance in soybean Jackson is controlled by a single dominant gene," Crop Science 46:1606-1608, and Hill, C. B., et al. (2006), "A single dominant gene for resistance to the soybean aphid in the soybean cultivar Dowling," Crop Science 46:1601-1605 disclose two previously-discovered soybean aphid resistance genes, Rag₁ in Dowling and another gene in Jackson.

A trait that maps to soybean Linkage Group F is root-knot nematode resistance. (Tamulonis, J. P., et al. (1997), "DNA marker analysis of loci conferring resistance to peanut root-

knot nematode in soybean," Theor. Appl. Genet. 95:664-670.) Jeong, S. C. et al., "Cloning And Characterization Of An Rga Family From The Soybean Molecular Linkage Group F," in an Abstract published by Plant & Animal Genome VIII Conference, Town & Country Hotel, San Diego, Calif., Jan. 9-12, 52000 at a website address with the usual www prefix followed by intl-pag.org/8/abstracts/pag8255.html and in Yong G. Yu, Glenn R. Buss, and M. A. Saghai Maroof (1996), "Isolation of a superfamily of candidate disease-resistance genes in soybean based on a conserved nucleotide-binding site," PNAS, 1093:11751-11756, discloses that the soybean chromosomal region on linkage group F flanked by the markers K644 and B212 contains several virus, bacteria, fungus and nematode resistance genes.

Conventional plant breeding for insect resistance tradition- 15 ally relied on screening whole plants for resistance directly with live insects and assessing insect population development or plant damage caused by insect feeding, or indirectly with techniques that measure insect feeding behavior, such as Electrical Penetration Graph (EPG). Implementation of these 20 techniques requires a certain amount of time and specialized space, such as in a greenhouse or plant growth room. More efficient and cost-effective molecular genetic and polymerase chain reaction (PCR) techniques, with the development of DNA markers, enable breeders to significantly increase 25 throughput and efficiency in screening plants for traits that are tightly linked to DNA markers, by screening genomic DNA of plants in the laboratory. There are numerous examples of the use of this technology to select plants with certain traits in breeding programs, including insect resistance. Other publi- 30 cations directed to marker-identification of soybean aphid resistance include Li, Y, et al., "Soybean aphid resistance genes in the soybean cultivars Dowling and Jackson map to linkage group M," Molecular Breeding (in press); Hill, C. B., et al. (2006), "Soybean aphid resistance in soybean Jackson is 35 controlled by a single dominant gene," Crop Science 46:1606-1608; Hill, C. B., et al. (2006), "A single dominant gene for resistance to the soybean aphid in the soybean cultivar Dowling," Crop Science 46:1601-1605; Li, Y., et al. (2004) "Effect of three resistant soybean genotypes on the 40 fecundity, mortality, and maturation of soybean aphid (Homoptera: Aphididae)," Journal of Economic Entomology 97:1106-1111; Hill, C. B., et al. (2004) "Resistance to the soybean aphid in soybean germplasm and other legumes," p. 179, World Soybean Research Conference, Foz do Iguassu, 45 PR, Brazil; Hill, C. B., et al. (2004), "Resistance to the soybean aphid in soybean germplasm," Crop Science 44:98-106; and Hill, C. B., et al. (2004), "Resistance of Glycine species and various cultivated legumes to the soybean aphid (Homoptera: Aphididae)," Journal of Economic Entomology 50 97:1071-1077). Additional methods and molecular tools are needed to allow breeding of A. glycines resistance into highyielding G. max soybean varieties.

All publications referred to herein are incorporated herein by reference to the extent not inconsistent herewith.

SUMMARY

A method is provided for determining the presence or absence in a soybean germplasm of a gene for resistance to 60 the soybean aphid, *Aphis glycines*. The aphid resistance trait has been found to be closely linked to a number of molecular markers that map to linkage group F. The gene conferring the resistance trait is designated "Rag2" pending approval of the Soybean Genetics Committee. The Rag2 gene was originally 65 discovered in the resistance source Sugao Zairai (PI200538). ("PI" stands for "plant introduction" and this PI number

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refers to the USDA depositary accession number.) The trait of resistance to *Aphis glycines* is also found in other varieties as described hereafter.

The Rag2 gene, is non-allelic with the Rag1 gene previously found in the soybean cultivar Dowling (Hill, C. B. et al., (2006), "A single dominant gene for resistance to the soybean aphid in the soybean cultivar Dowling," Crop Science 46:1601-1605). Similar to Rag1, when present in soybean plants, the Rag2 gene conditions strong resistance to the soybean aphid by preventing aphid colonization on plants through reduced aphid multiplication, survival, lifespan, and development of nymphs to adults. Expression of resistance is dominant over susceptibility in heterozygous plants containing both forms of the gene. Resistance controlled by Rag2 is effective against all known soybean aphid biotypes.

The location of the Rag2 gene was mapped to linkage group F on the soybean genetic map and it is closely flanked by two DNA markers called simple sequence repeats (SSR), namely Soyhsp176 and Satt510, which are tightly linked to the gene. The tight linkage of the two DNA markers with Rag2 enables soybean breeders to efficiently identify plants that have the soybean aphid resistance gene in progeny of their crosses without having to inoculate plants with aphids.

Use of the technology to identify the presence of the Rag2 gene facilitates and expedites the development of new soybean aphid-resistant cultivars using conventional breeding methods without genetic engineering, by back crossing the Rag2 gene into current, adapted soybean cultivars, converting them to new soybean aphid resistant soybean cultivars. This technology, combined with the technology to identify Rag1 and the related gene covered in U.S. Patent Publication No. 20060015964 enables the development of soybean cultivars with more than one resistance gene to maximize resistance to the soybean aphid.

In accordance with the present method, the Rag2 gene for resistance to Aphis glycines co-segregates with molecular markers with which it is linked on linkage group F, most preferably, Satt510 and Soyhsp176. Additional markers that are also useful for identifying the presence of the Rag2 gene include Sat_120, Sat_234, and Sat_297. The Rag2 gene has been found to map to a locus that lies between the markers Satt510 and Soyhsp176. Other markers of linkage group F may also be used to identify the presence or absence of the gene. Preferably flanking markers are used for identifying the presence of the Rag2 gene for marker-assisted breeding. In one embodiment, the markers used map within about 20 cM, and preferably within about 3 cM to about 10 cM of the Rag2 gene locus (which contains the Rag2 gene), or within about 20 cM and preferably within about 3 cM to about 10 cM of Satt510 or Soyhsp176.

The information disclosed herein regarding Rag2 locus is used to aid in the selection of breeding plants, lines and populations containing *Aphis glycines* resistance for use in introgression of this trait into elite soybean germplasm, i.e., germplasm of proven genetic superiority suitable for cultivar release.

Also provided is a method for introgressing a soybean *Aphis glycines* resistance gene into non-resistant soybean germplasm or resistant soybean germplasm that is more or less resistant than that of PI200538. According to the method, nucleic acid markers linked to the Rag2 gene are used to select soybean plants containing a Rag2 locus. Plants so selected have a high probability of expressing the trait *Aphis glycines* resistance. Plants so selected can be used in a soybean breeding program. Through the process of introgression, the Rag2 gene locus is introduced from plants identified using marker-assisted selection into other plants. According

to the method, agronomically desirable plants and seeds can be produced containing the Rag2 gene locus from germplasm containing the Rag2 gene.

The Rag2 gene locus is defined as the DNA between flanking markers Satt510 and Soyhsp176.

Particular examples of sources of Rag2 resistance (aphid resistance conferred by the Rag2 gene) are provided by soybean cultivar Sugao Zarai (PI200538) and progeny thereof carrying the Rag2 gene locus.

Also provided herein is a method for producing an inbred soybean plant adapted for conferring, in hybrid combination, Aphis glycines resistance. First, donor soybean plants for a parental line containing the Rag2 gene are selected. According to the method, selection can be accomplished via nucleic acid marker-associated selection as explained herein. Selected plant material may come from, among others, an inbred line, a hybrid, a heterogeneous population of soybean plants, or simply an individual plant. According to techniques well known in the art of plant breeding, this donor parental 20 line is crossed with a second parental line. Preferably, the second parental line is high yielding. This cross produces a segregating plant population composed of genetically heterogeneous plants. Plants of the segregating plant population are Rag2 gene locus are selected for further breeding until a line is obtained that is homozygous for resistance to Aphis glycines at the Rag2 locus. This further breeding may include, among other techniques, additional crosses with other lines, hybrids, backcrossing, or self-crossing. The result is an inbred line of soybean plants that are resistant to Aphis glycines and also have other desirable traits from one or more other inbred lines.

The method can also include producing inbred lines having 35 both Rag trait resistance from Rag trait loci on linkage group M as described in U.S. Patent Publication No. 20060015964 (including Rag1 aphid resistance) and Rag2 aphid resistance from linkage group F, as well as traits derived from elite soybean lines. This method comprises crossing soybean 40 plants having Rag2 resistance with soybean plants having Rag1 gene resistance and additional Rag gene resistance conferred by a gene or gene found on linkage group M, and testing for the presence of the aphid resistance traits from both linkage groups F and M using marker-assisted selection, and 45 then making additional crosses with elite lines. As is known in the art, the aphid resistance traits from linkage groups F and M can be stacked in this manner, along with other desirable traits from the elite line(s), into a new soybean cultivar with the intention to increase the durability and effective lifetime of the aphid resistance trait by increasing the difficulty and time for the soybean aphid to produce genetic variants that can overcome both resistance genes.

Soybean plants, seeds, tissue cultures, variants and mutants having Aphis glycines resistance produced by the foregoing methods are also provided herein.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 shows a published soybean genetic linkage F composite map and anchored markers. The map has been broken into thirteen consecutive vertical sections, FIG. 1A through FIG. 1M.

FIG. 2 is a linkage map of a portion of soybean linkage 65 group F (LGF) showing the locations of the soybean aphid resistance gene Rag2. The location of Sat_297 has been

designated as 0, measuring from which the location of Rag2 is shown at 18 cM, midway between Satt510 at 13 cM and Soyhsp176 at 23 cM.

DETAILED DESCRIPTION

"Allele" is any of one or more alternative forms of a gene, all of which alleles relate to one trait or characteristic. In a diploid cell or organism, the two alleles of a given gene occupy corresponding loci on a pair of homologous chromosomes.

"Backcrossing" is a process through which a breeder repeatedly crosses hybrid progeny back to one of the parents (recurrent parent), for example, a first generation hybrid F₁ with one of the parental genotypes of the F₁ hybrid.

"Cultivar" and "cultivar" are used synonymously and mean a group of plants within a species (e.g., Glycine max) that share certain genetic traits that separate them from the typical form and from other possible varieties within that species. Soybean cultivars are inbred lines produced after several generations of self-pollination. Individuals within a soybean cultivar are homogeneous, nearly genetically identical, with most loci in the homozygous state.

"Gene" means a specific sequence of nucleotides in DNA screened for the Rag2 gene locus. Those plants having the 25 that is located in the germplasm, usually on a chromosome, and that is the functional unit of inheritance controlling the transmission and expression of one or more traits by specifying the structure of a particular polypeptide or controlling the function of other genetic material. In the present instance, the Rag2 gene for resistance to Aphis glycines has been found on major soybean linkage group F flanked by markers Satt510 and Soyhsp176. The Rag2 gene may be isolated by one skilled in the art of genetic manipulation without undue experiments by means known to this art including PCR cloning utilizing the adjacent Satt510 and Soyhsp176 primer sequences, or primer sequences from other markers flanking the gene as described herein, by positional cloning using BACs (bacterial artificial chromosomes), or other methods. See, e.g., Wu, et al., "A BAC and BIBAC-based Physical Map of the Soybean Genome" (2004) Genome Res. February; 14(2):319-26, which describes the use of BACs in mapping the soybean genome. Contiguous BACs lying between Soyhsp176 and Satt510, and in which the Rag2 gene is present, may be found in BAC libraries known to the art, such as The Soybean GBrowse Database.

> "Germplasm" means the genetic material with its specific molecular and chemical makeup that comprises the physical foundation of the hereditary qualities of an organism. As used herein, germplasm includes seeds and living tissue from which new plants may be grown; or, another plant part, such as leaf, stem, pollen, or cells, that may be cultured into a whole plant. Germplasm resources provide sources of genetic traits used by plant breeders to improve commercial cultivars.

"Hybrid plant" means a plant offspring produced by cross-55 ing two genetically dissimilar parent plants.

"Inbred plant" means a member of an inbred plant strain that has been highly inbred so that all members of the strain are nearly genetically identical.

"Introgression" means the entry or introduction by hybridization of a gene or trait locus from the genome of one plant into the genome of another plant that lacks such gene or trait

"Molecular marker" is a term used to denote a nucleic acid or amino acid sequence that is sufficiently unique to characterize a specific locus on the genome. Examples include restriction fragment length polymorphisms (RFLPs) and simple sequence repeats (SSRs). RFLP polymorphisms are

found when base substitutions, additions, deletions or sequence rearrangements occur between restriction endonuclease recognition sequences. The size and number of fragments generated by one such enzyme is therefore altered. A probe that hybridizes specifically to DNA in the region of 5 such an alteration can be used to rapidly and specifically identify a region of DNA that displays allelic variation between two plant varieties. SSR markers occur where a short sequence displays allelic variation in the number of repeats of that sequence. Sequences flanking the repeated sequence can 10 serve as polymerase chain reaction (PCR) primers. Depending on the number of repeats at a given allele of the locus, the length of the DNA segment generated by PCR will be different in different alleles. The differences in PCR-generated fragment size can be detected by gel electrophoresis. Other 15 types of molecular markers are known. All are used to define a specific locus on the soybean genome. Large numbers of these have been mapped. Each marker is therefore an indicator of a specific segment of DNA, having a unique nucleotide sequence. The map positions provide a measure of the relative 20 positions of particular markers with respect to one another. When a trait is stated to be linked to a given marker it will be understood that the actual DNA segment whose sequence affects the trait generally co-segregates with the marker. More precise and definite localization of a trait can be obtained if 25 markers are identified on both sides of the trait. By measuring the appearance of the marker(s) in progeny of crosses, the existence of the trait can be detected by relatively simple molecular tests without actually evaluating the appearance of the trait itself, which can be difficult and time-consuming, 30 requiring growing up of plants to a stage where the trait can be expressed.

Another type of molecular marker is the random amplified polymorphic DNA (RAPD) marker. Chance pairs of sites complementary to single octa- or decanucleotides may exist 35 in the correct orientation and close enough to one another for PCR amplification. With some randomly chosen decanucleotides no sequences are amplified. With others, the same length products are generated from DNAs of different indifor every individual in a population. The variable bands are commonly called random amplified polymorphic DNA (RAPD) bands.

Another type of molecular marker is the target region amplification polymorphism (TRAP) marker. The TRAP 45 technique employs one fixed primer of known sequence in combination with a random primer to amplify genomic frag-

A further type of molecular marker is the single nucleotide polymorphism (SNP) marker, in which DNA sequence varia- 50 tions that occur when a single nucleotide (A, T, C, or G) in the genome sequence is altered are mapped to sites on the soy-

Other molecular markers known to the art, as well as phenotypic traits may be used as markers in the methods 55 described herein.

"Linkage" is defined by classical genetics to describe the relationship of traits that co-segregate through a number of generations of crosses. Markers on the same chromosome are linked to one another, meaning that they are inherited as a unit 60 unless there is recombination between markers. Genetic recombination occurs with an assumed random frequency over the entire genome. Genetic maps are constructed by measuring the frequency of recombination between pairs of traits or markers. The closer the traits or markers lie to each 65 other on the chromosome, the lower the frequency of recombination, the greater the degree of linkage. Traits or markers

are considered herein to be linked if they generally co-segregate. A 1/100 probability of recombination per generation is defined as a map distance of 1.0 centimorgan (1.0 cM). Preferably, markers useful for screening for the presence of Rag2 Aphis glycines resistance map to within 20 cM of the trait, and more preferably within 10 cM of the trait.

A second marker that maps to within 20 cM of a first marker that co-segregates with the Rag2 trait and generally co-segregates with the Rag2 trait is considered equivalent to the first marker. Any marker that maps within 20 cM and more preferably 10 cM of the Rag2 trait belongs to the class of preferred markers for use in screening and selection of soybean germplasm having the Rag2 Aphis glycines resistance trait. A number of markers are known to the art to belong to linkage group F on which the Rag trait is found. A number of markers are proprietary markers known only to certain of those skilled in the art of soybean plant breeding. A proprietary marker mapping within 20 cM, and preferably within 10 cM, of any publicly known marker specified herein is considered equivalent to that publicly-known marker.

"Linkage group" refers to traits or markers that generally co-segregate. A linkage group generally corresponds to a chromosomal region containing genetic material that encodes the traits or markers.

"Locus" means a chromosomal region where a polymorphic nucleic acid or trait determinant or gene is located.

"Polymorphism" means a change or difference between two related nucleic acids. A "nucleotide polymorphism" refers to a nucleotide that is different in one sequence when compared to a related sequence when the two nucleic acids are aligned for maximal correspondence. A "genetic nucleotide polymorphism" refers to a nucleotide that is different in one sequence when compared to a related sequence when the two nucleic acids are aligned for maximal correspondence, where the two nucleic acids are genetically related, i.e., homologous, for example, where the nucleic acids are isolated from different strains of a soybean plant, or from different alleles of a single strain, or the like.

"Marker assisted selection" means the process of selecting viduals. With still others, patterns of bands are not the same 40 a desired trait or desired traits in a plant or plants by detecting one or more nucleic acid polymorphisms from the plant, where the nucleic acid polymorphism is linked to the desired trait.

> "Plant" means plant cells, plant protoplast, plant cell or tissue culture from which soybean plants can be regenerated, plant calli, plant clumps and plant cells that are intact in plants or parts of plants, such as seeds, pods, flowers, cotyledons, leaves, stems, buds, roots, root tips and the like.

> "Probe" means an oligonucleotide or short fragment of DNA designed to be sufficiently complementary to a sequence in a denatured nucleic acid to be probed and to be bound under selected stringency conditions.

> "Rag2-derived resistance" means resistance in a soybean germplasm to Aphis glycines that is provided by the heterozygous or homozygous expression of the Rag2 gene within the Rag2 locus mapped between the SSR markers Satt510 and

> "Rag phenotype" means resistance to Aphis glycines by soybean germplasm, as demonstrated by resistance to Aphis glycines after inoculation with same according to the methods described herein. Rag2 phenotype means such aphid resistance conferred by the Rag2 gene.

> "Rag soybean plant" means a plant having resistance to Aphis glycines that is derived from the presence and expression of at least one Rag gene, or that is shown to have a Rag gene. Rag2 soybean plant means a plant having such aphid resistance conferred by the Rag2 gene.

"Self-crossing or self-pollination" is a process through which a breeder crosses hybrid progeny with itself, for example, a second generation hybrid F_2 with itself to yield progeny designated $F_{2:3}$, meaning the progeny from an individual F_2 generation plant.

As used herein, the terms "segregate," "segregants," "co-segregate," "hybrid," "crossing," and "selfing" refer to their conventional meanings as understood in the art (see, for instance, Briggs, F. N. and Knowles, P. F. and, *Introduction to Plant Breeding* (Reinhold Publication Corp., New York, N.Y., 10 1967).

Markers that "flank" the Rag2 gene are markers that occur one to either side of the Rag2 gene. Flanking marker DNA sequences may be part of the gene or may be separate from the gene.

The method for determining the presence or absence of the Rag2 gene, which confers resistance to the soybean aphid *Aphis glycines* in soybean germplasm, comprises analyzing genomic DNA from a soybean germplasm for the presence of at least one molecular marker, wherein at least one molecular 20 marker is linked to the Rag2 trait locus, and wherein the Rag2 trait locus maps to soybean major linkage group F and is associated with resistance to the soybean aphid *Aphis glycines*. The term "is associated with" in this context means that the Rag2 locus containing the Rag2 gene has been found, 25 using marker-assisted analysis, to be present in soybean plants that show or are capable of showing resistance to *Aphis glycines* in live aphid bioassays as described herein.

Aphis glycines resistance associated with the Rag2 gene was found in PI200538 and can also occur in the following 30 soybean germplasm accessions that are resistant to all known soybean aphid biotypes: PI71506; PI88508, Showa No. 1-4; PI230977; PI437696, San-haj-hun-mao-huan-dou; PI499955, PI507298, Sokoshin (Kamigoumura); PI518726, Bao jiao huang; PI548237, T260H; PI548409, Sato; 35 PI567391, Jiang se huang dou; PI567541B; PI567598B; 587552, Nan jing da ping ding huang yi 1; PI587617, Jin tan qing zi; PI587656, Huang dou; PI587663, Zhong chun huang dou; PI587666, Er dao zao; PI587669, Zan zi bai; PI587677, Xiao li huang; PI587685, Da li huang 2; PI587693, Yu shan 40 dou; PI587702, Qing pi dou; PI587717, Xiang yang ba yue zha; PI587732, Ying shan ji mu wo; PI587759, Song zi ba yue cha; PI587763, Jing huang 36; PI587775, Tong shan si ji dou; PI587800, Ying shan da li huang; PI587816, Bai mao dou; PI587824, Ying shan qing pi cao; PI587840, Du wo dou; 45 PI587861, Da qing dou; PI587870, Huang pi dou; PI587871, Bao mao dou; PI587873, Feng wo dou; PI587876, Xi mao dou; PI587897, Qing pi dou; PI587899, Ba yue bai; PI587905, Xiao huang dou; PI587972, Chang zi dou; PI588000, Shi yue huang; PI588040, Shan xing dou; 50 PI594421, Da du huang dou; PI594425, Xiao cao huang dou; PI594431, Chang pu qing dou; PI594499, Luo ma aluo; PI594503, Mu gu hei chi huang dou; PI594514, Hua lian dou, PI594554, Huang pi tian dou; PI594573, Lu pi dou; PI594592, Shi yue xiao huang dou; PI594595, Ba yue da 55 huang dou (jia); PI594703, Qing pi dou -1; PI594707, Da hei dou; PI594822, Xi huang dou; PI594868, Huang dou; and PI594879, Huo shao dou. The Rag2 gene can also be found in progeny of the foregoing varieties and in other varieties by methods set forth herein.

Other sources of *A. glycines* resistance include the *G. max* varieties: PI87059, Moyashimame; PI417084A, Kumaji 1; PI508294; PI548445, CNS; PI548480, Palmetto; PI548657, Jackson; PI548663, Dowling; PI567543C; PI567597C; PI587553A; PI587559B, Dan to he shang tou jia; PI587664B, 65 Shan zi bai; PI587668A, Hui mei dou; PI587674A, Ba yue bai; PI587682A, Da li huang 1; PI587684A, Ai jiao huang;

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PI587686A, Xi li huang 1; PI587687A Xiao li dou 1; PI587700A, Da qing dou; PI587723A, Ying shan ji mu wo; PI587844C, Tong cheng hei se dou; PI587863B, Liu yue bai; PI587877A, Jiu yue zao; PI587891A, Qi yue ba; PI594426A, Tie jiao huang; PI594426B, Tie jiao huang; PI594427A, Ba yuemang; PI594557B, Lao shu dou; PI594560B, Xia shui huang; PI594586A; PI594666B, Liu yue mang 5; PI594711B, Qing huang za dou 3; PI594751A, Long zhou dong feng dou; PI594864, Yang yan dou; PI603521; PI603530A; PI603538A; PI603640; PI603644; PI603655; PI603650; PI605771; PI605823; PI605855; and PI605902, and progeny thereof. G. soja varieties: G3; JS1; L4; PI518282, S12 Taichung 38; PI518281, Taichung 37; PI573059, and PI573071 and progeny of these varieties, are also sources of A. glycines resistance. These varieties may contain the Rag1 and related aphid resistance gene on linkage group M, and/or can contain the Rag2 gene, or different soybean aphid resistance genes. Resistance that is controlled by Rag1 or Rag2 in these and other varieties can be confirmed by marker-assisted selection as described herein.

Any one of the foregoing varieties or their progeny bearing a Rag gene may be used in the methods described herein, and any combination thereof is considered to be a class of varieties useful in the methods provided herein.

Preferably a marker used to determine the presence or absence of a Rag gene is selected from the group consisting of Satt510, Soyhsp176, Sat_234, Sat_297, and any marker that maps to within at least about 10 to about 20 cM of any of said markers.

Any marker assigned to soybean linkage group F may be useful for this purpose. Exemplary markers of linkage group F include: Satt510, R045_1, Rsv1, Sat_154, BLT053_7, B212_1, Rpv1, Soyhsp176 and L28831, and markers that map within about 3 to about 10 cM, or in another embodiment, within about 10 to about 20 cM, of any of the foregoing.

A further class of markers useful in the present methods include: Ubiquitin, BLT025_1, P157_1, Mng157_1, A757_1, Sat_229, Satt114, L063_1, A186_1, Rpg1, Sat_234, K644_1, L28831, Soyhsp176, Rpv1, B212_1, BLT053_7, Sat_154, Rsv1, R045_1, L050_14, Satt510, Sat_317, K007_2, Rps3, cr321_2, Sct033, Sat_120, Satt335, and Satt334, and markers that map within about 10 to about 20 cM of any of the foregoing.

A further class of markers useful in the present methods include: A401_1, A806_1, K390_1, Sat_309, Satt374, Satt516, Satt425, Mng228_1, Gy5, G248_2, Bng075_1, K002_1, Satt595, B202_1, Sat_133, K265_1, K314_1, Bng004_1, HSP_2, Satt663, Sat 103, Bng118_1, Sat_297, Ubiquitin, BLT025_1, P157_1, Mng157_1, A757_1, Sat_229, Satt114, L063_1, A186_1, Rpg1, Sat_234, K644_1, L28831, Soyhsp176, Rpv1, B212_1, BLT053_7, Sat_154, Rsv1, R045_1, L050_14, Satt510, Sat_317, K007_2, Rps3, cr321_2, Sct033, Sat_120, Satt335, and Satt334, A245_1, Satt362, B174_3, A708_1, Sct_188, Pa2, Sat_375, Satt072, B1, Sat_313, Bng190_1, cr207_2, E049_2, and Satt490, and markers that map within about 10 to about 20 cM of any of the foregoing.

A further class of markers useful in the present methods include: GMRUBP_SSR, Satt325, Sat_390, M8E6mr1, Satt146, Satt586, Satt569, Satt343, DOP_A04, Sat_387, Satt193, G214_13, Satt030, K250_1, Satt649, j11_1, Sat_262, BLT030_1, Satt145, 0PAN06, Satt269, Satt346, Satt252, Satt149, BLT010_1, AW186493, Satt423, BE806387, Sat_240, Satt206, Satt659, Sat_039, W1, BLT057_1, Sat_298, DUBC767, C0L2-1, Satt160, A401_1, A806_1, K390_1, Sat_309, Satt374, Satt516, Satt425, Mng228_1, Gy5, G248_2, Bng075_1, K002_1, Satt595,

SSR locus

Satt252

Satt140

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B202_1, Sat_133, K265_1, K314_1, Bng004_1, HSP_2, Satt663, Sat_103, Bng118_1, Sat_297 Ubiquitin, BLT025_1, P157_1, Mng157_1, A757_1, Sat_229, Satt114, L063_1, A186_1, Rpg1, Sat_234, K644_1, L28831, Soyhsp176, Rpv1, B212_1, BLT053_7, Sat_154, 5 Rsv1, R045_1, L050_14, Satt510, Sat_317, K007_2, Rps3, cr321_2, Sct033, Sat_120, Satt335, and Satt334, A245_1, Satt362, B174_3, A708_1, Sct_188, Pa2, Sat_ 375, Satt072, B1, Sat_313, Bng190_1, cr207_2, E049_2, SATT490, Shr, L195_2, Satt144, cr409_1, Sat_197, 10 K014_2, B148_1, Sat554, A566_1, Ngm26b, Satt657, Cgy1, Ngm22, Satt218, Satt522, AW756935, T092_1, Sat_ 090, Bng172_1, Satt656, A083_3, Sat_417, K102_2, Sat_ 074, Ngm23b, Ngm44b, Satt395, and 0PAV06b, and markers that map within about 10 to about 20 cM of any of the 15 foregoing.

Updated information regarding markers assigned to soybean linkage group F may be found on the USDA's Soybase website. Table 1 provides current information on the Genbank location, location in Linkage Group F, and Accession Nos. of 20 markers useful in the methods disclosed herein. Sequence information pertaining to the markers can be found on Genbank using the gi#. Table 2 provides upper and lower primer sequences for these markers. Note that FIG. 2 indicates a different order for the markers shown. It should be understood 25 that up-to-date information regarding markers on Linkage Group F can be used in the methods disclosed herein as it becomes available.

TABLE 1

SSR locus	GenBank gi#	cM Position in LG	GenBank Accession#
GMRUBP	18741	0.00	V00458
Sat_390	31044745	1.79	CC453915
Satt146	14969861	1.92	BH126358
Satt325	14970019	2.23	BH126516
Satt343	14970037	3.04	BH126534
Sat_387	31044742	3.11	CC453912
Satt569	14970238	3.35	BH126735
Satt193	14969903	3.42	BH126400
Satt586	14970255	3.63	BH126752
Satt030	14969810	3.95	BH126307
Satt649	31044834	5.36	CC454004
Sat262	31044627	9.69	CC453797
Satt145	14969860	10.65	BH126357
Satt269	14969968	11.37	BH126465
Satt348	14970041	15.29	BH126538

12 TABLE 1-continued

cM Position

in LG

GenBank

Accession #

BH126450

BH126361

Markers on Linkage Group F

GenBank

gi#

14969953

14060864

Satt423 14970105 20.56 BH126602 AW186493 6455810 21.04 AW186493 BE806387 10237499 22.97 BE806387 Sat_240 31044608 25.58 CC453778 Satt659 31044844 26.71 CC454014 Satt205 14969915 26.98 BH126412 Sat_309 15243073 27.87 BH146207 Sat_309 31044671 41.47 CC453841 Satt374 14970064 43.01 BH126672 Satt316 14970107 43.44 BH126686 Satt516 14970189 44.42 BH126686 Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_297 31044650 59.60 CC453830 Sat_297 31044650 59.60 CC453768 Sat_114 14969885 63.69 BH126332 <th< th=""><th>Satt149</th><th>14969864</th><th>18.13</th><th>BH126361</th></th<>	Satt149	14969864	18.13	BH126361
BE806387 10237499 22.97 BE806387 Sat_240 31044608 25.58 CC453778 Satt659 31044844 26.71 CC454014 Satt205 14969915 26.98 BH12612 Sat_039 15243073 27.87 BH146207 Sat_298 31044661 32.32 CC453831 Satt160 14969875 33.19 BH126372 Sat_309 31044671 41.47 CC453841 Satt374 14970064 43.01 BH126561 Satt425 14970107 43.44 BH126604 Satt516 14970189 44.42 BH126606 Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_297 31044660 59.60 CC453803 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332	Satt423	14970105	20.56	BH126602
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Satt659 31044844 26.71 CC454014 Satt205 14969915 26.98 BH126412 Sat_039 15243073 27.87 BH146207 Sat_298 31044661 32.32 CC453831 Satt160 14969875 33.19 BH126372 Sat_309 31044671 41.47 CC453841 Satt374 14970064 43.01 BH126561 Satt425 14970189 44.42 BH126604 Satt516 14970189 44.42 BH126686 Satt595 14970264 50.24 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969806 50.78 BH126275 Sat_297 31044660 59.60 CC453830 Sat_297 31044600 59.60 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sa	BE806387	10237499	22.97	BE806387
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Sat_039 15243073 27.87 BH146207 Sat_298 31044661 32.32 CC453831 Satt160 14969875 33.19 BH126372 Sat_309 31044671 41.47 CC453841 Satt374 14970064 43.01 BH126561 Satt425 14970107 43.44 BH126604 Satt516 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126672 Sa	Satt659	31044844	26.71	CC454014
Sat_298 31044661 32.32 CC453831 Satt160 14969875 33.19 BH126372 Sat_309 31044671 41.47 CC453841 Satt374 14970107 43.44 BH126561 Satt516 14970189 44.42 BH126686 Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_317 31044573 68.91 CC453705 Satt510 14970184 71.41 BH126673 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sa	Satt205	14969915	26.98	BH126412
Satt160 14969875 33.19 BH126372 Sat_309 31044671 41.47 CC453841 Satt374 14970064 43.01 BH126561 Satt425 14970107 43.44 BH126604 Satt516 14970189 44.42 BH126686 Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_151 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453705 Satt331 14970276 74.13 BH126773 Sa	Sat_039	15243073	27.87	BH146207
Sat_309 31044671 41.47 CC453841 Satt374 14970064 43.01 BH126561 Satt425 14970107 43.44 BH126604 Satt516 14970189 44.42 BH126666 Satt595 14970264 50.24 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969878 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_293 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126526 Sa	Sat_298	31044661	32.32	CC453831
Satt374 14970064 43.01 BH126561 Satt425 14970107 43.44 BH126604 Satt516 14970189 44.42 BH126606 Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126720 Satt334 14970029 77.70 BH126526 Sa	Satt160	14969875	33.19	BH126372
Satt425 14970107 43.44 BH126604 Satt516 14970189 44.42 BH126686 Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126729 Satt334 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Sa	Sat_309	31044671	41.47	CC453841
Satt516 14970189 44.42 BH126686 Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 149697028 78.06 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970028 78.06 BH126525 S	Satt374	14970064	43.01	BH126561
Satt595 14970264 50.24 BH126761 Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044600 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Satt_120 14969793 75.97 BH126526 Satt334 14970028 78.06 BH126526 Satt362 14970028 78.06 BH126782 Satt072 14969823 87.01 BH126782 S	Satt425	14970107	43.44	BH126604
Sat_133 14969806 50.78 BH126303 Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt334 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970028 78.06 BH126525 Satt072 14969823 87.01 BH126320 Sa	Satt516	14970189	44.42	BH126686
Satt663 31044848 56.17 CC454018 Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt334 14970029 77.70 BH126526 Satt362 14970028 78.06 BH126525 Satt362 14970028 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453845 Sa	Satt595	14970264	50.24	BH126761
Sat_103 14969778 57.77 BH126275 Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126526 Satt334 14970028 78.06 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Sa	Sat_133	14969806	50.78	BH126303
Sat_297 31044660 59.60 CC453830 Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126590 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970028 78.06 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Sa	Satt663	31044848	56.17	CC454018
Sat_229 31044598 62.79 CC453768 Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970028 82.83 BH126782 Satt072 14969823 87.01 BH126782 Satt072 14969823 87.01 BH126320 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 S	Sat_103	14969778	57.77	BH126275
Satt114 14969835 63.69 BH126332 Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126525 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969889 102.08 BH126356 Sat_197 31044568 103.51 CC453738	Sat_297	31044660	59.60	CC453830
Sat_234 31044603 66.55 CC453773 SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738	Sat_229	31044598	62.79	CC453768
SOYHSP176 169984 68.44 M11317 Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126920 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt254 14970224 111.89 BH126721 <td< td=""><td>Satt114</td><td>14969835</td><td>63.69</td><td>BH126332</td></td<>	Satt114	14969835	63.69	BH126332
Sat_154 31044535 68.91 CC453705 Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 149700285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt657 31044568 103.51 CC453738 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126692	Sat_234	31044603	66.55	CC453773
Satt510 14970184 71.41 BH126681 Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126525 Satt362 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126721 Satt522 14970195 119.19 BH126692	SOYHSP176	169984	68.44	M11317
Sat_317 31044678 72.97 CC453848 Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969889 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692	Sat_154	31044535	68.91	CC453705
Sct_033 14970276 74.13 BH126773 Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt22 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935	Satt510	14970184	71.41	BH126681
Sat_120 14969793 75.97 BH126290 Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265	Sat_317	31044678	72.97	CC453848
Satt335 14970029 77.70 BH126526 Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt654 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC453041	Sct_033		74.13	BH126773
Satt334 14970028 78.06 BH126525 Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC453041 Sat_417 31044771 135.95 CC453941	Sat_120			
Satt362 14970053 82.83 BH126550 Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC453041 Sat_417 31044511 142.35 CC453681	Satt335		77.70	BH126526
Sct_188 14970285 85.33 BH126782 Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Satt334		78.06	
Satt072 14969823 87.01 BH126320 Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Satt362	14970053	82.83	BH126550
Sat_375 31044731 88.09 CC453901 Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Sct_188	14970285	85.33	BH126782
Sat_313 31044675 91.87 CC453845 Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Satt072	14969823	87.01	BH126320
Satt490 14970164 97.97 BH126661 Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Sat_375	31044731	88.09	CC453901
Satt144 14969859 102.08 BH126356 Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Sat_313	31044675	91.87	CC453845
Sat_197 31044568 103.51 CC453738 Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Satt490	14970164	97.97	BH126661
Satt554 14970224 111.89 BH126721 Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Satt144	14969859	102.08	BH126356
Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Sat 197	31044568	103.51	CC453738
Satt657 31044842 116.91 CC454012 Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681	Satt 554	14970224	111 89	BH126721
Satt218 14969925 117.65 BH126422 Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681				
Satt522 14970195 119.19 BH126692 AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681				
AW756935 7686224 124.88 AW756935 Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681				
Sat_090 14969768 130.64 BH126265 Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681				
Satt656 31044841 135.12 CC454011 Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681				
Sat_417 31044771 135.95 CC453941 Sat_074 31044511 142.35 CC453681				
Sat_074 31044511 142.35 CC453681				
-				
Satt395 14970081 146.42 BH126578	_			
	Satt395	14970081	146.42	BH126578

TABLE 2

	Marker Seque	ences
SSR locus	Upper primer sequence (5'>3'	Lower primer sequence (5>3')
GMRUBP	CTGGCGTGCTAAAAGTA [SEQ ID NO: 1]	GGACAGATTTGATCAATAATT [SEQ ID NO: 2]
Sat_390	GCGTAGATGCTTATAATCGACCCTAACAAT	FGCGCGAGGATCCCATAAAAAAAGTAAAATAG [SEQ ID NO: 4]
Satt146	AAGGGATCCCTCAACTGACTG [SEQ ID NO: 5]	GTGGTGGTGAAAACTATTAGAA [SEQ ID NO: 6]
Satt325	GCGGGGTATTAAGGGAAAACAAAA [SEQ ID NO: 7]	GCGTAAACGAACAATCACTTCATA [SEQ ID NO: 8]
Satt343	CATGGCGGAAAGCGAAACA [SEQ ID NO: 9]	TCCCAATTCACCTCTTCA [SEQ ID NO: 10]

TABLE 2-continued

Marker Sequences					
SSR locus	Upper primer sequence (5'>3')				
Sat_387	GCGGAATTTACCAGTTTATAATATTGCTGA [SEQ ID NO: 11]	GCGTACTAAATATTCAAAGACTCAAAGAGAA [SEQ ID NO: 12]			
Satt569	GCGCAAATTGCTTCACGCATCCAAAT [SEQ ID NO: 13]	GCGGCCTACTATAGTGAAGGGTATA [SEQ ID NO: 14]			
Satt193	GCGTTTCGATAAAAATGTTACACCTC [SEQ ID NO: 15]	TGTTCGCATTATTGATCAAAAAT [SEQ ID NO: 16]			
Satt586	GCGGCCTCCAAACTCCAAGTAT [SEQ ID NO: 17]	GCGCCCAAATGATTAATCACTCA [SEQ ID NO: 18]			
Satt030	AAAAAGTGAACCAAGCC [SEQ ID NO: 19]	TCTTAAATCTTATGTTGATGC [SEQ ID NO: 20]			
Satt649	TTACTGGCCGTGTTTACCCGTGTAA [SEQ ID NO: 21]	GCGGACGTTATAAGATTTTTTTATCATG [SEQ ID NO: 22]			
Sat_262	GCGTTTGCATTAGGGATTATCTAGTTTATGA [SEQ ID NO: 23]	GCGGGTTAGAACATTCTTAGTTAGCTCCAG [SEQ ID NO: 24]			
Satt145	AGCATATGGGATACAAGTGATTAG [SEQ ID NO: 25]	CGGTGTTGGTGTGTATGT [SEQ ID NO: 26]			
Satt269	GCGTGCCAGGTAGAAAATATTAG [SEQ ID NO: 27]	GCGGTTTTTCACTTTTCAAAATTC [SEQ ID NO: 28]			
Satt348	GCGCTTAGTAATGGTTCCCACAGATAA [SEQ ID NO: 29]	GCGGTGATATCTAGCAACACAA [SEQ ID NO: 30]			
Satt252	GCGAATTTGGATTAATTAAATTTATG [SEQ ID NO: 31]	GCGCTCGGTCCTCTCAAATAAGGTCTC [SEQ ID NO: 32]			
Satt149	TTGCACATTCTTTTTGGTAAACAGTCATAA [SEQ ID NO: 33]	GTTGGAGGCCATAGTCACATTAATCTTAGA [SEQ ID NO: 34]			
Satt423	TTCGCTTGGGTTCAGTTACTT [SEQ ID NO: 35]	GTTGGGGAATTAAAAAAATG [SEQ ID NO: 36]			
AW186493	GCGGTGATCCGTGAGATG [SEQ ID NO: 37]	GCGGAAAGTAGCACCAAGAG [SEQ ID NO: 38]			
BE806387	GCGACCCCTTTTGTCTTCTT [SEQ ID NO: 39]	GCGGAGGCCAGAGATGAA [SEQ ID NO: 40]			
Sat_240	GCGGGCAGAAGTCTAATGAATGTGAAATGA [SEQ ID NO: 41]	GCGGTTGTGACCGAAATAGATGTTATTTAAT [SEQ ID NO: 42]			
Satt659	GCGGCTCAACTTCGTGTAACAAG [SEQ ID NO: 43]	GCGCATCGGTAACTATCTAATATTCGTA [SEQ ID NO: 44]			
Satt206	GCGCATGTGAAAAGAATGAGATTATGTA [SEQ ID NO: 45]	GCGTCCAAACTCATCCTTAAGGTATT [SEQ ID NO: 46]			
Sat_039	CAAGAATAATCTAAAGGTACACTT [SEQ ID NO: 47]	AGTTAAAAAACCCACACAC [SEQ ID NO: 48]			
Sat_298	GCGCGTCGAAGCAAAAATTAAA [SEQ ID NO: 49]	GCGGCGAAACCCACAAAGCATA [SEQ ID NO: 50]			
Satt160	TCCCACACAGTTTTCATATAATATA [SEQ ID NO: 51]	CATCAAAAGTTTATAACGTGTAGAT [SEQ ID NO: 52]			
Sat_309	GCGAACGGATATATACCCATAAATTTTCATG [SEQ ID NO: 53]	GCGTCATCCAATATAACAATTGTTAAAGTCA [SEQ ID NO: 54]			
Satt374	AACATTTGCCGAAAAAAATAACTATGATG [SEQ ID NO: 55]	GCGTATCAATTAAGATCCATTAAGTG [SEQ ID NO: 56]			
Satt425	GCGCAATTAAGATCCCTAAGTGATT [SEQ ID NO: 57]	GCGGCTTTTCACTCTTCTTTTATTATT [SEQ ID NO: 58]			
Satt516	GCGTTAGCACTATTTTTTTACAAGA [SEQ ID NO: 59]	GCGCCGTTCCTCTTTACTTTAT [SEQ ID NO: 60]			

TABLE 2-continued

	Marker Sequences				
SSR locus	Upper primer sequence (5'>3')				
Satt595	GATGGGAAGCAAACAAGAAG [SEQ ID NO: 61]	AACCCCCTCCCCTAAAT [SEQ ID NO: 62]			
Sat_133	GCGCACATCTTAACTCAAATAATTGATAAAG [SEQ ID NO: 63]	GCGTTCAATTGGATTTGATGAAATTTTAAAT [SEQ ID NO: 64]			
Satt663	GCGTCATGCAATGTTGTATAAT [SEQ ID NO: 65]	GCGACTGCAGATAACTTGACTGGTAGT [SEQ ID NO: 66]			
Sat_103	ACTGGGAATCCATTTCTTGTTA [SEQ ID NO: 67]	AAAGAACTTTCAATCAAATGTTGTG [SEQ ID NO: 68]			
Sat_297	GCGTGAAAATAAATACATAGACATCCACCAT [SEQ ID NO: 69]	GCGTTTTAACACGCATCAACACTCTTC [SEQ ID NO: 70]			
Sat_229	GCGTGTGCTACTTCACATCTTGAGAGAAAGA [SEQ ID NO: 71]	GCGAGGGTTTAGAAAAAGATTCACCAAATAT [SEQ ID NO: 72]			
Satt114	GGGTTATCCTCCCCAATA [SEQ ID NO: 73]	ATATGGGATGATAAGGTGAAA [SEQ ID NO: 74]			
Sat_234		GCGGAAACCATCCTTATATGTCAATTGCTCA [SEQ ID NO: 76]			
SOYHSP176	TTTTTGTTTAAGTTACTGTACTGT [SEQ ID NO: 77]	GCTAGTCTTCTACAACCTTCTA [SEQ ID NO: 78]			
Sat_154	GCGTCAGGGTCAAGTCATCTAACA [SEQ ID NO: 79]	GCGGACGCATTTCCTATTGATCAAG [SEQ ID NO: 80]			
Satt510	GCGAGTTTCGCCGTTACCACCTCAGCTT [SEQ ID NO: 81]	CCCTCTTATTTCACCCTAAGACCTACAA [SEQ ID NO: 82]			
Sat_317	GCGACAGTCCCAATACCATTAACAAGT [SEQ ID NO: 83]	GCGTCCTTAGGTACCTAGAATAATTCTTCAC [SEQ ID NO: 84]			
Sct_033	CTTTTAAATTATAATAGCATGATCT [SEQ ID NO: 85]	TGCTAATTTAGATTACGTTATGT [SEQ ID NO: 86]			
Sat_120	CATATAAAAATGGTCCTCTCACATA [SEQ ID NO: 87]	GCTTGAGCAACTTACAATTCACT [SEQ ID NO: 88]			
Satt335	CAAGCTCAAGCCTCACACAT [SEQ ID NO: 89]	TGACCAGAGTCCAAAGTTCATC [SEQ ID NO: 90]			
Satt334	GCGTTAAGAATGCATTTATGTTTAGTC [SEQ ID NO: 91]	GCGAGTTTTTGGTTGGATTGAGTTG [SEQ ID NO: 92]			
Satt362	GCGTTGTTGTTTCAAATGTATTTTAGTT [SEQ ID NO: 93]	GCGGACGGATCATCAAACCAATCAAGAC [SEQ ID NO: 94]			
Sct_188	TTCAACCATGTCATAAAAT [SEQ ID NO: 95]	CTCACTCCTCCATAAAAAT [SEQ ID NO: 96]			
Satt072	GGAAAGAATCAGCAAAAT [SEQ ID NO: 97]	CCCCCACATAAATAATAAA [SEQ ID NO: 98]			
Sat_375	GCGTGTTAATGATTGCATAAGGTTCG [SEQ ID NO: 99]	GCGTGTCAAAAGAAACTCAATAAAGAAAAAT [SEQ ID NO: 100]			
Sat_313	GCGTATTCCCTTAACAAAATTAAAGTTTCAC [SEQ ID NO: 101]	GCGCGTCAGCCTAACAAAAAGAATAAAAT [SEQ ID NO: 102]			
Satt490	GCGGCACGAGTCAACTTTCTGTTTCCT [SEQ ID NO: 103]	GCGGAAGAAGATTTTCGTTTTTAT [SEQ ID NO: 104]			
Satt144	CGTCGCCATCACTATGAGAA [SEQ ID NO: 105]	CCATCTTGAGCAGAGTTTGAAGTT [SEQ ID NO: 106]			
Sat_197		GCGGTTAACAGCCAAGTTCTTTC [SEQ ID NO: 108]			
Satt554	GCGATATGCTTTGTAAGAAAATTA [SEQ ID NO: 109]	GCGCAAGCCCAAATATTACAAATT [SEQ ID NO: 110]			

TABLE 2-continued

Marker Sequences				
SSR locus	Upper primer sequence (5'>3')	Lower primer sequence (5>3')		
Satt657	GCGCATTTGGACTTTTACTTC [SEQ ID NO: 111]	GCGACGATGTTAATTGGTAGAATC [SEQ ID NO: 112]		
Satt218	TCAATCAACAAAAACATAATTCTTC [SEQ ID NO: 113]	ATTTGTGTTTTGTTTTAGCTCTCTA [SEQ ID NO: 114]		
Satt522	GCGAAACTGCCTAGGTTAAAA [SEQ ID NO: 115]	TTAGGCGAAATCAACAAT [SEQ ID NO: 116]		
AW756935	GCGGCTGGTGATTGTGTAAT [SEQ ID NO: 117]	GCGTAATATAGTTTTGTATTGAAAT [SEQ ID NO: 118]		
Sat_090	CTCGCTGCTACTGGTC [SEQ ID NO: 119]	AAGAATGCGTTGGATTTA [SEQ ID NO: 120]		
Satt656	GCGTACTAAAAATGGCAATTATTTGTTG [SEQ ID NO: 121]	GCGTGTTTCAGTATTTGGATAATAGAAT [SEQ ID NO: 122]		
Sat_417	GCGAATATGGCGTTGAAAATAGTGAT [SEQ ID NO: 123]	GCGACCCAGATTCTGTGCTAAGA [SEQ ID NO: 124]		
Sat_074	GGGTGAGAAATACATGCAACTTACA [SEQ ID NO: 125]	GGGCATCAAAATTGATATTAAATGTCTAA [SEQ ID NO: 126]		
Satt395	CGCGCTAGTTGAATGAATGT [SEQ ID NO: 127]	GCGCATTGAGGAATTTTTAT [SEQ ID NO: 128]		

Other types of markers such as SNP markers, for example, as described in Jeong, S. C. and Saghai Maroof, M. A. (2004), "Detection and genotyping of SNPs tightly linked to two disease resistance loci, Rsv1 and Rsv3, of soybean," Plant Breeding 123:305-310, mapping close to Rag2 on linkage group F are also useful in the methods described herein.

Sequences for specific markers useful in the present methods are provided below (taken from the USDA Cregan Soymap website):

Sat_297:

```
[SEQ ID NO: 129]
      gatecectea geetageett cagatgtgge etgaceagag ageattgaat gaacageaeg
  61 ttccttttct tgctccagca ccgtcataca gtggagggat ggttccacct caaggaatgt
      atccatcttc tgattggagt gggtatcatc aggtaccttt gaatccatat taccctcccg
 181 gtgttccttt cccgcatttt ccagctgccc atatgaatca cccgatgtac aaggctgcag
      atataccagg acatcaacca ccaccatctg atgagtatcc cgagagacct ggccaacctg
 301 aatgccagca tttcgtta
Sat_234:
                                                                    [SEQ ID NO: 130]
   1 taacgcgaaa gggggaacat cttatatgaa taataataaa tggagaaaag gaaaagaatc
  61 acaggttcca ggttttttcc ttttataccc tccttttctt cctaaattct gaggtttcac
 121 cataaccata ttqqqatc
Soyhsp176:
                                                                   [SEQ ID NO: 131]
   1 gaattetgaa attgggtett tttgtgggea etttttgatg tttttgttta agttactgta
      ctgtgggcca caaaacgtat agatcaaagt agtaataata atattgatta aatgatatat
      atatatatat atatatat atatctagaa ggttgtagaa gactagctag aacgtacgta
      ttcgtgtgga gaagtcctga agtttatcga atcatctaaa actgctaaaa tagcaaacaa
 181
 241 cattatattg taaacaatat ttttctggaa catacaagag tatcctttca cttcctttaa
      atacctcgag tgtccccatt gacatcatca aacaagagaa gagttacaga atttcctgtt
 301
 361 tacgatetea ttacaatttt geaactttea aagettatta getaaagtaa cateaaaaga
 421
      tgtcattgat tccaagtatt ttcggtggcc caaggagcaa cgtgttcgat ccattctcac
      tcgatatgtg ggatcccttc aaggattttc atgttcccac ttcttctgtt tctgctgaaa
 541 attctgcatt tgtgaacaca cgtgtggatt ggaaggagac ccaagaggca cacgtgctca
 601
      aggctgatat tccagggctg aagaaagagg aagtgaaggt tcagattgaa gatgataggg
 661 ttcttcagat tagcggagag aggaacgttg agaaggaaga caagaacgac acgtggcatc
 721
      gcgtggaccg tagcagtgga aagttcatga gaaggttcag attgccagag aatgcaaaag
      tggagcaagt aaaggcttgt atggaaaatg gggttctcac tgttactatt ccaaaggaag
 841 aggttaagaa gtctgatgtt aagcctatag aaatctctgg ttaaacttgg tttcactgaa
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901 aatcgtgaga gcttttaaat ttgctttgtt gtaataagtg tcctttgtct tgtgttccaa
  961 tggtgatttt gagaaagatc atacaattgt gccttgtgtt gttgtgcaag tgtaattgaa
 1021 gtgaataaaa aattaacacc tgctttcaga aaattttgct gtgtgtcatt gtcatcgaat
 1081
      atgtgatgta ggcaagaaat agaccgtgaa aataatatct gacatttggc taattgcttt
 1141 tgttatgctg agacactcta tgtgaaataa ctgcatttat catgttccat cttcttaata
 1201 caagaagtca ataccaatgt cttaccaaat taagataaca ggttgatttg gactcatcaa
 1261 agtgcagccc tttatttgga ctcatcaaag tgcagcacta aagggttttg ttaactagca
 1321 agttcagagc atcatttaag taattaaaag aaaaaatatt aaatatataa atcataagat
 1381 gatatcaaaa aattcatgaa cagtctcttc atttttttc aataaaaata tttttatttt
      aattttttaa aataatatcc tcataacatt ggtttaactc ccaagtttaa aatttactag
 1441
 1501 tgctagataa attctctaag ataatgtata gataaaata agataaatta gaaaattttt
 1561
     aaggagagat ttttttttat aaaaattagg tatatgtatt ggttttagtt tacagagaaa
 1681 taaatottaa tattttttt gacagaatto t
Satt510 (BAC-cultivar Faribault):
                                                                [SEQ ID NO: 132]
    1 geogtegeet tageeggage tgeaggetee gtgeettget eegeegeeat
   61 categories tegetector toteleat gaetgeoritt gaegitttaa gattetatat
  121 ataqtttqca tttcatqaat tattattcca aaataatata taqaqaqata ataaactqtt
  181 agattqcqaq tttcqccqtt accacctcaq cttattttat qattattatt attattatta ttattattat
  241 tattattatt attattatta ttattattat tattaattqt tqtaqqtctt aqqqtqaaat aaqaqqqatt
  301 ttgatcctct ctacattttt attgttaatt atgtaatgct atatattatg tatgggtata atttagatcc
  361 agtcatttta tgtttctcat gttctttttt ttaatttatc acttctataa agaaaataac ttaaactcaa
  421 aatacttata ataacatago tgatacattt atattatato cactaaatta tttgatatat gagcagtato
  481 gtagtggtat aggtttgaat gtcagagg
Sat_120:
                                                                [SEQ ID NO: 133]
   1 taaaqctqca ccaqctaqca tttccttqat atcaataccc tqcaatqcaq ctqqaaqtaa
   61 acccacaget gacatttetg aagteetace accaacecaa teaaacatag gaaacegage
  121 taaccatccc tctattctag cagcggtatc caacagagaa ttttcttgag taattgcaac
  181 accctgtttt gagaattgca gccctgcatc tctgaaggct ttccgtactt ctagtagacc
  241 attgcgggtt tcaggtgtgc ctccgctctt agaaatgaca attacaagag tagttgccag
  301 ttcaggtcct agttgagcaa tttgatgatc aatcccagca ggatc
Sat_375:
                                                                [SEQ ID NO: 134]
    1 aatcattaac atataccatt agaatatgtt aatgattgca taaggttcgg gcacccacta
   121 ctattagaat atgttattct cagtcttagt ttattttaga cttttagatt ttgagtagtt
  181 acatattaac attctaaata gtgcaaatac tatattgaaa attcattatt tttctttatt
  241 gagtttcttt tgacatatta taattacatt acttagatag actacttata tttctttctg
  301 tatatatgat aaggtgtatt actaacccca ctagagctac aactacaact aaagaaataa
  361 tataaaacta tgaatatcaa tcttctgtgt tttcatttaa ttatattcgg ttataaaaca
  421 ataacagctc ataaaacaat aattattgaa atttaaaaatc c
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-continued

Markers that map closer to the Rag2 locus are preferred over markers that map farther from the Rag2 locus for use in the present methods. The markers may be any type of mapped molecular marker or phenotypic trait known to the art, including restriction fragment length polymorphism (RFLP) markers, target region amplification polymorphism (TRAP) markers, random amplified polymorphic (RAPD) markers, simple sequence repeat (SSR) markers, single nucleotide polymorphism (SNP) markers, and isozyme markers.

In one embodiment of the methods described herein, markers flanking the Rag2 locus are used in the marker-assisted selection processes as described herein. The genomic DNA of soybean germplasm is preferably tested for the presence of at least two of the foregoing molecular markers, one on each side of the Rag2 locus. Most preferably, the two markers are 55 Soyhsp176 and Satt510. Markers that map close to Soyhsp176 and Satt510 can also be used, provided they fall to either side of the Rag2 locus. Preferably, one of said at least two molecular markers is within at least about 3 to about 10 cM, or about 10 to about 20 cM of Satt510, and another of said 60 at least two molecular markers is within at least about 3 to about 10 cM or about 10 to about 20 cM of Soyhsp176, and to ensure that the markers used flank the Rag2 locus, one of said at least two molecular markers within at least about to 3 about 10 cM or about 10 to about 20 cM of Satt510 should be farther 65 than that distance from Soyhsp176, and another of said at least two molecular markers within at least about 3 to about

10 cM or about 10 to about 20 cM of Satt510 should be farther than that distance from Soyhsp176.

A method described herein for reliably and predictably introgressing soybean *Aphis glycines* resistance into non-resistant soybean germplasm or into less or differently-resistant soybean germplasm comprises: providing a first soybean germplasm that has Rag2-gene-derived resistance to *Aphis glycines*; providing a second soybean germplasm that lacks Rag2-gene-derived resistance to *Aphis glycines*; crossing the first soybean germplasm with the second soybean germplasm to provide progeny soybean germplasm; screening said progeny germplasm to determine the presence of Rag2-gene-derived resistance to *Aphis glycines*; and selecting progeny that tests positive for the presence of Rag2-gene-derived resistance to *Aphis glycines* as being soybean germplasm into which germplasm having Rag2-gene-derived resistance to *Aphis glycines* has been introgressed.

The second soybean germplasm that lacks Rag2-gene-derived resistance to *Aphis glycines* can be germplasm that lacks resistance to *Aphis glycines* entirely, or can be germplasm that has *Aphis glycines* resistance derived from another source, such as a Rag1 or related gene as described in U.S. Patent Publication No. 2006/0014964.

Preferably, the screening and selection are performed by using marker-assisted selection using a marker on major linkage group F as described above.

The screening and selection can also be performed by exposing plants containing said progeny germplasm to aphids

of the species Aphis glycines in a live aphid bioassay and selecting those plants showing resistance to aphids (or if the second germplasm already carries Aphis glycines resistance from a different gene, selecting those plants showing resistance to an Aphis glycines biotype that can overcome resistance that is present in the second germplasm) as containing soybean germplasm into which germplasm having Rag2gene-derived resistance to Aphis glycines has been introgressed. The live aphid assay may be any such assay known to 10 the art, e.g., as described in Hill, C. B., et al., "Resistance to the soybean aphid in soybean germplasm" (2004) Crop Science 44:98-106, Hill, C. B., et al., "Resistance of Glycine species and various cultivated legumes to the soybean aphid 15 (Homoptera: Aphididae)" (2004) J. Economic Entomology 97(3)1071-1077, "Li, Y. et al., "Effect of three resistant soybean genotypes on the fecundity, mortality, and maturation of soybean aphid (Homoptera: Aphididae)" (2004) J. Economic Entomology 97(3):1106-1111, Hill, C. B., et al., "A single 20 dominant gene for resistance to the soybean aphid in the soybean cultivar Dowling" (2006) Crop Science 46: 1601-1605, or Hill, C. B., et al. "Soybean aphid resistance in soybean Jackson is controlled by a single dominant gene" (2006) 25 Crop Science 46: 1606-1608, or as described in the Examples hereof. A preferred method includes placing aphid-infested plant parts on vegetative cotyledon (VC) stage plants and rating aphid population and plant damage weekly. As described herein, a 0 to 4 scale, where 0=no aphids present, 1=few solitary live or dead aphids (dead aphid bodies) present, 2=several transient aphids (aphids possibly probing for a suitable feeding site) present with some viviparous aptera surrounded by a few nymphs, 3=dense colonies, and 35 4=dense colonies accompanied by plant damage, including leaf distortion and stunting, may be used.

The screening and selection may also be done by methods including hybridizing nucleic acid from plants containing progeny germplasm to a nucleic acid fragment comprising a 40 Rag2 gene, and selecting those plants having germplasm that hybridizes to the nucleic acid fragment as having resistance to *Aphis glycines*.

A method described herein for breeding a soybean plant homozygous for the Rag2 Aphis glycines resistance gene that 45 is a cultivar adapted for conferring, in hybrid combination with a suitable second inbred, Rag2 resistance to Aphis glycines, comprises selecting a first donor parental line possessing the desired Rag2 Aphis glycines resistance, said first donor parental line comprising a Rag2 Aphis glycines resis- 50 tance gene located on major linkage group F; crossing the first donor parental line with a second parental line that is generally high yielding in hybrid combination to produce a segregating plant population of genetically heterogenous plants; screening the plants of the segregating plant population for 55 the Rag2 gene; selecting plants from the population having the gene; and breeding by self-crossing the plants containing the gene until a line is obtained that is homozygous for the locus containing the Rag2 gene and adapted for conferring, in hybrid combination with a suitable second inbred, Rag2 resis- 60 tance to Aphis glycines.

The screening and selection are preferably performed by using marker-assisted selection as described above, but may also be performed by live aphid bioassay as described above, selecting those plants showing resistance to aphids as containing soybean germplasm having a Rag gene. When it is known that the only source of aphid resistance in the plant

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material comes from a plant having Rag2 resistance, it can be concluded that the resistance shown in live aphid bioassays is Rag2 resistance. The screening and selection may also be done by hybridizing nucleic acid from plants containing said progeny germplasm to a nucleic acid fragment comprising the Rag2 gene and selecting those plants whose germplasm hybridizes to the nucleic acid fragment as having an aphid resistance gene.

As the parental line having Rag2 soybean aphid resistance, any soybean line known to the art or disclosed herein as having Rag2 soybean aphid resistance, as described above, may be used. In addition, without undue experimentation, varieties set forth in Table 10 known to have soybean aphid resistance can be tested using marker-assisted analysis as described herein for the presence of the Rag2 gene, thus identifying additional lines for use in the breeding methods described herein.

Also provided herein are soybean plants produced by any of the foregoing methods:

Isolated nucleic acid fragments comprising a Rag2 gene are also provided herein. The nucleic acid fragments comprise at least a portion of nucleic acid belonging to linkage group F, and further comprise nucleotide sequences falling between molecular markers Satt510 and Soyhsp176. They are capable of hybridizing under stringent conditions to nucleic acid of a soybean cultivar having Rag2 resistance to *Aphis glycines*.

Vectors comprising such nucleic acid fragments, expression products of such vectors expressed in a host compatible therewith, antibodies to the expression product (both polyclonal and monoclonal), and antisense nucleic acid to the nucleic acid fragment are also provided herein.

Also provided herein are soybean plants having Rag2 resistance to *Aphis glycines* comprising a Rag2 gene and produced by introgression of DNA containing the gene into a soybean germplasm lacking the gene in its genome, and progeny of said soybean plants.

Seed of a soybean germplasm produced by crossing a soybean cultivar having Rag2 *Aphis glycines* resistance in its genome with a soybean cultivar lacking the Rag2 gene in its genome, and progeny thereof, is also provided herein. Such seed, from BC3 or BC4 generations derived from crosses with aphid resistant Sugao Zairai (PI200538)×Ina or ×Williams 82 F_2 plants, is made available through the University of Illinois.

EXAMPLES

Example 1

Genetic Analysis of Rag2 in PI200538

Crosses were made between PI200538 and two susceptible soybean cultivars, Ina and Williams 82. The parents, F₁ and F₂ plants were tested in a choice test in the greenhouse using the methods described in Hill, C. B., Y. Li, and G. Hartman (2006), "A single dominant gene for resistance to the soybean aphid in the soybean cultivar Dowling," Crop Science 46:1601-1605. Three weeks after infestation, aphid colonization was visually rated using the following scale: 0=no aphids present, 1=few solitary live or dead aphids (dead aphid bodies present), 2=several transient aphids present with some viviparous aptera surrounded by a few nymphs, 3=dense colonies, and 4=dense colonies accompanied by plant damage.

PI200538 plants had ratings of 0, I, or 2 with a rating of 1 being most frequent. Ina and Williams 82 plants had ratings of 3 or 4. Progeny from crosses between PI200538 and the

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susceptible parents were considered to be resistant with ratings of 0 to 2 and susceptible with ratings of 3 or 4. F₁ plants were all resistant to the soybean aphid, indicating that resistance was dominant over susceptibility. X² analyses on the segregation of F₂ plants (Table 3) indicated that a single 5 dominant gene conditioned resistance.

TABLE 3

Genetic analysis of the segregation of F_2 plants in two Ina × PI200538 and three Williams $82 \times PI200538$ F_2 populations for resistance to the soybean aphid Observed F_2 segregation						10	
Cross	F ₂ Family	Resistant	Susceptible	X ² (3:1)	P	15	
Ina × PI200538	4401	39	14	0.06	0.81		
	4741	75	23	0.12	0.73		
	Totals			0.18	0.91		
	Pooled	114	37	0.02	0.89		

0.16 0.69 Heterogeneity Williams 82 × 4791 88 30 0.01 0.92 PI200538 4792 67 0.39 0.53 19 4793 48 0.00 1.00 16 Totals 0.40 0.94 Pooled 155 49 0.75 0.10 Heterogeneity 0.29 0.86

 $F_{2:3}$ progeny from F_2 plants derived from plants in two Ina×PI200538 (Table 5) and three Williams 82×PI200538 (Table 6) F₂ populations were evaluated for resistance to the 30 soybean aphid. To have high confidence that all possible susceptible segregants were detected, only F_{2:3} families that had a minimum of 11 viable plants were included in the genetic analysis of F₂ plant soybean aphid resistance genotypes. A maximum of 20 F₃ plants from an F₂ plant were 35 included in the genetic analyses.

TABLE 4

Genetic analysis of the segregation of F _{2:3} families,	ies,
derived from plants in two Ina × PI200538 F ₂ populations,	lations,
for F ₂ plant soybean aphid resistance genotype	e

F ₂ family	F ₂ plant phenotype	F_2 plant genotype	No. of F _{2:3} families	X ² (1:2:1)	Р
4401	Resistant	RR (all F ₂₋₃ plants resistant)	4		
		Rr (resistant and susceptible	13		
		F _{2:3} plants)			
		rr (all F _{2:3} plants susceptible)	0		
	Susceptible	RR (all F _{2:3} plants resistant)	0		
		Rr (resistant and susceptible	0		
		F _{2:3} plants) rr (all F _{2:3} plants susceptible)	5		
		11 (an 1 2:3 plants susceptione)	,	0.82	0.66
4741	Resistant	RR (all F ₂₋₃ plants resistant)	14	0.62	0.00
		Rr (resistant and susceptible	20		
		F _{2:3} plants)			
		rr (all F _{2:3} plants susceptible)	0		
	Susceptible	RR (all F ₂₋₃ plants resistant)	0		
		Rr (resistant and susceptible	0		
		F ₂₋₃ plants)			
		rr (all F _{2:3} plants susceptible)	5		
				4.18	0.12
Totals				5.00	0.08
Pooled				1.62	0.44
Heterogeneity				3.37	0.07

TABLE 5

Genetic analysis of the segregation of $F_{2:3}$ families, derived from plants in three Williams $82 \times PI200538 \ F_2$ populations, for F2 plant soybean aphid resistance genotype

F ₂ family	F ₂ plant phenotype	F ₂ plant genotype	No. of F _{2:3} families	X ² (1:2:1)	P
4791	Resistant	RR (all F _{2:3} plants resistant) Rr (resistant and	17 43		
		susceptible F _{2:3} plants) rr (all F _{2:3} plants susceptible)	0		
	Susceptible	RR (all F _{2:3} plants	1		
		resistant) Rr (resistant and	0		
		susceptible $F_{2:3}$ plants) rr (all $F_{2:3}$ plants plants susceptible)	13	2.62	0.27
4792	Resistant	RR (all F _{2:3} plants	10	2.02	0.27
		resistant) Rr (resistant and susceptible	33		
		F _{2:3} plants) rr (all F _{2:3} plants susceptible)	1		
	Susceptible	RR (all F _{2:3} plants resistant)	0		

TABLE 5-continued

Genetic analysis of the segregation of $F_{2:3}$ families, derived from plants in three Williams $82 \times PI200538 F_2$ populations, for F_2 plant soybean aphid resistance genotype

F ₂ family	F ₂ plant phenotype	F ₂ plant genotype	No. of F _{2:3} families	X ² (1:2:1)	P
		Rr (resistant and susceptible F _{2:3} plants) rr (all F _{2:3} plants susceptible)	0	1.07	0.25
4793	Resistant	RR (all F _{2:3} plants	14	1.97	0.37
		resistant) Rr (resistant and susceptible	25		
		F _{2:3} plants) rr (all F _{2:3} plants susceptible)	0		
	Susceptible	RR (all F _{2:3} plants resistant)	0		
		Rr (resistant and susceptible	3		
		F _{2:3} plants) rr (all F _{2:3} plants	7		
		susceptible)		3	0.22
Totals Pooled Heterogeneity				7.59 4.34 3.25	0.06 0.11 0.07

Results of the $\rm F_2$ genetic analyses indicated that there was a single, dominant gene in PI200538 that conditioned resistance to the soybean aphid. The results of the $\rm F_3$ genetic analyses for the InaxPI200538 and WilliamsxPI200538 crosses supported the single, dominant gene hypothesis.

Crosses were made between the cultivars Dowling, possessing Rag1, and Jackson, that likely also possess Rag1, and PI200538, and their $\rm F_2$ progeny were evaluated for soybean aphid resistance to determine if Rag1 and the gene in PI200538 were allelic or the same gene. Segregation of resistant and susceptible $\rm F_2$ plants significantly fit a 15:1 pattern, expected for the segregation of two different, non-allelic dominant genes (Table 6). The results indicated that the gene in PI200538, tentatively called Rag2, is non-allelic and unique from Rag1.

TABLE 6

Genetic analysis of the segregation of F₂ progeny for soybean aphid resistance from crosses between Dowling and PI200538 and Jackson × PI200538

Observed F- segregation

Cross Susceptible $X^2(15:1)$ P Resistant 39 Dowling x 3 0.81 PI200538 Jackson × 71 6 0.310.58 PI200538 Totals 0.37 0.83 110 9 0.19 0.66 Pooled Heterogeneity 0.67

26

Example 2

Identification of Linked SSR Markers and Soybean Map Location of Rag2

Subsets of 90 F₂ plants were randomly selected from the combined F2 populations of each of the crosses Inax PI200538 and Williams 82×PI200538 for mapping the location of Rag2 in the soybean genetic map. DNA was extracted from each of the plants in each of the two subsets and polymerase chain reaction (PCR) was carried out using simple sequence repeat (SSR) markers developed by Dr. Perry Cregan, USDA-ARS. The PCR products were evaluated on gels as previously described in: Wang, D. J., et al., (2003), "A low-cost, high-throughput polyacrylamide gel electrophoresis system for genotyping with micro satellite DNA markers," Crop Science 43:1828-1832.

Initial SSR marker screening to identify markers that were polymorphic between the parents of the crosses and that 20 could be associated with the soybean aphid resistance gene was done with genomic DNA extracted from the parents and separate DNA samples from 10 randomly selected susceptible F₂ plants that were bulked from each cross subset. In order to minimize the number of soybean SSR markers to screen for polymorphisms and association with resistance, knowledge of the potential association between resistance to aphids and root knot nematodes, as put forward in Hill, C.B., Y. Li, and G. L. Hartman, (2004), "Resistance to the soybean aphid in soybean germplasm," Crop Science 44:98-106 was 30 exploited to select markers from soybean linkage groups (LG) E and F. Genes for resistance to peanut root knot nematode, found in PI200538, were mapped to LGs E and F (Tamulonis, J. P., et al. (1997), "DNA marker analysis of loci conferring resistance to peanut root-knot nematode in soybean," Theoretical and Applied Genetics 95:664-670). Two LG M SSR markers, Satt435 and Satt463, tightly linked to Rag1 (U.S. Patent Publication No. 20060015964), were also included in the screen as a check for genetic allelism between Rag1 and Rag2.

Six LG F SSR markers were found to be polymorphic between Ina and PI200538, associated with soybean aphid resistance, and linked to Rag2. The location of Rag2 in relation to the six SSR markers was generated with Joinmap 3.0, a genetic mapping software application, after entering the genotype data for the six LG F SSR markers, the F₂ resistance phenotype data, and available F₂ genotype data for each of the 90 F₂ plants in the Ina×PI200538 F2 mapping population. Tight linkage of Soyhsp176 and Satt510, both within 5 centimorgans (cM) on opposite sides of Rag2, was shown. With the location of Sat_297 taken as zero, and Rag2 at 18 cM, the additional markers were: Sat_234 at 5 cM, Soyhsp176 at 23 cM; Satt510 at 13 cM; Sat_120 at 26 cM and Sat_375 at 40 cM.

Subsequently, genotype data from 45 F2 plants from a cross between Ina×PI200538 was analyzed. Segregation of markers Sat_120 and Sat_375 did not fit the expected F2 1:2:1 ratio for a co-dominant gene, and therefore were dropped from the linkage data described above. The linkage map resulting from this analysis is shown in FIG. 2.

Example 3

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Effectiveness of Rag2 Against Different Soybean Aphid Isolates

A soybean aphid population was found in Ohio that could colonize soybean plants possessing the resistance gene Rag1.

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soybean aphid population was a biotype that could overcome

the resistance expressed by Rag1 in soybean plants, distin-

guishing it from other soybean aphid isolates. Identification of a soybean aphid biotype that can overcome the resistance ⁵ gene Rag1. In non-choice tests (Table 7) and choice tests (Table 8), resistance expressed by Rag2 in PI200538 was as

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TABLE 9-continued

Number of aphids of 11 soybean aphid isolates on Dowling (Rag1) and PI200538 (Rag2) 7 and 10 days after infestation

Soybean line	Aphid isolate	Number of aphids (7 days)	Number of aphids (15 days)
	Secor, Indiana from R. cathartica	0	0
	Springfield from R. frangula	17	4
	Stratton, Indiana from R. cathartica	8	42
PI200538	Black Hawk from R. cathartica	23	82
(Rag2)	Black Hawk III from R. cathartica	1	0
	Bronson, Michigan from R. cathartica	0	0
	Hy 47 & 64, Illinois from <i>R. cathartica</i> Irish Hills from <i>R. cathartica</i>	3	3
	Jolliette College from R. cathartica	0	0
	Pit, Indiana from R. cathartica	0	0
	Rock II from R. cathartica	0	0
	Secor, Indiana from R. cathartica	2	0
	Springfield from R. frangula	1	0
	Stratton, Indiana from R. cathartica	2	0

effective against the Ohio biotype as an isolate from Illinois.

Number of aphids per plant 10 and 15 d after infestation with the Ohio and Illinois aphid isolates across two non-choice tests.

TABLE 7

		10 days infestation	15 days after infestation	
Soybean genotype	Illinois Isolate	Ohio Isolate	Illinois Isolate	Ohio Isolate
Dowling (Rag1)	8cd [†]	146a	12de	586ab
Williams82	231a	209a	726a	574abc
LD05-16611 (Rag1)	10cd	215a	774a	548ab
Jackson	7d	191a	11de	396ab
Dwight	146a	178a	363ab	332b
PI567541B	6abcd	57b	5abcde	82c
PI567597C	31b	12c	18d	7e
PI200538 (Rag2)	8cd	8cd	9de	7e

 † Means followed by the same letters in the 10 d after infestation columns or the 15 d after infestation columns are not significantly different by the least significant different test (P = 0.05).

TABLE 8

with the Ohio isolate in a choice test.	
Number of aphids plant ⁻¹ 10 days after infestation	
177a	
168a	
166a	
156a	
110b	
34c	
31cd	
22d	
	te in a choice test. Number of aphids plant ⁻¹ 10 days after infestation 177a 168a 166a 156a 110b 34c 31cd

 $^{\uparrow}Means$ followed by the same letters in a column are not significantly different by the least significant different test (P = 0.05).

Results of a preliminary, un-replicated, non-choice test of 11 soybean aphid isolates collected from their primary hosts, *Rhamnus cathartica* and *R. frangula*, at different locations in the central USA, indicated that Rag2 provided effective resistance against all of them in PI200538 (Table 9).

TABLE 9

Numbe	r of aphids of 11 soybean aphid isolates on PI200538 (Rag2) 7 and 10 days after in		Rag1) and	55
Soybean line	Aphid isolate	Number of aphids (7 days)	Number of aphids (15 days)	
Dowling	Black Hawk from R. cathartica	2	6	60
(Rag1)	Black Hawk III from R. cathartica	1	1	
	Bronson, Michigan from R. cathartica	0	0	
	Hy 47 & 64, Illinois from R. cathartica	7	3	
	Irish Hills from R. cathartica	4	3	
	Jolliette College from R. cathartica	5	15	
	Pit, Indiana from R. cathartica	18	126	65
	Rock II from R. cathartica	10	26	

Example 4

Soybean Accessions that Possess Rag2

A set of 80 soybean germplasm accessions that were resistant to an Illinois soybean aphid isolate were challenged with the Ohio soybean aphid biotype that can overcome Rag1. The accessions listed in Table 3 had resistance not significantly different from PI200538. It is can therefore be deduced that these accessions also possess Rag2 (or possibly another gene effective against the Ohio biotype but not Rag1.)

TABLE 10

List of soybean accessions that had resistance equal to PI200538

71506 88508 Showa No. 1-4 200538 Sugao Zarai 230977 437696 San-haj-hun-mao- huan-dou 499955 507298 Sokoshin Kamigoumura 518726 Bao jiao huang 548237 T260H 548409 Sato 567391 Jiang se huang dou 567541B 567598B 587552 Nan jing da ping dir huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	0.0 1.7 1.0 1.3 1.0 1.0 1.0 1.0 1.5 1.6 1.7 1.7 2.0 g 1.0
200538 Sugao Zarai 230977 437696 San-haj-hun-mao-huan-dou 499955 Sokoshin Kamigoumura 518726 Bao jiao huang T260H 548237 T260H Sato 567391 Jiang se huang dou 567541B S67598B 587552 Nan jing da ping din huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.0 1.3 1.0 1.0 1.0 1.0 1.5 1.6 1.7 1.7 2.0 g
230977 437696 San-haj-hun-mao-huan-dou 499955 507298 Sokoshin Kamigoumura 518726 Bao jiao huang 548237 T260H 548409 Sato 567391 Jiang se huang dou 567541B 567598B 587552 Nan jing da ping din huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.3 1.0 1.0 1.0 1.0 1.5 1.6 1.7 1.7 2.0 g
437696 San-haj-hun-mao-huan-dou 499955 Sokoshin 507298 Sokoshin Kamigoumura 518726 548237 T260H 548409 Sato 567391 Jiang se huang dou 567541B 567598B 587552 Nan jing da ping dinuang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.0 1.0 1.0 1.0 1.5 1.6 1.7 1.7 2.0 1.0
huan-dou 499955 507298 Sokoshin Kamigoumura 518726 Bao jiao huang 548237 T260H 548409 Sato 567391 Jiang se huang dou 567541B 567598 S87552 Nan jing da ping dir huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.0 1.0 1.5 1.6 1.7 1.7 2.0 g
507298 Sokoshin Kamigoumura 518726 Bao jiao huang 548237 T260H 548409 Sato 567391 Jiang se huang dou 567541B S67541B 567545 Nan jing da ping din huang yi 1 587652 Nan jing da ping din huang yi 1 587664 Huang dou 587665 Huang dou 587666 Er dao zao 587669 Zan zi bai	1.0 1.5 1.6 1.7 1.7 2.0 g
Kamigoumura 518726	1.0 1.5 1.6 1.7 1.7 2.0 g 1.0
548237 T260H 548409 Sato 567391 Jiang se huang dou 567541B S67598B 587552 Nan jing da ping din huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.5 1.6 1.7 1.7 2.0 g 1.0
548409 Sato 567391 Jiang se huang dou 567541B S67541B 567598B S87552 Nan jing da ping di huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.6 1.7 1.7 2.0 g 1.0
567391 Jiang se huang dou 567541B 567598B 587552 Nan jing da ping din huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.7 1.7 2.0 g 1.0
567541B 567598B 587552 Nan jing da ping di huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.7 2.0 g 1.0
567598B 587552 Nan jing da ping din huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	g 2.0 1.0
587552 Nan jing da ping din huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	g 1.0
huang yi 1 587617 Jin tan qing zi 587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	8
587656 Huang dou 587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	1.0
587663 Zhong chun huang dou 587666 Er dao zao 587669 Zan zi bai	
dou 587666 Er dao zao 587669 Zan zi bai	1.0
587669 Zan zi bai	1.0
	1.0
FOR COR. 371 11.1	1.0
587677 Xiao li huang	1.0
587685 Da li huang 2	1.0
587693 Yu shan dou	1.3
587702 Qing pi dou	1.7
587717 Xiang yang ba yue:	ha 1.3
587732 Ying shan ji mu wo	1.1
587759 Song zi ba yue cha	1.0
587763 Jing huang 36	

TABLE 10-continued

30
TABLE 10-continued

	ybean accessions that had resis Rag2) against the Ohio soybea		_		ybean accessions that had resign (Rag2) against the Ohio soybear	
PI#	Name	Aphid Rating (0-4)	5	PI#	Name	Aphid Rating (0-4)
587800	Ying shan da li huang	1.0			dou	
587816	Bai mao dou	1.1		594514	Hua lian dou	1.1
587824	Ying shan qing pi cao	1.1		594554	Huang pi tian dou	1.1
587840	Du wo dou	1.1		594573	Lu pi dou	1.0
587861	Da qing dou	1.3	10	594592	Shi yue xiao huang	1.0
587870	Huang pi dou	1.3			dou	
587871	Bao mao dou	1.0		594595	Ba yue da huang dou	1.1
587873	Feng wo dou	1.7			(jia	
587876	Xi mao dou	1.0		594703	Qing pi dou –1	1.1
587897	Qing pi dou	2.1		594707	Da hei dou	1.3
587899	Ba yue bai	1.0	15	594822	Xi huang dou	1.0
587905	Xiao huang dou	1.0	13	594868	Huang dou	1.0
587972	Chang zi dou	1.0		594879	Huo shao dou	1.5
588000	Shi yue huang	1.1				
588040	Shan xing dou	1.0				
594421	Da du huang dou	1.6		Although	methods and cultivars	have been described in
594425	Xiao cao huang dou	1.1	20	detail for no	irnoses of clarity and i	ınderstanding it will be
594431	Chang pu qing dou	1.1	20	detail for purposes of clarity and understanding, it wis clear to those skilled in the art that equivalent cultivars, n		
594499	Luo ma aluo	1.1			-	
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The invention claimed is:

1. A method for reliably and predictably introgressing soybean *Aphis glycines* resistance conferred by a Rag2 gene into non-resistant soybean germplasm or less resistant soybean germplasm comprising:

providing a first soybean germplasm that has resistance to *Aphis glycines* conferred by said Rag2 gene;

providing a second soybean germplasm that lacks said resistance;

crossing the first soybean germplasm with the second soybean germplasm to provide progeny soybean germplasm;

analyzing said progeny germplasm by marker-assisted selection (MAS) to:

determine the presence of Rag2 resistance to *Aphis gly-cines* by detecting a resistance to *Aphis glycines* (Rag2) locus that maps to soybean linkage group F of said soybean germplasm, wherein said Rag2 locus is flanked on opposite sides by markers Soyhsp176 and Satt510, and wherein the Rag2 locus comprises allelic DNA sequences that control resistance to *Aphis gly-cines*; and

determine the presence or absence of said allelic sequences linked to the Rag2 gene coding for resistance to *Aphis glycines* in said germplasm; wherein the presence or absence of said allelic sequences linked to said gene is determined by identifying a

- region of DNA that displays allelic variation between said first and second soybean germplasm linked to said Rag2 locus; and
- determining that said gene coding for Rag2 resistance is present in said soybean germplasm when said region of DNA comprises a polymorphism linked to Rag2 resistance and present in PI 200538; and
- determining that said gene is not present in said germplasm when said region of DNA does not comprise said polymorphism linked to Rag2 resistance and present in PI 200538; and
- selecting progeny that tests positive for the presence of Rag2 resistance to *Aphis glycines* as being soybean germplasm into which germplasm having Rag2 resistance to *Aphis glycines* has been introgressed.
- 2. The method of claim 1 wherein said allelic form of DNA is identified by MAS comprising selecting a marker from the group consisting of Satt510, Soyhsp176, Sat_234, and Sat_297 and other markers that map to within 20 cM of any of these markers on linkage group F.
- 3. The method of claim 1 wherein said second soybean 20 germplasm has resistance to *Aphis glycines* conferred by a gene on soybean linkage group M.
- **4**. The method of claim **1** wherein said second soybean germplasm provides a desirable agronomic trait.
- **5**. The method of claim **1** wherein said second soybean ₂₅ germplasm does not contain a determinant for *Aphis glycines* resistance.

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- 6. The method of claim 1 wherein said first soybean germplasm that has resistance to *Aphis glycines* is germplasm of a cultivar having Rag2 *Aphis glycines* resistance selected from the group consisting of Pl200538; Pl437696; Pl499955; Pl507298; Pl548237; Pl567391; Pl587656; Pl587669; Pl587775; Pl587870; Pl587871; Pl587899; Pl587972; Pl588000; Pl588040; Pl594431; Pl594499; Pl594573; Pl594707; Pl594822; Pl594879; and progeny thereof having Rag2 *Aphis glycines* resistance.
- 7. The method of claim 6 wherein said soybean germplasm that has resistance to *Aphis glycines* is of PI200538.
- 8. The method of claim 1 for the production of an inbred soybean cultivar adapted for conferring, in hybrid combination with a suitable second inbred, resistance to *Aphis glycines* conferred by the Rag2 gene locus, wherein said second soybean germplasm is high yielding in hybrid combination with said first soybean germplasm that has said resistance to *Aphis glycines*; comprising the further step of self-crossing the plants that tested positive for the presence of said *Aphis glycines* resistance until a line is obtained that is homozygous for said *Aphis glycines* resistance and adapted for conferring, in hybrid combination with a suitable second inbred, said resistance to *Aphis glycines* resistance conferred by the Rag2 gene locus.

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