

# De novo assembly of soybean wild relatives for pan-genome analysis of diversity and agronomic traits

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Wild relatives of crops are an important source of genetic diversity for agriculture, but their gene repertoire remains largely unexplored. We report the establishment and analysis of a pan-genome of *Glycine soja*, the wild relative of cultivated soybean *Glycine max*, by sequencing and *de novo* assembly of seven phylogenetically and geographically representative accessions. Intergenomic comparisons identified lineage-specific genes and genes with copy number variation or large-effect mutations, some of which show evidence of positive selection and may contribute to variation of agronomic traits such as biotic resistance, seed composition, flowering and maturity time, organ size and final biomass. Approximately 80% of the pan-genome was present in all seven accessions (core), whereas the rest was dispensable and exhibited greater variation than the core genome, perhaps reflecting a role in adaptation to diverse environments. This work will facilitate the harnessing of untapped genetic diversity from wild soybean for enhancement of elite cultivars.

Annual wild soybean (*Glycine soja* Sieb. & Zucc.) is the closest relative and antecedent of cultivated soybean (*Glycine max* (L.) Merr.), one of the world's primary sources of plant protein and vegetable oil. Whereas *G. max* has lost substantial genetic diversity through successive bottlenecks owing to domestication and selection for traits to increase yield under intensive human cultivation, *G. soja* is distributed across a broad geographical range (24–53° N, 97–143° E) and has adapted to a variety of ecological conditions<sup>1</sup>, providing a promising source of novel genes for soybean improvement needed to respond to rapid population growth and environmental changes.

Since the publication of the soybean genome (var. Williams 82 (GmaxW82))<sup>2</sup>, attempts have been made to characterize *G. soja* through genome resequencing<sup>3–5</sup>. However, resequencing is limited in terms of capturing many types of structural variation, especially for crop wild relatives, which are more genetically diverse than their domesticated counterparts. High levels of sequence similarity are required to map resequencing short reads to a reference genome at the expense of losing information from more diverged genomic regions. Presence-absence variation (PAV) and copy number variation (CNV), which are often associated with agronomic traits<sup>6</sup>, may also

be missed<sup>7</sup>. Moreover, a single genome is insufficient to represent the genomic content of a predominantly selfing (autogamous) species, such as *G. soja*, in which individuals are distinct from one another due to low levels of genetic exchange and recombination<sup>8</sup>. Thus, the *de novo* construction of a pan-genome for a species, consisting of a core genome shared among individuals and individual-specific or partially shared dispensable genome, is necessary to capture the majority of genetic diversity within a species<sup>9</sup>.

To fully characterize the genomic content and molecular evolutionary history of *G. soja*, the undomesticated gene pool of soybean<sup>8</sup>, we constructed a pan-genome by sequencing and assembling seven *G. soja* accessions *de novo*. Within this pan-genome, the dispensable gene set was found to have evolved more rapidly and be more variable than the core gene set, and genes under selection exhibited relatively little overlap between the seven *G. soja* lineages, indicating that local adaptation may have affected nonoverlapping sets of genes. Some of the genes with structural variation between *G. soja* and GmaxW82, based on homology and co-localization with mapped quantitative trait loci (QTLs), may be associated with adaptation to various abiotic and biotic stresses. Our data also push the divergence between *G. max* and

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*G. soja* back to ~0.8 million years ago (mya), from a recent estimate of ~0.27 mya<sup>4</sup>. This work illustrates the value of *de novo* assemblies in building and characterizing large eukaryotic pan-genomes, and provides candidate genes for introgression into cultivated soybean for crop improvement.

## RESULTS

### *De novo* genome assemblies of seven *G. soja* accessions

Seven *G. soja* accessions, representing the geographical adaptation within the species, were selected and named GsojaA to GsojaG (Supplementary Figs. 1 and 2 and Supplementary Table 1). Six accessions represent major genetically distinct clusters (North, Huanghuai and South regions of China, and Japan, Korea and Russia) of *G. soja* and 87.1% of the genetic diversity<sup>8</sup>. Another was selected from the Northeast region (NER) of China, a predicted soybean domestication center<sup>10</sup>, to better represent the geographic distribution of *G. soja*. Each genome was sequenced with Illumina HiSeq2000 using a combination of libraries with insert sizes of 180 bp, 500 bp and 2 Kbp (Supplementary Table 2), for an average of 111.9 fold coverage (Table 1). The estimated genome sizes ranged from 889.33 Mbp for GsojaG (93.6% of the GmaxW82) to 1,118.34 Mbp for GsojaD (117.7% of GmaxW82). Variation in genome size could be partially explained by differences in abundance of repetitive sequences (Supplementary Fig. 3).

Each genome was assembled using SOAPdenovo<sup>11</sup>, a *de novo* genome assembler based on a de Bruijn graph algorithm, and resulted in contig N50 sizes ranging from 8 to 27 Kbp and scaffold N50 sizes ranging from 17 to 65.1 Kbp (Table 1). The assemblies were validated by alignment to GmaxW82, covering more than 94% of the 54,175 genes (Glyma1.1 annotation, <http://www.phytozome.net>, Table 1). The coverage of GmaxW82 genes by assembled genomes was either higher than (5 *G. soja* genomes) or comparable to (2 *G. soja* genomes) the coverage by aligning resequencing reads. As such, the current assemblies enabled accurate detection of variation and comparative analyses within genic regions. Additional mate-pair libraries with large inserts would further improve assembly size, thereby resolving variation in nongenic regions.

Protein-coding genes were annotated for each genome by integrating homology searches, mRNA expression evidence, and *ab initio* prediction (Supplementary Table 3). After correction for gene fragments due to incomplete assembly, we estimated an average of 55,570 genes per genome (Table 1), slightly more than the 54,175 genes in GmaxW82. The quality of the annotation was supported by the observation that 89.24–91.93% of genes had at least one ortholog in the GmaxW82 genome and 63.94–71.19% of the predicted genes were expressed, as determined by RNA-seq of mixed tissues of corresponding accessions. In addition, a majority (96.15–97.16%) had protein homologs in other plant genomes (Supplementary Table 4).

Underestimation of gene number may result from lack of evidence to support prediction. More likely, however, the gene number may be inflated as it includes genes split across contigs and genes on separately assembled haplotypes.

### Variation between *G. soja* and GmaxW82

To identify variation between *G. soja* genomes and GmaxW82, we designed a computational pipeline that takes advantage of the sequencing reads as well as assembled genomes to catalog variation including single-nucleotide polymorphisms (SNPs), insertions or deletions (indels), CNV and PAV (Supplementary Fig. 4). There were 3.63–4.72 million SNPs in individual samples, with 0.12–0.15 million in coding sequence regions (CDS). This included 1,764 loci where *G. soja* had a stop codon and GmaxW82 did not, and 2,286 loci where GmaxW82 had a stop codon and *G. soja* did not (Supplementary Table 5). The assembly-based method enabled us to find more SNPs than by resequencing alone, especially in divergent regions where unassembled short sequencing reads are difficult to be mapped (Fig. 1a and Supplementary Figs. 5 and 6). Even though *G. soja* is a predominantly selfing species, three of the seven (GsojaA, GsojaC and GsojaD) had relatively high heterozygous SNP rates (Supplementary Table 5). The high rate of heterozygosity may have complicated *de novo* assembly and resulted in inflated gene numbers due to divergent haplotypes being represented as separate scaffolds in the assembly.

We detected 0.50–0.77 million indels in *G. soja* as compared to GmaxW82, of which 93.3% (70/75) of randomly selected indel alleles were validated by Sanger sequencing (Supplementary Table 6 and Supplementary Figs. 7 and 8). Although many indels were trinucleotides, 2,989–4,181 resulted in frameshifts (Supplementary Fig. 9 and Supplementary Tables 7 and 8). For example, three indels were identified in one of the two homologs of *Spiral2* (*spr2*) (*AT4G27060*), a key microtubule gene for directional cell elongation that is associated with the right-handed helical growth in *Arabidopsis*<sup>12</sup> (Fig. 1b). In *Glyma02g25230*, the *Spiral2*-homolog, three guanines were deleted at positions 695, 707 and 714 in all seven *G. soja* genomes, resulting in five amino acid changes in one of the HEAT-repeat motifs. Thus, *Glyma02g25230* is a candidate gene that may be responsible for the change from the twining growth habit found in *G. soja* to erect growth found in *G. max* (Supplementary Fig. 10).

We identified CNV in genic regions and found hundreds of genes that had either gained or lost copies in individual *G. soja* accessions compared to GmaxW82. Of 1,978 genes affected in the *G. soja* accessions, 1,179 had CNV loss, 726 had CNV gain and 73 had both CNV loss and gain (Supplementary Tables 9 and 10). Gene Ontology (GO) analysis indicated that genes related to abiotic and biotic stress tolerance, such as resistance (R)-genes with nucleotide-binding site (NBS) or NBS-leucine-rich repeat (LRR) domains and transcription factors, were significantly ( $P < 0.01$ , chi-squared test) enriched in genes

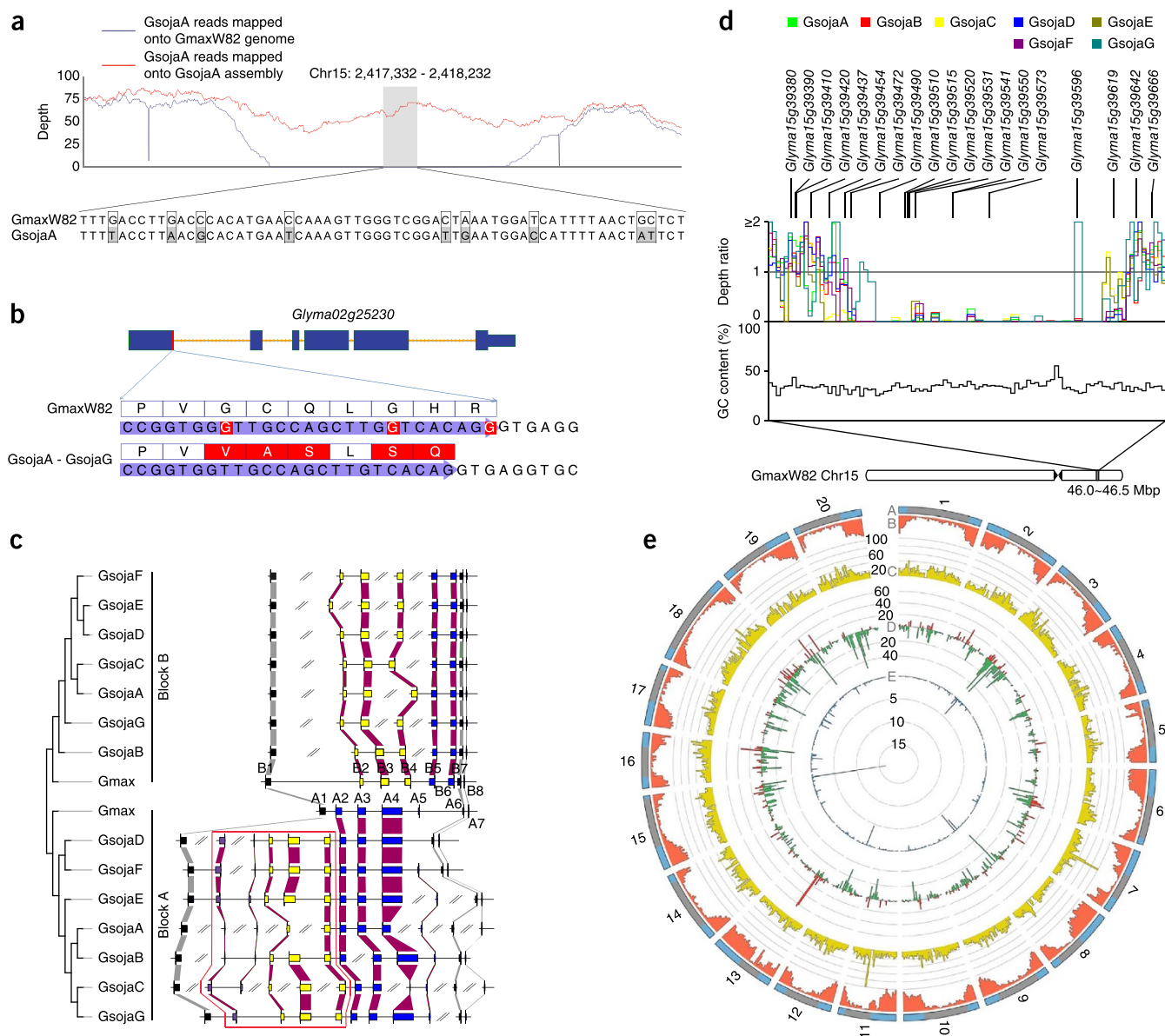
**Table 1** Sample information, assembly and annotation for seven *G. soja* accessions

ID	GsojaA	GsojaB	GsojaC	GsojaD	GsojaE	GsojaF	GsojaG
Origin	Zhejiang, China	Ibaraki, Japan	Chungchong Puk, Korea	Shandong, China	Shanxi, China	Heilongjiang, China	Khabarovsk, Russia
Fold sequencing depth (x)	117.7	115.6	122.7	136.3	103.1	83.7	104
Estimated genome size (Mbp)	981.04	1,000.8	1,053.78	1,118.34	956.43	992.66	889.33
Assembled genome size (Mbp)	813	895	841	985	920	886	878
Contig N50 (Kbp)*	9	22.2	8	11	27	24.3	19.2
Scaffold N50 (Kbp)	18.3	57.2	17	48.7	65.1	52.4	44.9
GmaxW82 gene coverage by alignment (%)	94.98	94.46	95.89	94.97	94.75	94.65	93.25
GmaxW82 gene coverage by assembly (%)	94.95	96.26	95.08	96.15	96.64	96.42	96.2
Predicted gene number	58,756	56,655	60,377	62,048	58,414	57,573	58,169
Refined gene number	55,061	54,256	56,542	57,631	55,901	54,805	54,797

affected by CNV. This indicates that these genes are evolutionarily labile and may be involved in adaptation to local environments and/or interaction with pathogens.

In total, 2.3–3.9 Mbp of *G. soja*-specific PAV (defined as >100 bp and <95% identity) was present in the *G. soja* genomes (Supplementary Table 11), less than the 8.3 Mbp reported for a single

previously resequenced *G. soja* genome<sup>4</sup>. Our lower PAV discovery rate may be due to the use of more stringent filtering criteria, including the removal of unassembled sequences in *G. max* and putative microbial sequences. There were 338 genes found to have at least 50% of their coding sequences composed of *G. soja*-specific sequences (Supplementary Table 12 and Supplementary Fig. 11). Of these



**Figure 1** Examples of variation between *G. soja* and *G. max*. **(a)** A diverged region on chromosome 15. Top panel, coverage of GmaxW82 (blue) and GsojaA genome (red) by GsojaA reads. Comparison between GmaxW82 and *G. soja* sequences show nine SNPs in a 62-bp fragment at bottom. This highly diverged region could not be aligned by mapping GsojaA reads to GmaxW82. **(b)** Indels in *Glyma02g25230*. GmaxW82 gene structure is shown at top (exons in blue) and five amino acid changes caused by three indels in *G. soja* are shown in red at bottom. **(c)** *G. soja*-specific PAV. Syntenic genomic region on chromosome 14 (Block A) compared to its homolog on chromosome 02 (Block B). Genes in each block are numbered A1 to A7 and B1 to B8, respectively. An 8-Kbp *G. soja*-specific region is outlined in red, in which the three genes absent in the *G. max* region but with homologs on block A are in yellow, and the two genes with no homologs either on block A or elsewhere in *G. max* are in purple. Genes exhibiting relatively conserved synteny between species are marked in blue. The anchor point of each block is shown in black. Homologous genes are connected by magenta shading and the anchor points of each block are connected by gray shading. A phylogenetic tree was built using genes in red (*Glyma02g38310*). **(d)** *G. max*-specific PAV. Depth distribution of *G. soja* genomes on GmaxW82 chromosome 15 (46–46.5 Mbp) is shown with y axis representing the depth ratio on top and GC content below. The sequence depth ratio was calculated using the average depth in 10-Kbp windows for each *G. soja* accession over the genome-wide average. Seven of the 19 genes with low read depth in *G. soja* accessions were identified as *G. max*-specific genes, for example, *Glyma15g39573*. **(e)** Distribution of genetic variation across the soybean genome. A, chromosome ideograms for *G. max*; B, gene density; C, distribution of genes containing large-effect mutations; D, CNV (red for gain, green for loss); and E, *G. max*-specific PAV. For circle C, D and E, numbers represent percentages of genes with corresponding mutations over total genes in 1-Mbp window.



genes or genomic loci affected by putative PAV, 37 were selected for PCR-based genotyping across GmaxW82 and the seven *G. soja* accessions (using 43 primer pairs). All 37 loci were concordant with the predicted PAV that was present in *G. soja* and absent in *G. max* (Supplementary Tables 13 and 14 and Supplementary Fig. 12). Of the *G. soja*-specific PAV genes, 34 were inferred to be involved in defense response, 33 in cell growth and 15 in photosynthesis, by GO category enrichment analysis (Supplementary Table 15).

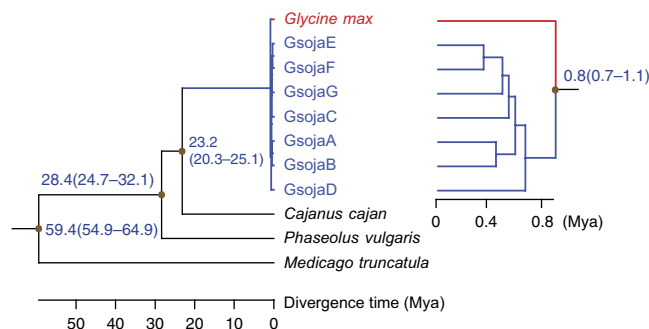
As an example of genic PAV, an 8-Kbp *G. soja*-specific region was identified in all seven accessions (Block A) on chromosome 14 and validated by PCR genotyping (Supplementary Fig. 13). A paralogous region (Block B) on chromosome 02 in *G. soja* and *G. max* contained three genes missing in Block A that were related to biotic and abiotic stress tolerance or plant development (*Glyma02g38290*, homolog of *ZRZ/ZF14* or *BCD1*; *Glyma02g38300*, homolog of *SUT4* and *Glyma02g38310*, homolog of *GLY1*)<sup>13,14</sup> (Fig. 1c). Of the 56 resequenced *Glycine* accessions<sup>3–5</sup>, 17 of 27 *G. soja* accessions and three of 29 *G. max* accessions contained this putative *G. soja*-specific sequence, indicating that this PAV is segregating in both *G. soja* and *G. max* but the frequency has been reduced in *G. max*, perhaps due to the domestication bottleneck.

The total length of *G. max*-specific PAV (1.86 Mbp) was less than that of *G. soja*-specific PAV. The GC content for these regions (32.9%) was not significantly different from the whole genome average (34.1%), rejecting the possibility of sequencing bias. A total of 16 genes were identified to contain *G. max*-specific sequences (>50% of the length of the CDS affected), including four genes that were entirely *G. max*-specific (Fig. 1d). This number was less than the 712 found in an earlier comparison of a single *G. soja* accession and GmaxW82 (ref. 4), which may be due to increased sampling of *G. soja* diversity and more stringent criteria. Based on KEGG analysis of the four *G. max*-specific genes, *Glyma01g37051* is involved in energy metabolism (ko00190 and ko00194), *Glyma13g36351* and *Glyma15g39541* in lipid metabolism (ko00590, ko00591 and ko00592), and *Glyma15g39596* in cell growth and death (ko04210). *Glyma01g37051* is a homolog of *AtLFNR1* (ferredoxin-NADP<sup>+</sup>-oxidoreductase, AT5G66190), which in *Arabidopsis* is involved in photosystem I-dependent cyclic electron flow and affects rosette size<sup>15</sup>.

Variation was distributed unevenly throughout the genome (Fig. 1e). Genes in recombination-suppressed pericentromeric regions were more significantly ( $P < 0.01$ , chi-squared test) affected than those in chromosome arms (Supplementary Table 16), as was also observed in *Zea mays* (maize)<sup>16</sup>. The majority of genes affected by large-effect variations (including SNPs or indels causing stop codon gain or loss and frameshift), CNV or genome-specific sequences were rare events usually found in only one of the seven *G. soja* accessions. These rare variants may be useful in improving soybean cultivars through breeding, as has been indicated by studies in maize and wheat, in which rare alleles were found to be associated with QTLs for agronomic traits<sup>17</sup>.

### Evolution of the *G. max* and *G. soja* species complex

We next constructed a phylogenetic tree using 670 conserved, single-copy, gene orthologs from *G. soja*, *G. max* and three other sequenced legume species (Fig. 2). Using the divergence time between *Medicago* and *Glycine* as a calibration point<sup>18</sup>, we estimated that pigeon pea and soybean diverged ~23 mya, and common bean and soybean ~29.5 mya, similar to previous estimates<sup>18,19</sup>. We also estimated the sequenced *G. soja* accessions and *G. max* diverged ~0.8 mya, earlier than previously estimated (0.27 mya) based on a single *G. soja* accession<sup>4</sup>. This difference is likely due to the inclusion of more diverged *G. soja* accessions in the current study.



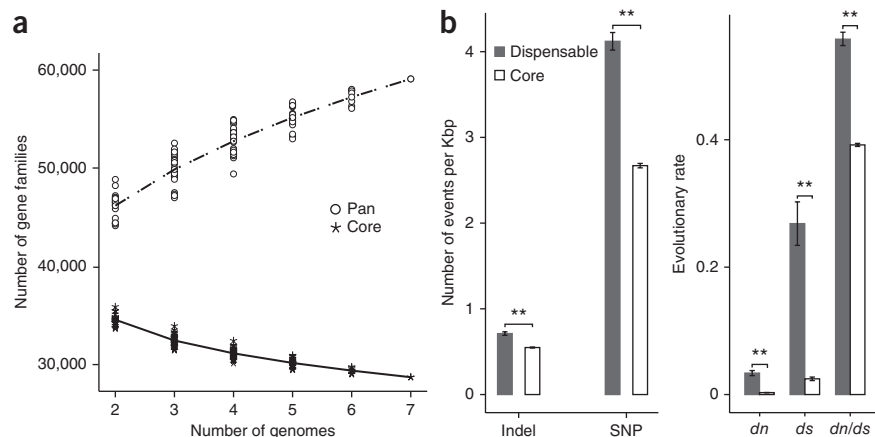
**Figure 2** Divergence among *G. max*, *G. soja* and selected legumes. Left, Relationship and divergence times among sequenced legume species, including *Cajanus cajan*, *Phaseolus vulgaris* and *Medicago truncatula*. Right, relationship of sequenced *G. soja* accessions and *G. max* (GmaxW82).

The divergence time between *G. soja* and *G. max* predates the estimated domestication time of soybean of ~5,500 ya<sup>20</sup>, therefore divergent selection may have contributed to the differentiation of the two subspecies before domestication of *G. max*. Using the branch-site likelihood ratio test, we identified 682 genes that showed evidence of positive selection in the GmaxW82 genome (Supplementary Table 17). These genes were enriched in diverse biological functional categories related to abiotic stress regulation, including stomatal complex development, voltage-gated potassium channel activity, proline metabolism and nitrate transport (Supplementary Table 18). In the *G. soja* accessions, 175–408 such genes were identified in individual wild soybean accessions and none was shared among all accessions. Only ten positive-selection genes were shared by at least three *G. soja* accessions and 132 such genes were shared by two accessions. Genes that underwent positive selection may have contributed to the adaptation of *G. soja* accessions to different environments, as evidenced by the species' wide range of geographical distribution and lack of shared positive-selection genes.

Changes within the *G. soja* pan-genome (total gene set) and core genome (genes shared among all seven genomes) were analyzed (Supplementary Fig. 14). The number of total genes increased as additional genomes were added and, in contrast, the number of shared genes decreased with additional *G. soja* genomes (Fig. 3a). The average pan-genome size of any two accessions accounted for 78.2% of that found using all seven accessions, confirming that a single genome does not adequately represent the diversity contained within *G. soja*. The pan-genome size was asymptotic with size of the seven genomes and an initial *G. soja* pan-genome with 59,080 gene families and an overall size 986.3 Mbp was constructed (Supplementary Table 19 and Supplementary Fig. 15). Nearly half (48.6%) of the gene families and 80.1% of sequences conserved across all seven *G. soja* genomes were core genomic units (Supplementary Figs. 14 and 15). Approximately half of the gene families (51.4% or 30,364) were present in more than one, but not all seven *G. soja* genomes, and represent the dispensable genome. As might be expected, unique gene families, those found only in one sample, were the least frequent and, among the seven accessions, GsojaD had the greatest amount of accession-specific gene families (Supplementary Fig. 14).

Owing to recent acquisition and deletion events<sup>21</sup>, the dispensable gene set was more variable than the core gene set. The frequency of SNPs in the core gene set was estimated to be 2.67 sites per Kbp, significantly less than 4.12/Kbp in the dispensable gene set ( $t$ -test,  $P < 0.01$ ). The same bias was observed in the frequency of indels

**Figure 3** *G. soja* pan-genome components and evolution of core and dispensable genomes. (a) Increase and decrease in gene families in pan-genome and core genome, respectively, with every additional *G. soja* genome. (b) Frequency of indels and SNPs (left panel) and *dn*, *ds* and *dn/ds* (right) in core and dispensable genomes. \*\* $P = 0.001$ , *t*-test. Error bars, mean  $\pm$  s.d.



(Fig. 3b and Supplementary Table 20). Moreover, the dispensable genes had significantly higher *dn*, *ds* and *dn/ds* ratio values than did core genes ( $P < 0.01$ , *t*-test; Fig. 3b), indicating that the dispensable genes have undergone weaker purifying selection and/or greater positive selection than core genes.

Core genes were enriched in biological processes including growth, immune system processes, reproduction, reproductive processes, cellular processes, and cellular component organization or biogenesis (Supplementary Table 21). Three categories were enriched in dispensable genes: receptor activity, structural molecule activity and antioxidant activity in molecular function. The core genes were more functionally conserved than the dispensable genes: 58.3% of the dispensable genes and 33.9% for the core genes set could not be assigned any functional annotation. Moreover, 95.5% of core genes had homologs in other species based on BLAST searches to 32 plant genomes (excluding soybean), significantly more than the dispensable gene set (83.5%, chi-squared test,  $P < 0.01$ ). These results confirm that lineage-specific genes evolve faster than genes that are shared between species, either by means of a higher evolutionary rate<sup>22</sup> or a higher gene loss rate<sup>23</sup>.

### *G. soja* versus *G. max*: genomic basis of agronomic traits

R-genes with NBS domains mediate effector-triggered immunity acting as detectors for pathogen virulence proteins<sup>24</sup>. The quantity of R-genes with a given domain architecture and the number of unique domain architectures varied between the two species. *G. soja* contained more R-gene domain architectures (25) than *G. max* (14) (Fig. 4a). The 11 lineage-specific R-gene domain architectures found in *G. soja* accessions were part of the dispensable genome, possibly reflecting adaptation to biotic stresses. The *G. soja* genome contained a broader range of NBS R-gene domain architectures than did *G. max*, but *G. max* had more NBS-encoding genes (460 versus 334 in GsojaA to 382 in GsojaD) (Supplementary Table 22). R-genes showed more copy number loss in *G. soja* accessions (88 lost versus 33 gained and 8 copy number gain/loss, Supplementary Table 23), several of which were known to confer resistance to soybean mosaic virus<sup>25</sup>, Asian soybean rust<sup>26</sup> and *Phytophthora* root rot<sup>27</sup>, and be involved in legume-rhizobia symbiosis<sup>28</sup>. In addition, five structurally diverged regions (located on chromosomes 03, 06, 07 and 18) within cultivated *G. max*<sup>29</sup>, had a high frequency of CNV in the pan-genome (Supplementary Fig. 16).

Genes controlling flowering time were likely involved in adaptation to new climatic regions. Comparing the *G. soja* genomes and GmaxW82, flowering time-related genes with large-effect variation were identified, including light receptor *PHYA*, floral integrator *FLOWERING LOCUS T* (*FT*) and meristem identity protein *LEAFY* (*LFY*) genes (Fig. 4b). Two of the four homologs of *PHYA* in soybean, *GmPHYA2* and *GmPHYA3*, correspond to major flowering loci, *E4* (ref. 30) and *E3* (ref. 31), respectively. We found mutations in *GmPHYA4* (*Glyma03g38620*), resulting in changes and deletions of

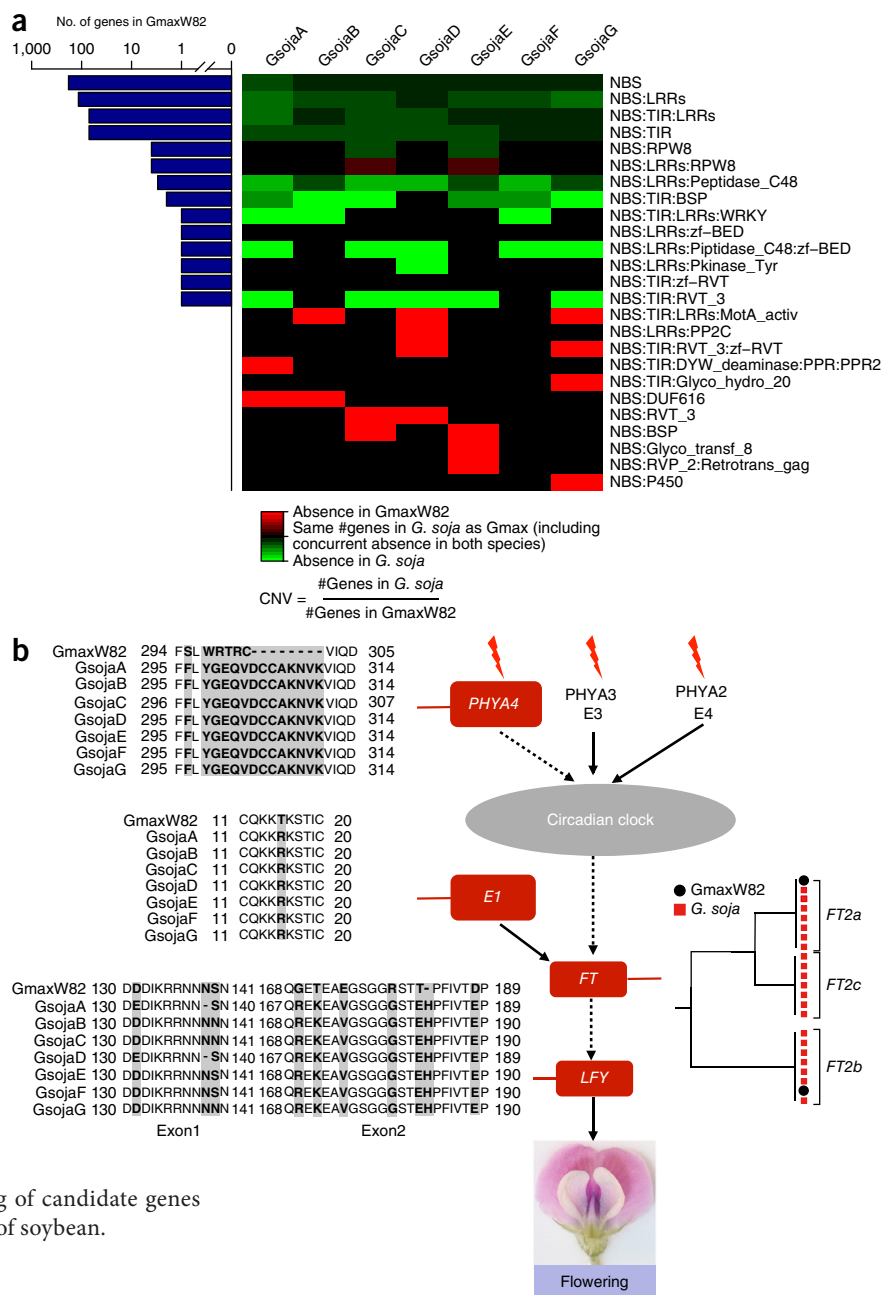
amino acid (AA) in the N-terminal P3/GAF domain (Supplementary Figs. 17 and 18). Among ten *FT* homologs in soybean, *GmFT2c* showed the closest relationship to *GmFT2a*, a candidate gene that coordinates flowering<sup>32</sup>. In GmaxW82, *GmFT2c* was missing ~50% of the N-terminal peptides (Fig. 4b and Supplementary Fig. 19); in *G. soja*, however, *GmFT2c* had a complete gene structure and was expressed. *LFY*, which regulates the timing of floral induction in *Arabidopsis*, had two putative orthologous copies in GmaxW82 [*GmLFY1* (*Glyma04g37900*) and *GmLFY2* (*Glyma06g17170*)]. Two 3-bp indels and nine nonsynonymous SNPs were identified in *GmLFY1* among the seven *G. soja* accessions, all in exons 1 and 2, forming variation hotspots (Fig. 4b). Both *GmFT2c* and *GmLFY1* were located in or near QTL, controlling the period of reproductive growth stage and maturity time<sup>33</sup>, thus providing gene candidates that may underlie flowering time in soybean.

Oil and fatty acid content of soybean has been intensely selected such that *G. max* produces nearly twice as much oil as *G. soja*<sup>34</sup>. A total of 1,332 genes were annotated in GmaxW82 based on homology to genes in 24 acyl lipid subpathways in *Arabidopsis* (Supplementary Table 24)<sup>35</sup>. Of these genes, 15.9% contained CNV and/or large-effect SNP and/or indels between the *G. soja* accessions and GmaxW82. Although five subpathways exhibited a significantly low frequency of variation ( $P < 0.05$ , Fisher's exact test; Supplementary Table 24), a number of gene variants were identified. Fatty acids are precursors for jasmonate production in the oxylipin pathway in plants following stress<sup>36</sup>. *SAG101* (*senescence-associated gene101*) in *Arabidopsis*, which encodes a triacylglycerol lipase to hydrolyze triacylglycerol to monoacylglycerol and produce free fatty acid, plays important roles in plant innate immunity against biotrophic pathogens<sup>37</sup>. In this study, large-effect SNP and/or indels were found in three clustered homologs of *SAG101* (*Glyma13g04561*, *Glyma13g04651* and *Glyma13g04571*) and are candidate genes for nearby QTL that affect oil content and confer resistance to *Phytophthora sojae*<sup>38</sup> (Supplementary Fig. 20).

Triacylglycerol is a major storage lipid in soybean and acyl-CoA: diacylglycerol acyltransferase (DGAT) is a key enzyme that catalyzes diacylglycerol to triacylglycerol, determining seed oil content in *Arabidopsis*<sup>35</sup>. Among 113 potential triacylglycerol biosynthesis genes, 10 had putative loss-of-function frameshifts due to large-effect SNP and/or indels, including the DGAT homolog (*Glyma16g21960*, *GmDGAT2B*) and a homolog of *Abscisic Acid Insensitive 4* (*ABI4*) (*Glyma14g06290*), a transcription factor that regulates triacylglycerol synthesis in *Arabidopsis* by activating *DGAT1* expression during nitrogen deficiency (Supplementary Fig. 21)<sup>39</sup>. A frame-shift mutation in *GmDGAT2B* was previously found to be associated with reduced seed oil concentration<sup>40</sup>. Another DGAT homolog, *DGAT1B*

## ARTICLES

**Figure 4** Variation of resistance genes and flowering time-related genes. **(a)** Heatmap for CNV of resistance-related PFAM gene categories across all accessions. Copy number ratio of gene categories in each box was calculated as the number of genes in each *G. soja* accession over number of genes in GmaxW82 within the same gene category. Red represents greater gene numbers in *G. soja* than GmaxW82, whereas green indicates fewer. Gene numbers for GmaxW82 for the first 14 gene categories are shown on left. In categories where GmaxW82 does not have any genes (gene number = 0), black indicates the absence of genes in a given *G. soja* accession, while red indicates the presence of genes in *G. soja* vs. none in GmaxW82. **(b)** Large-effect mutations and new gene copy in flowering time-related genes. In *PHYA4*, *E1* and *LFY*, amino acid changes resulting from large-effect SNPs and indels are shown. In the *FT* gene family, a new copy named *FT2c* was found in all seven *G. soja* accessions.



(*Glyma17g06120*), negatively associated with seed protein content<sup>40</sup>, was found to have been positively selected during domestication (protein and oil content are negatively correlated).

In contrast to *G. soja*, cultivated soybean has larger organ sizes, thicker stems and shorter plant height (Supplementary Fig. 10). A QTL region controlling plant height, lodging and yield spans ~1.2Mbp on chromosome 9 (ref. 41). In this region, four genes were identified with large-effect SNP and/or indels (Supplementary Table 25). GO analysis implicated all four genes in plant development, and work in other species has shown these genes to be either directly or indirectly involved in cytokinin metabolism<sup>42,43</sup>. The identification of large-effect SNP and/or indels between *G. soja* and GmaxW82 can facilitate discovery and cloning of candidate genes related to the domestication and improvement of soybean.

## DISCUSSION

Our work illustrates the advantage of *de novo* assembly in detecting genetic variation that would not have been found by resequencing alone. The majority (94%) of *G. max* genes were covered by our *G. soja* assemblies. Thus, we were able to identify structural variation, including 338 PAV, 1,978 CNV, and a series of SNPs and indels in highly divergent genic regions, where only assembled sequences could be mapped. Using this data set, we estimated that the divergence time between *G. soja* and *G. max* was nearly three times earlier than a previous estimate based on a single *G. soja* accession<sup>4</sup>.

The individual genomes and the pan-genome could be improved by additional sequencing of larger insert libraries to increase contig and scaffold sizes, which would allow full exploration of the nongenic parts of the genome. These improvements would also enable further interrogation of chromosome-scale structural variation, such as inversion and translocation events. Given the structural complexity of plant genomes, further advances in sequencing technology and bioinformatics tools are needed to overcome difficulties in genome

assembly and annotation to deal with highly similar transposable elements and other abundant repetitive sequences, heterozygosity and polyploidy<sup>44</sup>. Furthermore, the concept of a pan-genome should be expanded beyond the DNA level to include variation in gene regulation and expression.

This study confirms that a single genome does not adequately represent the diversity contained within a species<sup>8</sup>. We constructed a *de novo* assembly-based pan-genome of a crop and crop wild relatives, by sequencing seven representative wild soybean accessions from East Asia<sup>8</sup>, the center of *G. soja* diversity and domestication center of *G. max*. The *G. soja* pan-genome was 30.2 Mbp larger than the genome of a single resequenced accession<sup>4</sup>. The number of accessions required to represent the majority of genetic variation within a species is case-dependent<sup>45</sup>, and cataloguing a pan-genome will never be complete. The observation that the *G. soja* pan-genome size began to level off suggests that we have captured a representative portion of



the species' gene pool. Sequencing additional samples may further reveal allelic structure and variation within and between populations, thereby allowing investigation into local adaptation and migration. Increasing the sample size may also add to the dispensable genome, which is characterized by greater variation than the core genome and is potentially involved in environmental adaptation and organismal interactions<sup>46</sup>. Approximately 20% of the *G. soja* sequences and 51.4% of gene families were found to be dispensable, an interesting juxtaposition to the recently published pan-transcriptome of 503 maize samples where 82.7% of the representative transcript assemblies were dispensable<sup>47</sup>, supporting the prediction that an outbred species such as maize would have a larger dispensable genome than an autogamous species<sup>45</sup>.

The pan-genome of *G. soja* shows the extent of novel genes and alleles in wild relatives that can be introgressed into crops, which typically have lower genetic diversity as the result of domestication and breeding<sup>48,49</sup>. Candidate genes associated with adaptation to various abiotic and biotic stresses may contribute to increased resilience to climate variability in cultivated soybean. Genes with structural variation and large-effect variation between *G. soja* and *G. max* (GmaxW82) that associate with agronomic phenotypes, as inferred from homology and comparison with mapped QTLs, can be used to develop molecular markers for these segments and test new allelic combinations. Newly identified genetic variation for genomic regions that have been fixed in *G. max* can be used to design crosses to determine if these fixed regions underlie phenotypes of agricultural value, providing additional candidate genes for the development of new varieties. Further, these data should enable the crop breeding community to more effectively use molecular approaches, such as genomic selection, to reduce the yield drag often associated with introgression from wild species.

## METHODS

Methods and any associated references are available in the [online version of the paper](#).

**Accession codes.** Genome assembly data, BioProject: [PRJNA195632](#). Unassembled sequencing reads and raw sequencing reads of the transcriptome, SRA: [SRP040255](#).

*Note: Any Supplementary Information and Source Data files are available in the online version of the paper.*

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## AUTHOR CONTRIBUTIONS

L.Q., R.L., S.A.J., Ying-hui Li, G.Z., and J.M. conceived the study and jointly wrote the paper. Ying-hui Li, S.Z., X.S., Yan-fei Li, H.H., Z.L., K.W., Li-juan Zhang and R.C. provided DNAs and performed PCR validation and Sanger sequencing. G.Z., R.L., W.J., Z.Z., J.Z., Q.Z., Y.W., H.R., H.W., J.L. and Z.J. performed sequencing, genome assembly and genome annotation. Ying-hui Li, G.Z., W.J., J.Z., S.A.J., J.M., J.L., Y.G., Y.S., Z.Z., B.T., J.S., L.J., Liangtao Zheng, Y.H., G.K., C.K.L.Y., W.Z., R.G., W.L., S.C. and L.Q. performed comparative, population and evolutionary biology analyses.

## COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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## ONLINE METHODS

**DNA and RNA sequencing of seven *G. soja* accessions.** Seven *G. soja* accessions representing the array of geographical adaptation across the species, named GsojaA to GsojaG, respectively, were selected to construct pan-genome of *G. soja* species (Supplementary Figs. 1 and 2 and Supplementary Table 1). Of these, six represent the six major distinct clusters of *G. soja* species (North, Huanghuai and South regions of China, and Japan, Korea and Russia) and 87.1% of gene diversity as revealed by analysis of 99 simple sequence repeats<sup>8</sup>. To maximally represent the species' wide geographic distribution, an additional accession was selected from the Northeast region (NER) of China, one of predicted domestication centers<sup>10</sup>. Four Chinese *G. soja* accessions were from the Chinese National Soybean GeneBank (CNSGB), and the other three were obtained from the USDA-ARS Soybean Germplasm Collection (University of Illinois, Urbana, IL), provided by Randall Nelson. Short-insert (180 bp and 500 bp) and 2 Kbp mate-pair genomic DNA libraries were constructed for each soybean sample. RNA was extracted from eight tissues, including young leaf, flower, young pod (1 cm), pod shell (5 mm), pod shell (2 cm), seed (3 mm, half full and full). RNAs were purified and RNAs from different tissues for each accession were mixed equally to construct sequencing libraries. The libraries were paired-end sequenced on the Illumina HiSeq 2000 platform. The following types of reads were filtered out from subsequent analyses:

- (a) Reads with  $\geq 10\%$  unidentified nucleotides (N);
- (b) Reads with  $> 10$  nt aligned to the adaptor, allowing  $\leq 10\%$  mismatches;
- (c) Reads with  $> 50\%$  bases having phred quality  $< 5$ ; and
- (d) Putative PCR duplicates generated by PCR amplification in the library construction process (i.e., read 1 and read 2 of two paired-end reads that were completely identical).

A total of 779.2 Gb DNA (average 111.9 $\times$  coverage per sample) and 35.47 Gb (average 5.07 Gb per sample) of RNA sequence data were obtained.

**De novo assembly.** First, we generated a 17-mer depth distribution of short-insert paired-end reads using Meryl<sup>50</sup> and applied GCE<sup>51</sup> to estimate the genome sizes of individual *G. soja* accessions. Reads were preprocessed by ALLPATHS-LG<sup>52</sup> error correction module to remove base calling errors. We also used ErrorCorrection in SOAPdenovo<sup>11</sup> package to connect 180-bp library pair end reads and to generate longer sequences for assembly. Reads of 180-bp and 500-bp library were used for contig building, and all pair-end reads libraries were used to provide links for scaffold construction. GapCloser (v1.12) from SOAPdenovo<sup>11</sup> package was used for gap filling within assembled scaffolds using all pair-end reads. Finally, scaffold sequences, which can be aligned to bacterial genomes with identity  $\geq 95\%$  and e-value  $\leq 1e-5$ , were filtered.

**Genomic alignment and short read mapping.** Assembled scaffolds for all *G. soja* accessions were aligned to the reference genome (GmaxW82) using the NUCmer program from the MUMmer package<sup>53</sup>. The parameters used for the genome alignment were “-maxmatch -c 90 -l 40”. The alignment results were further filtered to retain only one-to-one alignment regions using the “delta-filter” program incorporated in MUMmer package<sup>53</sup>. Short reads were mapped to both GmaxW82 and *G. soja* genomes using BWA<sup>54</sup>. After BWA alignment, putative PCR duplicates were filtered using the ‘rmdup’ utility of SAMtools<sup>55</sup>.

**Gene prediction and annotation.** Genome assemblies were scanned for putative gene coding regions using the Augustus<sup>56</sup> package. Then protein sequences from *Glycine max*, *Lotus japonicas*, *Medicago truncatula* and *Arabidopsis thaliana* were mapped to our assemblies using TblastN<sup>57</sup> (1e-5) and refined by GeneWise<sup>58</sup>. RNA-seq reads were mapped to *G. soja* assemblies using TopHat (v2.0.7)<sup>59</sup> with default parameters to identify exon regions and splice positions, and transcriptome-based gene structure predictions were made by Cufflinks (v2.0.2)<sup>60</sup> with default parameters. Results by *ab initio*, homology-based and transcriptome-based predictions were combined with the Evidence Modeler (EVM) package<sup>61</sup> (EVM set). A second GeneWise<sup>58</sup> prediction was done by using only proteins from *G. max* to identify the best

gene model. To exclude putative false positive predictions, protein sequences from the final gene models were aligned against protein annotations from 18 plant species using BLAST<sup>57</sup> (1e-5), including *Glycine max*, *Cajanus cajan*, *Phaseolus vulgaris*, *Lotus japonicas*, *Medicago truncatula*, *Populus trichocarpa*, *Ricinus communis*, *Cucumis sativus*, *Malus domestica*, *Arabidopsis thaliana*, *Brassica rapa*, *Carica papaya*, *Vitis vinifera*, *Solanum lycopersicum*, *Musa acuminata*, *Sorghum bicolor*, *Zea mays* and *Oryza sativa*. Gene models with  $< 50\%$  alignment coverage of their best homologs were filtered out. Gene models with  $> 20\%$  CDS region covered by repeat contents or corresponding protein sequence matched to TE-related domains by HMMPfam<sup>62</sup> were also filtered out. Finally, we checked gene models that were split into small fragments due to the discontinuity of scaffolds and revised our prediction of genes in each of the seven *G. soja* genomes. Putative biological functions of individual genes were assigned according to the best BLAST<sup>57</sup> hits in the *Arabidopsis thaliana* proteome, and by searching publicly available databases including Pfam<sup>62</sup>, PRINTS, PROSITE, ProDom and SMART with InterProScan<sup>63</sup>. Gene Ontology (GO)<sup>64</sup> terms for individual genes were retrieved from the corresponding InterPro descriptions. We also mapped these genes to the KEGG pathway<sup>65</sup> to identify their best matched categories.

**Gene clustering.** The core and dispensable gene sets were estimated based on gene family clustering. Based on the OrthoMCL<sup>66</sup> clustering results, we extracted gene families that were shared between *G. soja* samples, which were defined as core gene families. Gene families that are missed in one or more soybean samples were defined as dispensable gene families.

**SNP and indel identification.** Homozygous SNPs and small indels (less than 100 bp) were extracted from the one-to-one genomic alignment results using MUMmer<sup>53</sup>. We detected heterozygous SNPs using SAMtools<sup>55</sup> based on alignments of short reads onto assembled *G. soja* genomes. Then we located these heterozygous SNP sites on GmaxW82 genome according to the one-to-one genome alignment results. For indels larger than 100 bp, we searched for all alignment gaps ( $> 100$  bp) in both *G. soja* genomes and GmaxW82 genome. Indels identified as gaps surrounded by one-to-one alignments with at least 100 bp in both ends were retained. We used EMBOSS Water<sup>67</sup> based on the standard Smith-Waterman algorithm<sup>68</sup> to revise the retained indel set by redoing the alignments of regions that contain multiple indels. Putative functional effects of SNPs and indels were annotated using the ANNOVAR package<sup>69</sup>. SNPs/indels causing stop codon gain, stop codon loss and frameshift were defined as large-effect mutations. Gene Ontology category enrichments for genes containing large-effect mutations were found using the FUNC package<sup>70</sup>. To validate our results, we randomly selected 51 SNPs and 75 indel events, which is, 173 SNP sites and 20 indel sites in seven *G. soja* accessions, for PCR-Sanger sequencing using the ABI 3730XL. Additional 73 SNP events at 25 SNP sites were selected for mass-spectrometric assay (Sequenom). Among these sites, 81 SNP and 64 indel events were only detected with the current approach and could not be found via resequencing approach alone.

**CNV detection.** To avoid abnormal depths in non-coding regions owing to repeat elements, we focused on CNV in gene regions. Only CDS regions were sampled as background for CNV detection. The mean sequencing depth ( $d_0$ ) and s.d. ( $s_0$ ) were calculated for the background data set. Sequencing depth of each gene's CDS was extracted, and the mean value ( $d_g$ ) and s.d. ( $s_g$ ) were calculated. A standard *t*-test was applied to genes with  $d_g$  below  $0.2d_0$  or above  $1.8d_0$ , and those with *P*-value  $\leq 1e-5$  were defined as CNV. Genes with high similarity with GmaxW82 chloroplast and mitochondrion (50% length covered by e-value  $\leq 1e-5$  and identity  $\geq 95\%$  BLAST<sup>57</sup> hits) were not included.

**PAV detection.** For sequences that could not be aligned to GmaxW82, we used BLAST<sup>57</sup> to realign them to the GmaxW82 genome, all whole-genome sequencing trace reads and bacterial artificial clone sequences of *G. max* from GenBank, and filtered sequence stretches with an identity larger than 95%. *G. soja*-specific sequences were obtained after excluding potential bacterial contamination based on BLAST<sup>57</sup> to the NT database. After filtering for putative repeat elements (80% coverage) through RepeatMasker (<http://www.repeatmasker.org/>)<sup>71</sup> using an in-house repeat database<sup>72</sup>, genes with  $> 50\%$  CDS regions covered by *G. soja*-specific sequences were defined as

*G. soja*-specific genes. Based on the short reads alignment results, blocks with no mapped reads by *G. soja* were defined as GmaxW82-specific sequences. Regions with distance less than 500 bp were merged into one block. Genes that overlapped these blocks with 50% length were considered as GmaxW82-specific sequences.

***G. soja*-specific sequence validation.** *G. soja*-specific sequences were validated using multiplex PCR amplification. Each amplification included two pairs of primers, *G. soja*-specific sequence (Supplementary Table 12) and a control sequence shared by *G. soja* and *G. max*. For the control sequence (C1, 218 bp in GmaxW82), the forward and reverse primers were GTCATTGTAACAGGTGGGAGA and ACTGCGACTTTATTAAGATAG (C1). For each sequence, one or two (different locations) multiplexed PCRs were tested.

**Phylogenetic analysis.** Protein coding genes from *G. max*, *Cajanus cajan*, *Phaseolus vulgaris*, *Medicago truncatula*, *Populus trichocarpa*, *Cucumis sativus*, *Arabidopsis thaliana* and the seven sequenced *G. soja* accessions were used for gene family construction. For genes with alternative splicing isoforms, the longest isoform for each gene was used. Protein sequences for genes were compared by using all-by-all BLASTP<sup>57</sup> (1e-5), and then OrthoMCL<sup>66</sup> was used to cluster genes into orthologous gene families. From the clustering results, 670 orthologous gene families, with exactly one copy from each genome, were classified and defined as conserved single-copy gene families. Protein sequences from the 670 gene families were aligned by MUSCLE<sup>73</sup>. Fourfold degenerative sites from the CDS alignments were extracted and concatenated for phylogenetic analysis. Phylogenetic trees were built by the Neighbor-joining method incorporated in MEGA package<sup>74</sup>. The divergence time between legume genomes were estimated using the 'mcmctree' program incorporated in the PAML<sup>75</sup> package. We identified the SNPs in 57 published soybean accessions<sup>3,4</sup> together with our seven *G. soja* accessions using SAMtools<sup>55</sup> and constructed a separate phylogenetic tree for these 64 soybean accessions.

**Construction of the *G. soja* pan-genome.** First, we defined the core genome, which is shared by all seven *G. soja* accessions. Based on genomic alignment results, the aligned regions on GmaxW82 genome for each *G. soja* accession contributed to the first portion of the *G. soja* core genome. For sequences that could not be aligned to GmaxW82, Mugsy<sup>76</sup> was used to make multiple alignments using the default parameters. From the multiple alignments, we identified sequences that were shared in seven samples, which contributed to the second portion of the core genome. Sequences shared by less than seven samples, plus those presented in only one sample, were defined as the *G. soja* dispensable genome.

**Identification of positive selection and domestication related genes.** The syntenic orthologous gene families were used to estimate selection pressure by using the branch-site model incorporated in the PAML package. Based on a maximum likelihood ratio test (LRT), we identified genes under positive selection in GmaxW82 and in each *G. soja* lineage. These genes were identified as positively selected according to the chi-squared test ( $P < 0.01$ , FDR < 0.05,  $df = 1$ ), and containing amino acid sites that were selected with a Bayes probability higher than 95%.

**R-gene classification.** All genes from the *G. max* and *G. soja* samples were annotated using the HMM model against the Pfam-A database<sup>62</sup>. Genes with the Pfam domain of PF00931 were defined as R-genes, classified by different Pfam domains with E-value of 0.001. Domains of PF00931 (NB-ARC) and PF01582 (TIR) were defined as NBS and TIR domains, respectively, whereas domains with leucine-rich repeat, including PF00560 (LRR\_1),

PF07725 (LRR\_3), PF12799 (LRR\_4), PF13306 (LRR\_5), PF13516 (LRR\_6), PF13504 (LRR\_7) and PF13855 (LRR\_8), were defined as LRR domains. TIR-only and TIR-X sequences were not included in this study. For R-genes with CNV, the expression analysis result and related traits reported in previous studies<sup>25–28,77–79</sup> were listed in Supplementary Table 23.

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