Broad-spectrum fungal resistance in sorghum is conferred through the complex regulation of an immune receptor gene embedded in a natural antisense transcript

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Abstract
Sorghum (Sorghum bicolor), the fifth most widely grown cereal crop globally, provides food security for millions of people. Anthracnose caused by the fungus Colletotrichum sublineola is a major disease of sorghum worldwide. We discovered a major fungal resistance locus in sorghum composed of the nucleotide-binding leucine-rich repeat receptor gene ANTHRACNOSE RESISTANCE GENE1 (ARG1) that is completely nested in an intron of a cis-natural antisense transcript (NAT) gene designated CARRIER OF ARG1 (CARG). Susceptible genotypes express CARG and two alternatively spliced ARG1 transcripts encoding truncated proteins lacking the leucine-rich repeat domains. In resistant genotypes, elevated expression of an intact allele of ARG1, attributed to the loss of CARG transcription and the presence of miniature inverted-repeat transposable element sequences, resulted in broad-spectrum resistance to fungal pathogens with distinct virulence strategies. Increased ARG1 expression in resistant genotypes is also associated with higher histone H3K4 and H3K36 methylation. In susceptible genotypes, lower ARG1 expression is associated with reduced H3K4 and H3K36 methylation and increased expression of NATs of CARG. The repressive chromatin state associated with H3K9me2 is low in CARG-expressing genotypes within the CARG exon and higher in genotypes with low CARG expression. Thus, ARG1 is regulated by multiple mechanisms and confers broad-spectrum, strong resistance to fungal pathogens.
IN A NUTSHELM

Background: Sorghum, the fifth most widely grown cereal crop globally, provides food security for millions of people. The fungal disease sorghum anthracnose is the most devastating disease of this crop, with yield losses ranging from 50 to 86%. Genetic resistance has been successful in limiting losses to diseases in some crops. This approach has largely relied on leveraging the natural variation in the plant innate immune system, which is effective in restricting infection or inhibiting the progress of disease symptoms after infection. According to the current paradigm, the two primary branches of the plant immune system are pathogen-associated molecular pattern (PAMP)-triggered immunity (PTI) and effector-triggered immunity (ETI; Jones and Dangl, 2006). PTI is a form of disease resistance initiated upon the recognition of PAMPs by plasma membrane-anchored pattern recognition receptors (PRRs) located on the surface of plant cell membranes. Perception of PAMPs by PRRs activates a battery of immune responses, including the production of antimicrobial defense compounds and pathogenesis-related proteins and the accumulation of reactive oxygen species (Nicain et al., 2009). This form of resistance is usually quantitative and often independent of the strain of the pathogen (Boller and Felix, 2009).

Plant pathogens often suppress PTI by deploying specific virulence effectors that interfere with PTI. In response, plants have evolved ETI, which is based on the recognition of effectors by structurally related but polymorphic intracellular immune receptors known as nucleotide-binding site leucine-rich repeat (NLR) proteins. The core of NLRs is the nucleotide-binding (NB) and leucine-rich repeat (LRR) domains with N-terminal coiled-coil (CC) or Toll/interleukin-1 receptor domains (Belkhadir et al., 2004). ETI activates stronger plant immune responses that confer resistance to strains of the pathogen that express particular effectors. The strong selective pressure on pathogens imposed by ETI results in frequent defeat of resistance by pathogens that acquire or lose new virulence effectors and thus escape recognition by NLRs (Jones and Dangl, 2006). Recent observations suggest that PTI and ETI represent a continuum of overlapping responses rather than being categorically distinct (Thomma et al., 2011).

Natural antisense transcripts (NATs) are noncoding RNAs that regulate gene expression in animal and plant cells through several mechanisms (Pelechano and Steinmetz, 2013). Transcription interference between the expression of protein-coding sense transcripts and the corresponding NAT (Silverman et al., 1992; Shearwin et al., 2005), chromatin modification (Csorba et al., 2014), RNA interference, and DNA methylation have all been implicated in regulating the expression of the sense transcripts (Faghihi and Wahlestedt, 2009; Magistri et al., 2012; Rinn and Chang, 2012). In plants, NATs regulate phosphate homeostasis in rice (Oryza sativa; Jabnoune et al., 2013), cell wall biosynthesis in barley (Held et al., 2008), cytokinin regulation in petunia (Zukko and Meyer, 2007), salt tolerance and fertilization in Arabidopsis (Borsani et al., 2005), fiber development in cotton (Wan et al., 2016), and drought tolerance in maize (Xu et al., 2017). Sorghum (Sorghum bicolor) is an important food crop that also serves as a source of animal feed, biofuel, and other industrial products. Sorghum anthracnose, caused by the fungal pathogen Colletotrichum sublineola (Cs), is the most devastating foliar disease of this crop (Ali and Warren, 1992). Although anthracnose-resistant germplasm is available, the specific resistance regulators and their mechanisms of function have been unclear. In the current study, by...
screening a collection of sorghum natural variants, we found that the sorghum genotype SC283 displays a high level of broad-spectrum resistance to several different Cs strains, whereas the genotype TAM428 is susceptible to many different strains of the fungus. Recombinant inbred lines (RILs) generated by crossing SC283 with TAM428 displayed clear-cut resistance or susceptible disease responses similar to the parental lines. Whole-genome resequencing of DNA from resistant and susceptible RILs defined a major anthracnose resistance locus in SC283 that also confers resistance to other fungal pathogens. The resistance locus is composed of ANTHROCNOS RESISTANCE GENE 1 (ARG1) encoding a canonical NLR that is nested in an intron of a unique NAT designated CARRIER OF ARG1 (CARG). DNA- and RNA-seq analysis revealed that in resistant RILs, the loss of CARG transcripts and a miniature inverted-repeat transposable element (MITE) insertion in the ARG1 promoter region were associated with significantly enhanced expression of the full-length nested ARG1 gene. In contrast, susceptible RILs produced two ARG1 transcripts encoding truncated NLR proteins concomitant with an increase in NAT expression. The identity of the resistance gene and the relationship between the loss of CARG and enhancement of ARG1 expression were validated in distinct sorghum natural variants that carry independent resistant and susceptible allele of the CARG and ARG1 genes. In addition, histone H3K4 and H3K36 methylation at the region of overlap between CARG and ARG1 and in the ARG1 exon is enriched in resistant alleles but reduced in susceptible alleles. The repressive chromatin state associated with H3K9me2 is low within the CARG exon in CARG-expressing genotypes and higher in genotypes with low CARG expression. In summary, we discovered an immune receptor gene residing in an intron of a noncoding RNA gene that is regulated by MITE elements and confers complete and broad-spectrum fungal resistance.

**Results**

The sorghum line SC283 displays broad-spectrum resistance to sorghum anthracnose caused by Cs

We screened diverse sorghum natural variants collected from different regions of the world for resistance to the hemibiotrophic fungal pathogen Cs by inoculation with a high concentration of fungal spore suspension and incubation under conditions that favor disease in the greenhouse (Supplemental Data Set S1). The sorghum genotype SC283 was resistant to 11 different Cs isolates from the USA and Africa, suggesting broad-spectrum resistance (Figure 1; Supplemental Data Set S2). The inoculated SC283 leaves remained healthy and displayed resistance with hypersensitive response (HR) after inoculation with the Cs strain Csgl2 (Figure 1, A and B). In contrast, the widely known susceptible line TAM428 lacked any apparent resistance response and showed extensive disease lesions, massively chlorotic leaf areas, and complete tissue collapse (Figure 1, A and B). At 2 weeks after inoculation, SC283 remained healthy with no symptoms of infection, whereas TAM428 plants were killed by the fungus (Figure 1C). Microscopic analysis of inoculated tissue after trypan blue staining revealed restricted fungal growth in SC283 but extensive growth in TAM428 (Figure 1D). Interestingly, SC283 also manifested enhanced resistance to pathogens in a field experiment in Western Ethiopia, where anthracnose is the most prevalent disease. Figure 1E presents fungal resistance of SC283 via natural infection at a specific location.

**Identification of fungal resistance locus through whole-genome resequencing**

RILs generated by crossing SC283 and TAM428 were used to identify the resistance locus in SC283 using an approach that combined bulked segregant analysis (BSA), whole-genome sequencing, and genetic mapping. We tested the responses of 209 RILs after Cs inoculation in the greenhouse, which identified 109 resistant and 100 susceptible RILs; these results are consistent with the expected 1:1 segregation ratio ($\chi^2 = 0.387$, $P > 0.05$; Supplemental Data Set S3). The responses to the fungus in these resistant and susceptible RILs were similar to those of the parental lines SC283 or TAM428. We selected 50 resistant and 50 susceptible individual plants based on at least six rounds of independent disease assays (Supplemental Data Set S3). A pair of DNA bulks was constructed by pooling DNA from the 50 resistant and 50 susceptible RILs that were then sequenced using Illumina HiSeq 2500. More than one billion paired-end reads were obtained, including 494 million resistant bulk (RB) reads and 513 million susceptible bulk (SB) reads (Supplemental Data Set S4). These paired-end short reads covered the sorghum genome at an average depth of 66× and 68× in the RB and SB pools, respectively. In parallel, reference sequences were built by sequencing eight sorghum cultivars, including the two parental lines of the RILs used in this study (Supplemental Data Set S4).

To determine the genomic region associated with resistance, we conducted BSA in the quantitative trait locus (QTL)-seq pipeline (Takagi et al., 2013), an approach combining BSA with whole genome resequencing that is often used to identify genes underlying both qualitative traits and QTLs. QTL-seq relies on an estimation of the single-nucleotide polymorphism (SNP) index in the RB and SB sequences in order to identify genomic region harboring the major QTL. More than 3 million SNPs were identified based on mapped reads for QTL analysis; these SNPs were unevenly distributed in the genome. We determined the SNP-index of each SNP using the QTL-seq pipeline (Supplemental Figure S1; Takagi et al., 2013) and calculated the $\Delta$(SNP-index) by subtracting the SNP-index of SB from that of RB (Supplemental Figure S1). As expected, the $\Delta$(SNP-index) was zero in most genomic regions, but a few regions exhibited positive or negative values, indicating differences from the sorghum BTx623 reference genome (Paterson et al., 2009; Supplemental Figure S1E). A $\Delta$(SNP-index) higher than 0.44 was observed in the region from 7.15 to 15.80 Mb on chromosome 7 with $P < 0.05$ under the null hypothesis.
Figure 1 Disease responses of sorghum lines SC283 and TAM428 to Cs and identification of the resistance locus. A, C, Disease response phenotypes at 7-day post inoculation (dpi) (A), and 14 dpi (C). B, Disease symptoms on infected leaves at 10 dpi. (D) Trypan blue staining of Cs inoculated leaves showing fungal structures on TAM428 but a lack of fungal growth in SC283. Inoculated leaf tissues were stained with trypan blue, and samples were examined under a microscope to visualize fungal material. Scale bars represent 20 μm. E, Resistance of SC283 to foliar pathogens in the field under natural infestation. The experiments were repeated at least three times with similar results. In all the disease response data shown in (A)–(D), the Cs strain Csgl2 was used. F, Identification of an anthracnose resistance QTL in SC283 through QTL-seq analysis of recombinant inbred lines. SNP-index plots of R-bulk (top) and S-bulk (middle), and Δ (SNP-index) plot (bottom) of chromosome 7 with statistical confidence intervals under the null hypothesis of no QTLs (green, P < 0.05; yellow, P < 0.01). The Δ (SNP-index) plot was obtained by subtracting the of S-bulk SNP-index from the R-bulk SNP-index for the RILs. Plants were scored as resistant or susceptible based on their disease symptom or resistance response phenotypes. The DNAs from resistant or susceptible plants were bulked to make separate resistant (R) and susceptible (S) DNA bulks. S-bulk, DNA from the susceptible plants, R-bulk, DNA from the resistant plants. SNP index and Δ (SNP-index) was determined as described.
This contrasting pattern of the SNP-index for RB and SB defined a major Cs resistance locus within the 7.15–15.80 Mb genomic region on chromosome 7 (Figures 1, F and 2, A; Supplemental Figure S1E).

Identification of candidate resistance gene(s) in the mapped anthracnose resistance locus
To further narrow the genomic region carrying the resistance locus, we developed 13 molecular markers that were polymorphic between the parental lines spanning the 6.2–18 Mb genomic interval that was defined to contain the anthracnose resistance locus. Phenotypic analysis identified recombination events between markers, which further narrowed the candidate genomic region (Figure 2, A and B; Supplemental Data Set S5). Based on these results, the anthracnose resistance locus was mapped to 7.80-kb genomic region flanked by InDel-16 and InDel-6 markers that showed a complete co-segregation with the disease phenotype (Figure 2B).

Next, to identify the specific Cs resistance gene, we annotated SNPs, insertions, and deletions in the ~1 Mb mapping interval containing 29 genes (see “Materials and methods”) after filtering low-quality sequences and SNPs with no polymorphisms in the parental lines (Cingolani et al., 2012). In all, sequence variants in 15 genes were closely analyzed, and most predicted genes were excluded based on a lack of significant polymorphisms (Supplemental Data Set S6). Importantly, sequence polymorphisms that have potential effects on gene function or the integrity of encoded proteins were mapped to two candidate genes: Sobic.007G085400 and Sobic.007G085350 (Figure 2C; Supplemental Data Set S6). The susceptible TAM428 genotype contained a sequence polymorphism that introduced a premature stop codon in the Sobic.007G085400 coding sequence. However, in the resistant SC283 genotype, two SNPs replaced the stop codon with a serine codon in this gene, restoring the open reading frame (ORF; Supplemental Data Set S6).

Sobic.007G085400 encodes a canonical NLR with N-terminal CC domains, NB site, and C-terminal LRR domains (Supplemental Data Set S7), making it an excellent candidate, and is hence designated ANTHRACNOSE RESISTANCE GENE1 (ARG1). This class of proteins function as intracellular receptors for effector proteins and are key determinants of ETI (Jones and Dangl, 2006). The sequence polymorphism in ARG1 is consistent between the bulks and parental lines (Figure 2D). The susceptible lines and SB carry the stop codon and the resistant lines and RB harbor the intact ARG1 ORF (Figure 2E). Interestingly, Sobic.007G085400 (ARG1) is nested in the intron of the second candidate gene, Sobic.007G085350 (Figure 2D). In addition, an 8-bp sequence deletion (GGCGACCT) in the first exon of Sobic.007G085350 at position 10,793,252 on chromosome 7 was identified in the resistant parent SC283 that was not present in the susceptible parent TAM428 (Figure 2E; Supplemental Data Set S6). The deletion in Sobic.007G085350 in SC283 was also present in RB but absent in the SB sequence (Figure 2E). Based on these genetic data, the polymorphism at Sobic.007G085350 is also considered to be the candidate causal sequence change that co-segregated with the resistance phenotype in SC283. The likely noncoding corresponding gene (Sobic.007G085350) is hereafter designated as CARRIER OF ANTHRACNOSE RESISTANCE GENE (CARG).

The ARG1 gene embedded in the NAT gene encodes an NLR receptor
Analyses of the genomic organization of the CARG–ARG1 locus revealed that CARG has two exons, interrupted by two introns, the second of which is quite large (Figure 3A). The ARG1 coding region is embedded in this large second intron. To delineate the boundaries of the CARG–ARG1 genomic and transcript sequences, we conducted 5’- and 3’-rapid amplification of complementary DNA ends (RACE). The 5’-untranslated regions (UTRs) of the CARG gene are 148 bp in TAM428 and 139 bp in SC283, and the DNA sequences in these UTRs are identical except for a 9-bp size difference due to an Insertion/Deletion (Indel). SC283 carries a shorter CARG gene 3’-UTR (867 bp) than that of TAM428 (1,254 bp; Figure 3A; Supplemental File S1). Both TAM428 and SC283 carry 740-bp ARG1 5’-UTRs with very high sequence similarity. However, in SC283, the 5’-UTR carries a 423-bp intron from positions −662 to −1,084 relative to the ARG1 start codon and a second, 33-bp intron between positions −288 to −320, which are lacking in TAM428. TAM428 and SC283 both have 151-bp ARG1 gene 3’-UTRs (Figure 3A; Supplemental File S1). To confirm that the UTR sequences were not artifacts from cDNA synthesis, we performed reverse transcription polymerase chain reaction (RT-PCR) using one primer in the UTRs and one in the coding regions of ARG1 or CARG. The PCR products were cloned and at least three individual clones were sequenced, revealing that the 5’-UTR of ARG1 and the 3’-UTR of CARG partially overlap in both genotypes (Figure 3A; Supplemental File S1).

Interestingly, the CARG–ARG1 genomic region was found to harbor MITE-related sequences that showed significant differences between the resistant and susceptible genotypes (Figure 3A; Supplemental File S1 and Supplemental Figure S2). The susceptible genotypes carry MITEs of 275 bp in the 5’-UTR, 151 bp in the second intron, and 248 bp in the 3’-UTR of CARG. The resistant lines carry the same 151-bp MITE in the second intron of CARG, as well as a different 420-bp MITE insertion in the 3’-UTR in this gene, which is also located in the first intron of ARG1 in this allele. Sequence analysis of the 5’-UTR of this allele suggested that the MITE insertion introduced a splice junction that results in splicing of the MITE from the transcript. The locations of the MITEs relative to the CARG–ARG1 genes are shown in Figure 3A. The 5’- and 3’-MITE sequences flanking the CARG–ARG1 genes show very limited sequence identity with each other, and the MITEs in the 3’-UTR of the CARG in the two alleles were also inserted into different positions.
Figure 2  Fine mapping and sequence comparisons in mapped genomic region identifies ANTHRACNOSE RESISTANCE GENE1. A, Genomic region harboring the anthracnose resistance locus on Chromosome 7 (7.15–15.80 Mb). B, Fine mapping of the anthracnose resistance locus based on analyses of molecular markers shown on the physical map. The anthracnose resistance locus was fine mapped to a ~780-kb genomic region flanked by InDel-16 and InDel-6 markers, which showed no recombination. InDel-16, InDel-4, and InDel-6 markers co-segregated with disease responses in the RILs. The numbers in brackets indicate the number of recombinants recovered. C, Gene models in the mapping region flanked by InDel-15 and InDel-17 markers (10,009–11,029 kb). The two candidate genes are shown in magenta. D, Comparison of DNA sequence reads of the candidate genes Sobic.007G085350 and Sobic.007G085400 from RB and SB and parental lines SC283 and TAM428 using Integrative Genomics Viewer (IGV). The DNA sequences were compared with the reference genome BTx623, and SNPs were marked as color bars. The sequences from the SB and TAM428 show no sequence polymorphism relative to the reference genome in the mapping region. The sequence of RB is identical to that of SC283, and these sequences are different from those of SB and TAM428. The genomic structure of the resistance locus showing Sobic.007G085350 (CARG) and Sobic.007G085400 (ARG1), which is nested in Sobic.007G085300. The coding and genomic regions with introns are based on predictions in the database (Phytozome V10, Sbicolor 313 V3.1), which was confirmed by our sequencing. Black bars indicate genomic region, blue bars indicate coding sequence, thin blue bars indicate UTRs, lines indicate introns, and arrows indicate the direction of transcription. E, Sequence variants in the Sobic.007G085350 (CARG) and Sobic.007G085400 (ARG1) genes between susceptible and resistant genotypes. The Sobic.007G085350 and Sobic.007G085400 genomic regions were amplified from different genotypes and sequenced to determine sequence polymorphisms and to identify additional alleles. The lower shows a sequence comparison between susceptible and resistant lines in Sobic.007G085400. Lower left, a 2-bp (AG) deletion at 10,797,728 was observed only in the susceptible Ai4 and SQR genotypes, which created a frameshift. Lower right, two SNPs at 10,797,005 and 10,797,006 in resistant lines create synonymous mutations that correct a stop codon in the Sobic.007G085400 gene that is in all the sensitive lines except for Ai4 and SQR. The codons and the corresponding amino acids are shown for resistant and susceptible lines. Dash indicates the deletion unique to Ai4 and SQR.
Figure 3 Genomic structure and RNA-seq analysis of the CARG and ARG1 genes and upstream regulatory elements. A, Genomic structure of the CARG–ARG1 locus. The lower panel shows RNA-seq analysis of the CARG–ARG1 locus. RNA-seq was conducted at 0, 24, and 48 h after inoculation with Cs. Sequence reads were viewed by IGV and mapped to the TAM428 and SC283 genomic sequences of CARG and ARG1. The transcript count data are shown at 0–200 scale for SC283 and 0–20 scale for TAM428 due to the elevated levels of the transcripts for ARG1 in SC283. The mapped transcripts, 5′- and 3′-RACE, qRT-PCR, and genomic sequence data were used to determine the exon, introns, and UTRs of CARG and ARG1. B, Expression of CARG and ARG1 in SC283 and TAM428 based on RNA-seq transcript count data. Error bars indicate the standard deviation of three libraries (error bars ± SD [n = 3]). Different letters indicate significant difference based on the least significant difference (LSD, P < 0.05). C ARG1 and D CARG expression in different sorghum lines. In (C) and (D), expression levels were analyzed by qRT-PCR in independent sorghum genotypes at 0 and 2 days after Cs inoculation. Data were normalized by the comparative cycle threshold method with Actin as the internal control and presented as relative expression. The data represent mean ± SD (n = 3 biological replicates per genotype, collected from one independent experiment). Similar results were obtained in two independent experiments.
and thus likely represent two independent insertion events (Supplemental Figure S2).

We mapped RNA-seq data that we generated from healthy and pathogen-inoculated TAM428 and SC283 lines to genomic sequences of the corresponding lines to determine the transcript boundaries of CARG and ARG1. The ARG1 transcripts were detected in both TAM428 and SC283, but the CARG transcript was observed only in TAM428 (Figure 3A). The RNA-seq further revealed that the basal expression level of ARG1 was significantly higher in SC283, with further increased after Cs inoculation, while CARG expression was significantly lower both before and after infection (Figure 3A). Significantly different transcript read counts were observed for CARG and ARG1 between TAM428 and SC283 (Figure 3B). In contrast, TAM428 exhibited higher CARG expression and much lower ARG1 expression than SC283 (Figure 3B). The ratio of expression of ARG1 and CARG was only two-fold different in the susceptible TAM428 line, compared to at least a 250-fold difference in SC283 based on RNA-seq data (Figure 3B), further supporting the hypothesis that the loss of CARG transcript in SC283 is correlated with the enhanced expression of ARG1.

The gene expression pattern observed from the RNA-seq data was confirmed using quantitative RT-PCR (qRT-PCR) with primers flanking introns in both the CARG and ARG1 genes. The expression level of ARG1 was significantly higher in six resistant genotypes and three resistant RILs carrying the CARG deletions than in lines where CARG is normally expressed (Figure 3C; Supplemental Figure S3A). qRT-PCR using primers flanking the second CARG intron confirmed CARG expression in the susceptible genotypes and the susceptible RILs SSD50, SSD61, and SSD65, all of which exhibited significantly higher levels of CARG expression than in the resistant genotypes (Figure 3D; Supplemental Figure S3B). A second primer pair flanking the first intron of CARG gave similar results (Supplemental Figure S3C). In every case, alleles that abrogate CARG gene expression were associated with both high levels of ARG1 expression and resistance to anthracnose. Both ARG1 and CARG were induced by fungal infection, with only ARG1 showing a significantly larger induction in both genotypes.

Validation of ARG1 through characterization of independent alleles

To provide further genetic evidence for a link between the observed phenotypes and sequence variation in the candidate genes, we searched whole genome sequences of several sorghum genotypes to identify additional alleles of the CARG and ARG1 genes. Sorghum lines carrying independent deletions and/or SNPs in the CARG and ARG1 genes were identified from analysis of 46 deep sequenced cultivars and land race genotypes available in public databases and inhouse generated sequences that have been tested for anthracnose disease resistance (Supplemental Data Set S1). Among these, five resistant sorghum lines, SC35C, BTx378, KS115, PI585749, and PI586439, carried the same 8-bp deletion in the CARG gene and an intact ARG1 gene, as was observed in SC283. The resistant lines also carried additional sequence alterations in the CARG gene that are distinct from those of SC283, providing additional genetic evidence for variation in these genetic backgrounds (Figure 2E; Supplemental Figure S4). On the other hand, in all 40 susceptible lines examined, mutations that disrupt ARG1 were linked to intact CARG genes. These susceptible lines each carried one of two distinct susceptible ARG1 alleles, one with a premature stop codon identical to TAM428 at position 508 amino acid (aa) or a second, independent ARG1 allele that introduced a stop codon at position 275 aa in Ai4 and SQR (Figure 2E; Supplemental Figure S4 and Supplemental Data Set S1). These ARG1 sequence variations between the susceptible and resistant genotypes are linked to polymorphisms observed in the 5’-upstream region and within the CARG gene sequences in all the genotypes studied. The tight linkage between two independently derived mutations that result in stop codons that are both associated with susceptibility provides strong evidence that a functional ARG1 gene is required for resistance.

The sequence variations among resistant and susceptible RILs and other independent sorghum genotypes were confirmed using molecular markers (Figure 4, A–C). The InDel-4 marker flanking the sequence deletion in the CARG (Sobic.007G085350) exon and InDel-5 within the CARG intron co-segregated with the resistance phenotype, which confirmed that the polymorphism in CARG is tightly linked to resistance on the same region of chromosome 7 (Figure 4, A–C). InDel-6, located in the ARG1 promoter region, also co-segregated with the resistant and susceptible phenotypes. Resistance was invariably observed in RILs and five other genotypes with different origins that carried intact ARG1 and the linked molecular markers (Figure 4, C–E).

The genotypes that carried the ARG1 mutant allele and the associated polymorphisms showed typical disease symptoms, such as chlorotic and necrotic lesions and black spots caused by fungal acervuli (fungal reproductive structures) to a greater or lesser degree (Figure 4, D and E; Supplemental Figure S5). The susceptible genotypes TAM428, BTx623, Tetron, SQR, PQ434, KP33-2, ZZL, IS9830, and Ai4 showed more severe and advanced disease symptoms, while 555 and SRN39 showed mild disease symptoms. These results are consistent with the presence of additional modifiers of disease resistance in these lines (Figure 4E; Supplemental Figure S5).

To determine the relationship between disease symptoms and fungal growth, we quantified fungal growth in inoculated plants using quantitative PCR (qPCR) amplification of the internal transcribed spacer (ITS) region of the fungal ribosomal DNA (rDNA). Overall, the fungal growth correlated well with disease symptoms (Figure 4F). These analyses confirmed the sequence data, and the disease responses were consistent in all genotypes tested. Thus, among the genes that map to the QTL region, only the CARG–ARG1 gene pair showed consistent sequence polymorphism between the two parental lines and between resistant and susceptible
Figure 4 Co-segregation of molecular markers, and pathogen responses of recombinant inbred lines and distinct sorghum lines. A–C, Genotyping of the resistant and susceptible recombinant inbred lines (A and B) and different sorghum lines with distinct origins (C) using Indel molecular markers in Sobic.007G085350 and Sobic.007G085400. InDel4, InDel5, and InDel6 represent three deletion polymorphisms, which were used to design PCR markers in the Sobic.007G085350 and Sobic.007G085400 genes and are polymorphic between resistant lines and susceptible lines. R RILs, resistant recombinant inbred lines; S RILs, susceptible recombinant inbred lines; SSD, single seed decent. D, E, Disease response phenotypes of sorghum lines carrying different \textit{CARG} and \textit{ARG1} alleles after inoculation with \textit{Cs} strain Csg12. SSD1, SSD3, and SSD4 are resistant RILs, while SSD50, SSD61, and SSD65 are susceptible RILs. SC283, BTx378, KS115, SC35, PIS85749, and PIS86439 are resistant genotypes that carry the \textit{CARG} deletion and intact \textit{ARG1}. TAM428, BTx623, Tetron, SQR, PQ434, KP33-2, ZZZ, IS9830, 555, and SRN39 are susceptible genotypes that have intact \textit{CARG} gene and mutated \textit{arg1}. F, Quantification of fungal growth. Fungal growth in infected leaves was determined by qPCR amplification of the \textit{Cs Internal Transcribed Spacer rDNA (Cs ITS)}. Relative DNA levels were calculated using \textit{SbActin (Sb Act)} as a reference gene. Data represent mean ± SD from three technical replicates. Different letters indicate statistically significant differences ($P < 0.05$, Student’s $t$ test). Similar results were obtained in three different independent experiments.
of the allele linked to Cs is dominant for disease resistance. Thus, recarg–ARG1 from a 3 resistant: 1 susceptible segregation ratio (segregation in the F2 population do not differ significantly with resistance. The values obtained from the analysis of resistant, with the examined, 114 individuals were susceptible and 295 were resistant, with the 409 F2 single plants tested were resistant. Of the 409 F2 and selfed progenies from the TAM428 x SC283 cross. All 10 F1 plants expressed both CARG and ARG1 alleles. We evaluated plants carrying the homozygous CARG deletion (carg/carg;ARG1/ARG1), CARG homozygous wild-type (CARG/CARG;arg1/arg1), and heterozygous plants (CARG/carg;ARG1/arg1) for Cs resistance by assessing disease symptoms and fungal growth (Figure 5, A–C). Both TAM428 and F2 CARG/CARG;arg1/arg1 plants displayed disease symptoms, including microscopic dark spots indicative of fungal acervuli and chlorotic leaves, which were quantified by measuring the area of the disease lesion relative to the total leaf area (Figure 5, A and B).

We also quantified fungal growth based on qPCR amplification of the ITS region of the fungal rDNA. The F2 carg/carg;ARG1/ARG1 and carg/carg;ARG1/arg1 plants were equally resistant based on both fungal growth and quantification of disease symptoms and shared comparable levels of resistance with SC283 plants (carg/carg;ARG1/ARG1; Figure 5, B and C). The F2 CARG/CARG;arg1/arg1 plants were significantly more susceptible than the F2 carg/carg;ARG1/ARG1 and carg/carg;ARG1/arg1 plants, but clearly less susceptible than the TAM428 plants carrying the same CARG/CARG;arg1/arg1 alleles (Figure 5, A–C). Similar differences were observed in disease symptoms and fungal growth when the various genotypes were drop inoculated in detached leaf assays (Supplemental Figure S6). These results suggest the presence of other factors in the SC283 background that modulate resistance. Overall, however, resistance was associated with the presence of ARG1 and susceptibility was attributed to the loss of the wild-type ARG1 allele in all genotypes evaluated. The above data show that the CARG–ARG1 locus determines resistance in SC283 and likely in other resistant sorghum genotypes as well.

We conducted gene expression analyses in the F2 plants. Of these plants, carg/carg;ARG1/ARG1 and carg/carg;ARG1/arg1 plants showed comparable levels of resistance, having only the HR, which is consistent with their similar levels of ARG1 expression (Figure 5, A and D). In all of these genotypes, ARG1 gene expression levels were negatively correlated with CARG expression (Figure 5, D–F). In CARG/carg;ARG1/arg1 plants, the level of ARG1 expression was comparable to that in carg/carg;ARG1/ARG1 plants, despite an intermediate level of CARG transcript. These results suggest that the expression of only one copy of CARG is not sufficient to affect overall ARG1 transcript levels. However, the inverse correlation between CARG and ARG1 expression suggests that enhanced resistance may partially result from a loss of cis-NAT CARG transcript, perhaps permitting a concomitant increase in expression of an intact ARG1 allele in the resistant genotypes.

The ARG1 allele in susceptible genotypes expresses alternatively spliced transcripts encoding truncated NLRs

To further confirm the ARG1 expression patterns in response to Cs infection identified by RNA-seq, we performed RT-PCR analysis using primers permitting amplification of full-length ARG1. The transcript levels of ARG1 in SC283 and TAM428 appeared to display good correlation with the RNA-seq data. A single pathogen-inducible ARG1 transcript was observed in SC283, which harbored CARG with an 8-bp deletion and lacked the MITE in the 5′-region of this transcript. However, TAM428, which expresses the CARG NAT, produced two variant ARG1 transcripts, both of which were pathogen inducible (Figure 6A). We sequenced all ARG1 transcripts from SC283 and TAM428 to determine the nature of the splice variants of the ARG1 transcript. Interestingly, the larger variant was comparable in size to the ARG1 transcript in the resistant genotypes but carried a stop codon in the middle of the ARG1 gene, as shown in Figure 2E. The second transcript was much smaller, skipped the LRR domain, and retained only the CC and NBs-ARC domains (Figure 6B; Supplemental Figure S7). We analyzed alternative splicing of ARG1 across all genotypes tested by our disease assay to determine whether this occurs in other susceptible genotypes. Two different ARG1 transcripts were observed in the 10 susceptible lines tested, while all six resistant lines produced a single ARG1 transcript (Figure 6C).

The CARG 3′-UTR in the susceptible genotype produces small RNAs

The CARG–ARG1 locus has an interesting genomic structure. The entire coding sequence of ARG1 is embedded in an intron of CARG. The 5′-UTR of ARG1 overlaps with the 3′-UTR of CARG (Figure 3A; Supplemental Figure S8A), raising the possibility of an interaction between the two complementary transcripts. Due to the overlapping CARG and ARG1 transcripts, there is a potential for formation of double-stranded RNA (dsRNA) and small RNAs (sRNA). We therefore conducted small RNA profiling of healthy and infected SC283 and TAM428 to identify sRNA sequences
that map to the CARG–ARG1 region that may regulate gene expression. A comparison of sRNAs in the two genotypes identified a cluster of sRNAs from a portion of the 3′-UTR of CARG of TAM428 but not of SC283. These sRNAs correspond to a MITE that is present in TAM428 but is missing from SC283, which has a different MITE that is spliced out of the SC283 ARG1 5′-UTR (Supplemental Figure S8A). Given that the MITE in TAM428 is present in very high copy numbers in the sorghum genome (Supplemental Figure S2), it is possible that these small RNAs are derived from transcripts that include this MITE elsewhere in the genome. Interestingly, however, the sRNAs were present in...
TAM428, but there was no sRNA accumulation in SC283, suggesting that at least some of these sRNAs are correlated with **CARG** expression (Supplemental Figure S8B). Interestingly, this MITE shows similarity to a hairpin variant of the MITE that expresses a putative pre-miRNA that is processed into sbi-miR6225 (miRBase, Version 21), which is similar to the small RNAs present at **CARG** (Supplemental Figure S9). The significance of these sRNAs needs to be determined in future studies.

### MITE sequences regulate **ARG1** gene expression

We studied the association between the MITE insertions and the expression of the **CARG** and **ARG1** genes via qRT-PCR (Figure 7A). A higher level of **CARG** expression was observed in the susceptible lines harboring a 275-bp MITE insertion immediately upstream of the **CARG** transcript than in the resistant lines lacking this MITE, suggesting that this MITE may be driving **CARG** expression. Similarly, in lines that carry the 420-bp MITE in the 5'-UTR intron of **ARG1**, higher expression of this gene was observed. In contrast, the 248-bp MITE insertion in the 3'-UTR of **CARG** found in the susceptible genotypes did not correlate with any significant induction of **ARG1** expression (Figure 7A). These results suggest that the 420-bp MITE in the 5'-UTR of **ARG1** positively regulates **ARG1** expression in the resistant genotypes, whereas the 248-bp MITE negatively regulates **ARG1** expression in the susceptible genotypes, perhaps due to induction of expression of the **CARG** NAT.

To further evaluate the roles of MITEs and the **CARG** NAT in regulating gene expression, we generated a series of constructs derived from the **CARG**–**ARG1** locus with swapped sequences between SC283 and TAM428 (Figure 7B). First, the **ARG1** promoter region in TAM428 was replaced with that from SC283 and vice versa (S1 and S2). Second, the **ARG1** coding region in TAM428 was replaced with the **ARG1** from SC283 (S3). Third, the **CARG** promoter region in SC283 was replaced with the **CARG** promoter from TAM428 (S4). The resulting constructs were individually transformed into Arabidopsis in parallel with constructs representing the native **CARG**–**ARG1** genes in SC283 and TAM428. We selected single copy transgenic plants for all six Arabidopsis lines and measured **CARG** and **ARG1** expression using qRT-PCR (Figure 7B). The expression of each gene reflected the promoter associated with it in each construct. Thus, constructs S1, S3, and S4, which all...
had the native CARG promoter from TAM428 driving CARG expression, had levels of expression of this gene similar to that of the native TAM428 construct. Similarly, S1 and S4, which had the native promoter of ARG1 from SC283, showed similar levels of expression of this gene to that of the SC283 native construct (Figure 7B). Thus, each promoter appears to function largely independently. However, ARG1 revealed contrasting expression tendencies to CARG in S2 and S3 (Figure 7B), suggesting that at the relatively low levels of expression of ARG1 driven by the TAM428 promoter, CARG expression can in fact repress ARG1 expression.

**Figure 7** Regulation of CARG and ARG1 genes by MITE and 5′-UTR intronic sequences. A, Genomic distribution of MITE insertions in the CARG–ARG1 locus (middle). The ARG1 gene is in the second intron of CARG. Black boxes denote the exon of ARG1, and gray boxes denote exons of CARG. The two genes are in the opposite orientation. The promoter regions (upstream from the ATG) are represented as a thick horizontal line; the yellow and orange vertical lines are indel; the gold, violet, blue, and magenta boxes indicate 275-, 151-, 248-, and 420-bp MITE insertions, respectively. The stop codon in the middle of the yellow and orange vertical lines are indel; the gold, violet, blue, and magenta boxes indicate 275-, 151-, 248-, and 420-bp MITE insertions, respectively. T35S: CaMV 35S terminator. Each construct was co-transfected with a CaMV 35S:luciferase construct into Arabidopsis protoplasts. GUS activities with or without chitin treatment in Arabidopsis protoplasts. White and gray arrows indicate the promoter of ARG1 and CARG, respectively. Chitin induces ARG1 expression in the resistant genotype. Sorghum plants were treated with 2-nM chitin (P < 0.05, Student’s t test). C, ARG1 MITE from the SC283 allele regulates basal and chitin-induced ARG1 expression. Schematics of the ARG1 and CARG promoter GUS reporter construct on the left and their GUS activities with or without chitin treatment in Arabidopsis protoplasts. White and gray arrows indicate the promoter of ARG1 and CARG, respectively. T35S: CaMV 35S terminator. Each construct was co-transfected with a CaMV 35S:luciferase construct into Arabidopsis protoplasts. Transfection efficiency was normalized relative to the expression of the Luciferase gene. Data are mean ± SD from triplicate experiments. Different letters indicate statistically significant differences (P < 0.05).
The exogenous application of chitin and the 5'-UTR intron regulate ARG1 expression

We tested the ability of the promoter sequences of ARG1 from TAM428 and SC283 carrying distinct MITEs to regulate gene expression. The CARG and ARG1 promoters were cloned into β-glucuronidase (GUS) reporter constructs, and each construct was transfected together with a CaMV 35S:luciferase (LUC) construct into Arabidopsis protoplasts. GUS activity was normalized to the fluorescence of luciferase, which was used to determine the relative transcriptional efficiency. The ARG1 promoter from SC283 (ARG1-P-SC:GUS) significantly increased GUS activity, but the ARG1 promoter from TAM428 (ARG1-P-TAM:GUS) yielded lower GUS activity (Figure 7C). A lower level of GUS activity was generally observed using the CARG promoters, but GUS activity was higher using the TAM428 construct compared to the SC283 construct. Interestingly, ARG1-P-SC:GUS activity increased in response to chitin (β-1,4 linked N-acetylgulcosamine) treatment relative to the mock-treated samples, while ARG1-P-TAM:GUS activity showed no significant change in response to chitin treatment (Figure 7C). The CARG promoter GUS constructs showed no altered response to chitin treatment in all assays tested.

Chitin fragments are PAMPs generated by plant chitinase activity during fungal infection that are subsequently perceived by plant cells to activate immune responses (Gong et al., 2020). Cs significantly induced ARG1 expression in SC283, which may partially be attributed to recognition of chitin fragments (Figure 3C). To confirm the responses to chitin observed in protoplasts, we treated detached TAM428 and SC283 leaves with 2-nM chitin and analyzed chitin observed in protoplasts, we treated detached谭428 and SC283 leaves with 2-nM chitin and analyzed chitin fragments (Figure 3C). To confirm the responses to chitin treatment in all assays tested.

The resistant allele of ARG1 contains two introns in the 5'-UTR that are missing in ARG1 from the susceptible allele. One of the two introns in SC283 ARG1 is likely spliced due to the presence of its unique MITE (Supplemental Figure S2). To examine the roles of these introns in regulating gene expression, we generated constructs with and without these sequences (Figure 7E). Three constructs with various deletions of the introns in the 5'-UTR of ARG1 from SC283 were fused to the GUS reporter gene in the pSAT1 vector. SC283-Del-1 contains the native ARG1 promoter and the 5'-UTR but lacks the 423-bp intron that includes the MITE element. SC283-Del-2 carries the native promoter and 5'-UTR but lacks the 33-bp intron, and SC283-Del-3 contains the native promoter and the 5'-UTR but lacks all introns. We transfected Arabidopsis protoplasts with these constructs, with the CaMV 35S:LUC construct used as an internal control. Protoplasts harboring all of the constructs showed a significant increase in GUS activity compared to the negative control, i.e., the pSAT1 vector, which contains the Nopaline synthase (Nos) promoter without the GUS reporter gene. SC283-Del-1 and SC283-Del-3 exhibited lower GUS expression compared to SC283, whereas GUS expression in SC283-Del-2 was similar to that of SC283. These results indicate that the 423-bp intron including the MITE in the 5'-UTR functioned as transcriptional activator in the protoplast transient expression assays (Figure 7E).

Permissive chromatin at the ARG1 locus is correlated with fungal resistance

The higher expression of ARG1 in resistant genotypes is linked to a loss of the CARG NAT. This raised the possibility that ARG1 is regulated by CARG noncoding RNA, which could result in DNA methylation, histone methylation, or transcriptional interference due to the opposite orientation of the two transcripts. We first examined DNA methylation status in the coding regions of ARG1 using bisulfite sequencing analyses. In general, there was no significant difference in DNA methylation between SC283 and TAM428 within the ARG1 exon (Supplemental Figure S10). However, we were not able to analyze DNA methylation in the promoter regions of ARG1 and CARG due to polymorphisms in the promoter sequences of these genes.

To further understand how ARG1 gene expression is regulated, we examined histone H3 lysine methylation (H3Kme) patterns within the CARG and ARG1 exons, a region upstream of CARG, as well as the region shared by the CARG and ARG1 transcripts (Figure 8A). H3K4 and H3K36 methylation marks are generally associated with active transcription, whereas H3K9 methylation is a repressive mark associated with transcriptional silencing (Kouzarides, 2007) and is often linked to both DNA methylation and NAT-mediated regulation of gene expression (Li et al., 2012; Bohmdorfer and Wierzbicki, 2015). In general, H3K9me2 is more prevalent in facultative heterochromatin in gene-rich regions and H3K9me3 is often associated with constitutive heterochromatin (Peters et al., 2002, 2003).

Chromatin immunoprecipitation (ChIP) was conducted using antibodies specific to H3K4, H3K36, and H3K9 di- and trimethylation, followed by qPCR designed to amplify precipitated products from the indicated regions of the ARG1 and CARG genes to determine the level of chromatin modifications at these loci. At the 5'-UTR of ARG1, which covers the CARG/ARG1 overlap region, H3K4me2, H3K4me3, and H3K36me3 levels were dramatically higher in the resistant genotypes SC283 and SSD4 and reduced in the susceptible genotypes TAM428 and SSD65 (Figure 8, B and C), closely tracking with the levels of expression of this ARG1 gene. The chromatin of the ARG1 exon was also significantly enriched for H3K4me2, H3K4me3, and H3K36me3 in the resistant genotypes, whereas the levels of these marks were reduced in the susceptible genotypes (Figure 8, B and C), correlating with the loss of ARG1 expression in these genotypes. In contrast, despite the lower ARG1 expression, H3K36me2 was
Figure 8 Changes in active and repressive Histone H3 lysine methylation at CARG–ARG1 chromatin. A, Schematics showing the CARG and ARG1 genomic region. The location of the primers at the coding (Exon) and CARG/ARG1 overlap region used to analyze the level of H3K4, H3K36, and H3K9 methylation by ChIP assays are indicated by bars. The gray and black boxes indicate UTRs and exons, respectively. B–D, ChIP-qPCR analysis at CARG–ARG1 locus with antibodies against (B) H3K4me2/3, (C) H3K36me2/3, and (D) H3K9me2/3. ChIP was conducted on chromatin extracts with antibodies that recognize different histone H3 lysine methylation marks, and IgG was used as a background control. Precipitated DNA was quantified by qPCR, and the DNA enrichment is shown as a percentage of IP/input. Multiple comparisons of mean values were performed using Tukey–Kramer honest significant difference test ($P < 0.05$), and different letters indicate significant differences. Two independent experiments were performed with similar results. Ab, Antibody. SSD4 and SSD65 are resistant and susceptible recombinant inbred lines, respectively.
enriched in the exon of ARG1 chromatin of the susceptible genotypes (Figure 8C). Although H3K36 methylation is commonly associated with active transcription, previous studies have reported that it is also implicated in alternative splicing (Luco et al., 2010), which was observed in ARG1 of the susceptible genotypes (Figure 6). These data suggest that H3K36 methylation plays a role in the alternative splicing of ARG1 in the susceptible genotypes. Consistent with higher CARG gene expression, the CARG exon contained much higher levels of H3K4m2, H3K4me3, H3K36me2, and H3K36me3 in the susceptible genotypes versus the resistant genotypes (Figure 8, B and C).

H3K9 methylation is a repressive mark that is often triggered by small RNA (Holoch and Moazed, 2015). In contrast to H3K4 and H3K36 methylation, H3K9me2 and H3K9me3 levels were higher in the CARG/ARG1 overlap region in the susceptible genotypes, which exhibited lower ARG1 expression (Figure 8D). However, there were no significant differences in H3K9me2 and H3K9me3 at the ARG1 exon in either genotype (Figure 8D). H3K9me2 and H3K9me3 levels at the CARG exon were significantly higher in the resistant lines in which CARG expression was reduced, whereas low levels of H3K9 methylation at the CARG exon were observed in the susceptible lines where CARG was highly expressed (Figure 8D). In all cases, the control experiment was conducted on the same IP protein DNA complex using the primers at the constitutive sorghum Actin gene (Sobic.001G112600), which showed no difference in the level of histone H3 lysine methylation (Supplemental Figure S11).

Due to the extensive polymorphisms of the upstream region of CARG in the resistant and susceptible genotypes, histone H3 lysine methylation was not examined in the 5′-upstream region of CARG. Overall, however, the patterns of histone lysine methylation that could be assayed correlated well with gene expression patterns, but it is not clear if these are the causes or consequences of the reduced gene expression.

ARG1 confers resistance to fungal pathogens with distinct pathogenesis strategies

NLR-mediated resistance is often linked to plant immune responses to biotrophic and hemibiotrophic pathogens with race specificity (Jones and Dangl, 2006). To determine the specificity of ARG1, we tested the different genotypes for resistance to target spot, a fungal disease of sorghum caused by the necrotrophic fungus Bipolaris sorghicola (Supplemental Figure S12A). Unexpectedly, the plant responses to B. sorghicola were similar to those for Cs. Similarly, ARG1 conferred resistance to sorghum rust disease caused by the biotrophic fungus Puccinia purpurea (Supplemental Figure S12B). This resistance is, therefore, broadly effective against three species of fungal pathogens with three distinct modes of infection and pathogenesis. Resistance to distinct groups of pathogens is unexpected given that NLRs are a class of proteins that are generally linked to race-specific resistance, and, in some cases, known to promote susceptibility to necrotrophic fungi (Coll et al., 2011). Therefore, we identified a single NLR gene that causes broad spectrum and complete resistance to multiple unrelated fungal pathogens.

Comparative analysis of the ARG1 gene reveals distinct evolutionary relationships between resistant and susceptible genotypes

ARG1 encodes a typical NLR protein with N-terminal CC, NB-ARC, and LRR domains (Supplemental Data Set S7). Sequence comparisons revealed that ARG1 shares the highest sequence identity (54.61%) with RPP13-like protein in the wild rice species Oryza brachyantha (Ob), for which no functional data are available. RPP13 proteins from other plant species are also related to ARG1, with the Arabidopsis RPP13 showing significantly lower sequence identity (27.15%). We conducted phylogenetic analyses including ARG1, ObRPP13, and 84 functionally validated CC-NLRs retrieved from RefPlantNLR (Kourelis et al., 2021) to explore the evolutionary relationships of ARG1 to other CC-NLRs from different species. This comprehensive analysis showed that ARG1 is closely related to ObRPP13 (Supplemental Figure S13), which is consistent with their high sequence identity. Interestingly, Arabidopsis NRG1.2 closely clustered with ARG1. However, ARG1 and NRG1.2 share only 23.73% sequence identity. NRG1 was discovered in Nicotiana benthamiana because it was required for N protein-mediated resistance to tobacco mosaic virus (Peart et al., 2005). Arabidopsis RPP13 is a typical NLR that recognizes the Hyaloperonospora parasitica effector protein ATR13, which triggers resistance to biotrophic pathogens (Rentel et al., 2008).

A total of 397 NLR-encoding genes were identified from predicted gene models in the sorghum genome (Supplemental Figure S14 and Supplemental File S2), most of which were located on three chromosomes (Chr2, Chr5, and Chr8). In contrast, the CARG-deduced amino acid or DNA sequence is unique to the sorghum genome, with no similarity to other sequences in the database. Proteomic analysis of SC283 and TAM428 lines identified peptides that map to the ARG1 protein in the resistant lines, but no polypeptide was identified that maps to the CARG ORF in any of the genotypes. Furthermore, the putative 89 aa polypeptide that would be produced by that ORF would be unique to sorghum. Thus, although we cannot exclude the possibility that the ORF in CARG is translated, these data suggest that it is not and that the CARG transcript functions as a noncoding RNA.

We next assessed the genetic relationship of the ARG1 gene among many resistant and susceptible lines for which sequences were available from databases and our sequencing data. The phylogenetic relationship inferred from Maximum-likelihood analysis revealed a clear separation between the
resistant and susceptible lines (Figure 9; Supplemental File S3). The resistant lines form three sub-clusters; SC283, SC35C, and BTx378 form one sub-cluster, while PI585749 and PI586439 form another sub-cluster. KS115 forms its own separate sub-cluster, which also includes the distantly related ARG1 from SC283, as shown by the large distance in the phylogenetic tree. The susceptible lines fell into five sub-clusters. TAM428 and 28 susceptible genotypes make up one sub-cluster. The second sub-cluster includes PQ434 on its own and a third sub-cluster includes only PI525695, which is not closely related to TAM428. The fourth sub-cluster is composed of SQR, Tetron, and KP33-2, while the last sub-cluster consists of SQR and Ai4 (both originally collected from China), which were closely related to each other but were relatively distantly related to the other sub-clusters. These results point to variation within both the resistant and the susceptible sorghum genotypes.

Consistent with the above analyses, alignment of ARG1 sequences of the 42 genotypes revealed that all resistant lines carry intact ARG1 and are very similar to SC283, whereas ARG1 from the susceptible lines were identical or nearly identical to ARG1 from TAM428 or to SQR and Ai4, which have premature stop codons at different sites from TAM428 (Figure 2; Supplemental Figure S4 and Supplemental File S4). The pattern of sequence variation in the ARG1 gene confirms the differences between the different genetic backgrounds coming from diverse origins, with the three subclusters in the resistant group showing clear
differences in sequences. Despite extensive sequence searches, we identified no genotype that links the intact CARG NAT from the resistant ARG1 allele.

**Discussion**

Anthracnose is a major foliar diseases of sorghum that completely kills plants in the absence of resistance genes (Sherriff et al., 1995). Both the molecular mechanisms and the genes that regulate plant immunity to this pathogen are poorly understood. Here, we identified the ARG1 gene, encoding a plant immune receptor that confers broad spectrum and complete resistance to several distinct fungal pathogens. ARG1 is nested in an intron of a unique NAT, designated CARRIER OF ARG1 (CARG), and the entire CARG and ARG1 locus is flanked by MITE sequences that regulate the expression of CARG and ARG1 transcripts. A combination of MITE insertions in the 5' and 3'-CARG results in repressed expression of ARG1. The antisense NAT shares very limited sequence complementarity with the sense ARG1 transcript apart from a short segment of 158 nucleotides in TAM428 and 159 nucleotides in SC283. The 3'-UTR of CARG in the susceptible genotype produces small RNAs whose functional significance is unclear. CARG and ARG1 are transcribed in opposite orientations and exhibit inverse expression levels. The expression of the ARG1 allele from susceptible genotypes is associated with an increase in the repressive H3K9 di- and trimethylation marks within the CARG/ARG1 overlap region, as well as a reduction of these modifications in the CARG exon in the susceptible genotype. Abrogation of CARG expression is associated with the derepression of ARG1 expression, which in turn correlates with increased histone H3K4 and H3K36 methylation levels within the single ARG1 coding exon. In susceptible cultivars, basal transcription of ARG1 and CARG is likely maintained through a combination of mechanisms including interference with transcription, dsRNA, NAT-mediated histone methylation, and repressive chromatin states.

It should be acknowledged that the primary lesion most likely to be responsible for susceptibility is the premature stop codons present in all of the susceptible genotypes. The loss of the conserved LRR domain likely results in a nonfunctional protein, and may also lead to nonsense-mediated decay of the ARG1 transcript, which could explain its reduced steady-state levels (Lejeune, 2017). The increased level of CARG in susceptible lines may thus be a consequence of the loss of transcriptional interference due to reduced levels of ARG1 in these genotypes. According to this scenario, the changes in expression levels and chromatin modifications would be a consequence rather than a cause of a mutation in ARG1 that results in a loss of ARG1 transcript. However, at least in our Arabidopsis transgenic plants, high levels of ARG1 expression do not reduce the level of CARG in cis (Figure 7B).

There are a number of lines of evidence that suggest an alternative hypothesis, in which the NAT is a key player in the differentiation between resistant and susceptible genotypes. First, we note that all susceptible genotypes have both MITE insertions flanking the CARG–ARG1 locus and point mutations in ARG1. The tight association between two genetic lesions in the NAT and the absence of both of them in the ARG1 gene suggest that both lesions are required for resistance: one permits expression of the resistance gene due to the loss of the NAT, and the other permits high levels of expression of a functional NBs-LRR gene. However, because the polymorphisms in the two genes have not been separated, it is not possible at this time to determine whether both are required for the production of large quantities of functional ARG1 protein. The most straightforward way to determine this would be to genetically modify a resistant genotype such that CARG is expressed at high levels in situ. If this modification results in a susceptible phenotype despite the presence of an intact ARG1 gene, it would be possible to conclude that the loss of the NAT is required for full resistance.

There are also other scenarios that are worth entertaining. NBs-LRR genes are often found at new locations in different accessions or related species (Luo et al., 2012), and many of these “transposed” genes are not functional, likely due to local sequence context. Indeed, ARG1 is not present at a syntenic position relative to sorghum in the rice, Brachypodium distachyon, barley, teff, or Setaria italica genomes and is missing altogether from the maize genome, suggesting movement of this gene at some point in its evolutionary history. It is quite possible that ARG1 moved to a new position in sorghum and did not provide disease resistance, perhaps due to the absence of a strong, inducible promoter. In some lineages (those with the Indel), a MITE insertion triggered expression of an long non-coding RNAs/natural antisense transcript (lncRNA NAT), which repressed any residual expression of ARG1, and point mutations resulted in a loss of functional ARG1 altogether. In other lineages, relaxed purifying selection could have then resulted in the polymorphisms that may contribute to ARG1’s current unique broad-spectrum resistance. Subsequent strong selective pressure caused by disease could then have led to selection for a MITE insertion into ARG1, which both enhanced expression and made it inducible. According to this scenario, providing full resistance to the susceptible lines would require correction of both the ARG1 and CARG lesions.

The nature of ARG1 exon skipping is also unusual in that the skipping or the production of two transcripts from the same genomic template occurs in the absence of obvious well-defined intronic sequences in the ARG1 gene. Many resistance genes are regulated by differential splicing where premature stop codons introduced by frame shifts result in variant transcripts encoding proteins lacking LRR repeats (Yang et al., 2014). However, the functions of these transcripts or truncated proteins in the susceptible backgrounds are unknown.

Proteins with canonical NLR protein structure mediate recognition of virulence effectors, which then activate a very strong and race-specific resistance that varies depending on the pathogen strain. ARG1 encodes a typical NLR, which in
SC283 and other resistant genotypes confers resistance to distinct pathogen groups. These include the obligate biotrophic fungus *Puccinia purpurea* (which causes sorghum rust), the hemibiotrophic fungus *Cs*, as well as the necrotrophic fungus *B. sorghicola* (which causes target spot in sorghum). Broad-spectrum resistance to multiple distinct pathogenic species with disparate virulence strategies and lifestyles is extremely uncommon. In fact, some NLRs are known to actually promote susceptibility to a variety of other necrotrophic fungi in sorghum and other plants (Lorang et al., 2007, 2012). It is possible that ARG1 recognizes a conserved effector that is common to different plant pathogen lineages. Alternatively, the derepression of ARG1 may activate an immune response that is broadly effective against many pathogens (Thomma et al., 2011). Ligand-independent resistance due to the derepression of ARG1 may also be possible. The broad-spectrum resistance in the sorghum cultivar BK7 was attributed to a QTL that maps to the ARG1 chromosomal region, supporting a significant role for ARG1 in existing sorghum cultivars (Felderhoff et al., 2016).

In eukaryotic cells, noncoding RNAs affect gene expression through transcription interference, RNA masking, dsRNA-dependent mechanism, RNA interference, or antisense-mediated methylation (Faghihi and Wahlestedt, 2009; Bohmdorfer and Wierzbicki, 2015). In Arabidopsis, the role of antisense transcripts (COOLAIR) in the cold-induced, epigenetic silencing of Arabidopsis *FLOWERING LOCUS C* (FLC), a regulator of the transition to reproduction, is linked to switching of chromatin states at *FLC* during vernalization (Csorba et al., 2014). Inference of transcription and consequent changes in chromatin have also been observed in other systems (Xue et al., 2014). Due to the complementarity of parts of the CARG 3′-UTR and ARG1 5′-UTR and the identification of small RNAs in the 3′-UTR of CARG, we suggest that the low levels of expression in susceptible genotypes may be due, at least in part, to sense–antisense interference, and that this process may result in the changes in chromatin modification that we observed in both genes.

H3K4 methylation of ARG1 is significantly enriched in genotypes that show high levels of the expression of ARG1, as are H3K4 and H3K36 in the exon of CARG in genotypes that express high levels of that gene. Consistent with this finding, we observed enrichment of the repressive H3K9 methylation mark in the exon of CARG in resistant genotypes in which the expression of this gene is low. However, analyzing chromatin changes in the CARG promoter is complicated by the fact that the actual promoter region of this gene is poorly defined and is largely composed of transposable elements. Indeed, of the 1,500 bp of sequence upstream of the transcription start site in CARG of TAM428, only 26 bp are nontransposon sequences.

Clearly, additional studies are required to determine the degree to which changes in the expression of CARG mediate ARG1 regulation, and the means by which changes in histone methylation caused, or are caused by, changes in gene expression. However, we did find clear evidence that changes in histone methylation are associated with changes in the expression of these two genes, although our comparative analyses of DNA and histone methylation was hampered by high polymorphism in the DNA sequence in resistant and susceptible genotypes.

Genetic studies have defined multiple loci that control resistance to *Cs* (Perumal et al., 2009; Murali Mohan et al., 2010; Felderhoff et al., 2016). However, the identification of specific resistance genes and their mechanisms of action has been slow in coming. Our findings are significant both because of their direct application for controlling widespread and economically significant sorghum diseases and because this pair of genes represents an unusual regulatory mechanism of a known class of immune receptors. Indeed, resistance associated with a loss of the NAT of an immune receptor gene is unique. In addition, the MITE insertion in the 5′-ARG1 regulatory region confers inducible gene expression, adding to a growing body of evidence that transposable elements can be a significant source of regulatory information (Lisch, 2013).

Regardless of the molecular and cellular mechanisms involved, the CARG–ARG1 locus provides a unique resistance locus that can be easily introgressed into a variety of sorghum cultivars using CARG–ARG1-specific molecular markers. The resistance provided by ARG1 allele confers strong resistance to at least 10 distinct Cs strains tested, as well as 2 other fungal species. Transgenic expression of ARG1 in susceptible but adapted varieties of crop plants may also provide broad-spectrum resistance. Genome editing of the CARG and ARG1 genes in improved and adapted cultivars in order to generate broad-spectrum resistance will considerably shorten the breeding cycle and will make it possible to more precisely determine the means by which this unusual locus is regulated. A better understanding of the regulatory relationship between the NAT, ARG1, and the flanking MITEs will also likely provide important insights into the means by which novel patterns of gene regulation can rapidly evolve in plant genomes in response pathogens.

**Materials and methods**

**Plant growth**

The sorghum (*S. bicolor*) RILs were generated by crossing SC283 and TAM428 and advanced through single seed descent to the F6 generation and were maintained by self-fertilization. A total of 209 RIL lines were evaluated six times consecutively in the greenhouse. Plant growth conditions, methods of inoculation, and disease response assessments were as previously described (Prom et al., 2009). *Arabidopsis thaliana* Columbia-0 (Col-0) wild-type and the CARG–ARG1 transgenic plants were generated as described previously (Mengiste et al., 1997) and grown in a growth chamber under a 12-h photoperiod with 140–150 μE m⁻² s⁻¹ of fluorescent light (Philips F32T8/ADV835/ALTO-T8) at 22°C and 60% relative humidity.
Preparation of fungal cultures and plant disease assays

The Cs strains Csg1 and Csg2 were obtained from Dr Lisa Vaillancourt (University of Kentucky, Lexington). The other Cs strains are from different regions in Ethiopia and Nigeria (Supplemental Data Set S1). All strains were cultured on potato dextrose agar plates at 25°C. Fungal spores were harvested from 15- to 20-day-old cultures and suspended in ddH2O. The suspension was filtered through two layers of cheesecloth, and the concentration of spores was adjusted to 10⁶ spores·ml⁻¹. The spore suspension was uniformly sprayed onto 3- to 4-week-old sorghum plants. Plants were kept in humidity chambers for 2 days and transferred to the greenhouse with a temperature setting of 28°C with a 16-h light duration and with occasional misting to maintain high humidity. Disease responses were scored by visual assessment of disease symptoms or resistance responses, chlorosis, and fungal growth in planta. The detached leaf disease assay for Cs was conducted by drop inoculation of spores on leaves placed on wetted absorbent or filter paper and incubated in sealed transparent trays. A drop (20 µL of 10⁶ spores·ml⁻¹) of suspension was deposited on each leaf and disease evaluated by measuring lesion area and fungal growth. Total genomic DNA was isolated from Cs-inoculated leaves of each genotype using quick DNA extraction buffer (200-mM Tris–Cl, pH 7.5, 250-mM NaCl, 25-mM EDTA, pH 8, and 0.5% SDS). Fungal growth was assessed by qPCR amplification of the fungal rDNA and sorghum Actin as an internal control.

Rust (Puccinia purpurea)-infected sorghum leaves were collected from the Agronomy Center for Research and Education, West Lafayette, Indiana. The rust inoculum was maintained on rust-susceptible genotypes in the greenhouse. Inoculations and disease assays were conducted as described (White et al., 2014).

The target leaf spot fungus B. sorghicola isolates were obtained from Dr Burt H. Bluhm (University of Arkansas). The strain was cultured, harvested, and plants inoculated using the same method described for Cs strains. The concentration of spores was adjusted to 4 × 10⁶ spores·ml⁻¹ and the plants were inoculated as previously described (Borges, 1983).

Trypan blue staining

Leaf tissue samples from inoculated plants were collected for staining with trypan blue to reveal fungal growth in leaf tissue. The leaves were cleared in acetic acid: ethanol (1:3, v/v) solution overnight, followed by clearing using acetic acid: ethanol: glycerol (1:5:1, v/v/v) solution B for 3 h. The tissue was then stained with trypan blue (0.01% trypan blue in lactophenol) overnight. The stained tissue samples were rinsed multiple times and preserved in 60% glycerol for microscopic observation (Nikon ECLIPSE Ci).

RNA-seq analysis

TAM428 and SC283 plants were grown in soil for 3 weeks and inoculated with Csgl2 (10⁶ spore·ml⁻¹). At 0, 24, and 48 h after inoculation, the fifth leaves were collected from three biological replicates (approximately six plants each). Total RNA isolation was performed as described in the protocol of the Spectrum Plant Total RNA Kit with on-column DNase digestion (Sigma-Aldrich, USA), treated with DNase, and purified using the RNA Clean & Concentration TM-25 (ZYMOL RESEARCH). The quality of the total RNA was determined by NanoDrop and an Agilent 2100 Bioanalyzer. For each sample, 3-µg total RNA was used to prepare the mRNA-seq library according to the TrueSeq RNA Sample Prep Kit protocol (Illumina). Library quality control and quantification were performed with an Experion DNA 1K Chip (Bio-Rad) and a Qubit fluorometer (Invitrogen), respectively. A total of 734,963,453 high-quality reads (average length = 99 bp) were generated using an Illumina HiSeq 2500 sequencer (Supplemental Data Set S2). For each library, 75 million 100-bp paired-end sequences were generated using an Illumina HiSeq 2500 sequencer. After removing low-quality sequences containing uncalled bases (Ns), we used Tophat2 (version 2.1.1) software (Kim et al., 2013) to align the RNA-seq reads against the reference genome of BTx623 (PhytozomeV10: Sbicolor_313_v3.1). Tophat2 alignment parameters were set to allow a maximum of two mismatches and to exclude reads mapping to more than one position on the reference genome. Moreover, only reads for which both pairs successfully aligned were considered. The gene counts were extracted using the HTSeq python tool (Anders et al., 2015). Differential expression analyses were performed using the EdgeR package (Robinson et al., 2010) using empirical Bayesian methods. To filter out weakly expressed genes, only those genes with a minimum expression level of 1 RPKM (reads per kilobase per million mapped reads) in three replicates were included in the analysis. Genes with a LogFC > 1 (2-fold change) and false discovery rate < 0.05 and FDR < 0.05 were considered to be differentially expressed between conditions. To assess the variability among samples, we performed hierarchical clustering and dispersion analysis based on biological coefficient of variation. Hierarchical clustering was performed based on Euclidean distances. Dispersion was conducted using top 2,000 values in the EdgeR software package.

Functional classification analysis

To annotate entire gene sets of the sorghum and Cs genomes accurately, all protein sequences were analyzed using InterProScan 5.8-49.0 (Jones et al., 2014). We then used agriGO (http://bioinfo.cau.edu.cn/agriGO/) and ReviGO (http://revigo.irb.hr/) (Du et al., 2010; Supek et al., 2011) to identify the putative biological functions and biochemical pathways for the differentially expressed genes (DEGs) and to find statistically overrepresented gene ontology terms. To expand our functional analysis of the DEGs, we used MapMan software (http://mapman.gabipd.org) for visualization and biochemical pathway overlays as previously described (Lei et al., 2014). For MapMan analysis, all gene identification labels were converted into Sbicolor_79 label.
based on Sbicolor 3.1 annotation files (PhytozomeV10: Sbicolor_313_v3.1 synonym).

DNA isolation and whole genome sequencing
Among the RILs, 50 resistant and 50 susceptible plants were selected and used to construct two DNA bulks (RB and SB). To build the reference sequence, eight sorghum cultivars (Supplemental Data Set S4) were sequenced. For DNA extraction, 100-mg fresh leaf was harvested from each selected seedling and DNA was isolated using a DNeasy Plant Mini Kit (Qiagen, USA). Approximately 100-ng DNA from each sample was combined to construct two independent DNA bulks. The two DNA bulks were purified with a DNA clean-up & Concentration Kit (ZYMO Research, USA). A genomic DNA library was prepared for each DNA bulk using an Illumina TruSeq DNA Sample Preparation Kit (Illumina Inc., San Diego, CA, USA) according to the manufacturer's protocol. Each DNA library was sequenced on the Illumina HiSeq 2500 sequencing platform.

Bulk DNA sequencing and QTL analysis
The raw DNA-seq reads were trimmed and filtered to remove low-quality sequences using Fastx-tools (Pearson et al., 1997). Reads with a quality threshold lower than 30 and those shorter than 40 bp were discarded. The short reads from the two DNA bulks that passed the quality control were aligned to the reference genome of BTx623 (Phytozome V10: Sbicolor_313_v3.1) using BWA software (Version 0.7.12; Li and Durbin, 2009). Reads that aligned to more than one position in the reference genome were filtered out. The files were converted to BAM files using SAM tools (Li et al., 2009), sorted, and compared to locate duplicate records using Picard software (http://picard.sourceforge.net). Re-alignment (BAQ) was done to avoid false SNP calls near InDels. The resulting files were applied to GATK SNP-calling (version 3.3; McKenna et al., 2010; DePristo et al., 2011). SNP annotation was performed using SnpEff (version 4.1; Cinoglanli et al., 2012) with the sorghum annotation file (PhytozomeV10: Sbicolor_255_v2.1.gene.gff3). A total of 11,170 variants, including 9,567 SNPs, 755 insertions, and 848 deletions, were annotated in the QTL region. QTL analysis was performed as described previously (Takagi et al., 2013). The sorghum reference sequence was reconstructed by replacing nucleotides in BTx623 with the 1,826,960 SNP positions identified between the eight cultivars by aligning the short reads to the reference genome of BTx623 (PhytozomeV10: Sbicolor_313_v3.1). SNP-index was calculated at all SNP positions with Coval. All steps were performed using the QTL-seq Framework 1.4.4 pipeline (http://genome-e.ibrc.or.jp/home/bioinformatics-team/mutmap; Takagi et al., 2013). Slide window analysis was applied to SNP-index plots with 2-Mb window size and 50-kb increment. Identification of InDels, primer design, and marker analysis
To identify potential InDel markers between SC283 and TAM428, we detected sequence polymorphisms between them using the genome browser Integrative Genomics Viewer. Primer3Plus (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi) was used to design PCR primers with a length of 18–27 bp, GC content of 40%–60%, and PCR products of 100–700 bp. For marker analysis, genomic DNA was isolated from each selected RIL and parental line using a DNeasy Plant Mini Kit (Qiagen, USA). PCR amplification was performed in a 20-µL reaction containing 1 × PCR Buffer (1.5 mM MgCl2, 0.1 mM each dNTP, 0.5 µM of forward/reverse primers, 20–40 ng of genomic DNA, and 1.0 U GoTaq DNA polymerase (Promega, USA).

ChIP-qPCR
ChIP experiments were performed as described previously with minor modifications (Saleh et al., 2008). Leaf tissues (1.5 g) from 3-week-old plants were fixed with 1% (v/v) formaldehyde for 40 min at room temperature, and the chromatin samples were sonicated to yield 200–1,000-bp fragments. After pre-clearing of the chromatin samples with salmon sperm DNA/protein A agarose beads (EMD Millipore), immunoprecipitations were carried out with the appropriate antibodies to histone lysine methylation and reverse-cross-linking overnight at 65°C. Immunoprecipitated DNA samples were purified using a silica membrane column (MACHEREY-NAGEL Inc.) and eluted in 60-µL elution buffer. For qPCR, 2 µL of DNA was amplified using SYBR Green Supermix (Bio-Rad) with specific primers, as listed in Supplemental Data Set S8. The data are presented as percentage of input values. The antibodies used for the ChIP experiments were: H3K4me2 (07-030, EMD Millipore), H3K4me3 (07-473, EMD Millipore), H3K9me2 (ab1220, Abcam), H3K9me3 (07-442, EMD Millipore), H3K36me2 (07-369-1, EMD Millipore), H3K36me3 (ab9050, Abcam), and IgG (sc-2027, Santa Cruz) as a negative control.

RNA isolation
Total RNA was isolated from Cs-inoculated leaves and mock- or 2-nM chitin-treated leaves of each sorghum genotype and from leaves of each Arabidopsis transgenic line using TRI reagent (Molecular Research Center Inc.) according to the manufacturer’s instructions. The total RNA concentration and quality were measured using a NanoDrop 2000c spectrophotometer (Thermo Fisher Scientific).

RT-PCR and RT-qPCR analysis
After DNase I treatment (NEB), cDNA was synthesized from 2 µg of total RNA using M-MLV Reverse Transcriptase (Promega) according to the manufacturer’s protocol. For RT-PCR, the PCRs were carried out using GoTaq DNA Polymerase (Promega) with the primers listed in Supplemental Data Set S8. The reactions (Applied Biosystems 2720 Thermal Cycler) for the ARG1 and Actin genes consisted of 25, 28, 31, and 34 cycles in three steps:
95°C for 30 s, 57°C for 30 s, and 72°C for 2 min (ARG1 gene) or 30 s (Actin gene). Amplified PCR products were loaded on 1.5%-2.0% agarose gels, and bands were visualized by ethidium bromide staining. For RT-qPCR, reactions were performed using SYBR Green Supermix (BIO-RAD) with the primers listed in Supplemental Data Set S8, following the manufacturer’s instructions.

Generation of swapping constructs of CARG–ARG1 loci and plant transformation
A total of 6 CARG–ARG1 loci constructs were made in the pEarleyGate 104 (pEG104) vector backbone using an In-Fusion HD Cloning Kit (Clontech). These six constructs, SC283, TAM428, S1, S2, S3, and S4, are schematically represented in Figure 7B. The full-length and partial DNA amplification of CARG–ARG1 loci were carried out by PCR to generate DNA fragments for P1, P2, and P3 from both SC283 and TAM428, P4 from TAM428, P5 from SC283, P6 from TAM428, and P7 from SC283. These DNA fragments contained 15-bp overlapping regions to allow homologous recombination between DNA fragments. The In-Fusion reaction was conducted using mixed appropriate DNA fragments and linearized pEG104 according to the manufacturer’s instructions. The primers used in this work are listed in Supplemental Data Set S8. All six constructs were transformed into Agrobacterium tumefaciens GV3101, and subsequently, 5-week-old plants were used for Agrobacterium-mediated transformation by the floral dip method (Clough and Bent, 1998).

Construction of the GUS reporter gene-fusion plasmid, transfection, and GUS activity assay
A Nael/EcoRI fragment of pSAT1-Pnos-Venus-N (E4042) modified by introducing the GUS reporter and removing the nos promoter was used as a vector for introducing the ARG1 and CARG promoters: the 1,635-bp TAM428 ARG1 promoter; the 1,723-bp SC283 ARG1 promoter; the 1,153-bp TAM428 CARG promoter; and the 1,194-bp SC283 CARG promoter. Intron-deleted promoter-5′-UTR products were generated by performing a PCR-driven overlap extension method. Each gene segment in the promoter-5′-UTR region was spliced by segment-specific primers. The primers except for ARG1-SC283-Nael (-1722F) and ARG1-EcoRI (-1R) generated overlapping regions by including nucleotides that span the junction of each segment. The second or third PCR generated the intronless versions of ARG1 promoter-5′-UTR products, which were then inserted into the same vector used for promoter-GUS construction. All constructs were verified by sequencing and the plasmid DNA for transfection prepared using a Qiagen Plasmid Midi Prep kit. Primers used for the constructions are listed in Supplemental Data Set S8.

Approximately 1 × 10⁵ Arabidopsis protoplasts in 0.2 mL of MMg solution (0.4-M mannitol, 15-mM MgCl₂, and 4-mM MES, pH 5.7) were mixed with 10 μg of each plasmid DNA and an equal volume of PEG solution (40% [wt/vol] PEG4000; Fluka, 0.2-M mannitol and 0.1-M CaCl₂) for 5 min at room temperature. After incubation, the protoplasts were washed three times with 1 mL of W5 solution (154-mM NaCl, 125-mM CaCl₂, 5-mM KCl, 5-mM glucose, and 2-mM MES, pH 5.7). The protoplasts were resuspended gently in 1 mL of W5 and incubated at room temperature for 16 h. Fluorescence intensity (GUS and luciferase activity) was determined using a Tecan Infinite M200 pro microplate reader (Tecan). Co-transfection with a CaMV 35S-luciferase plasmid was used to determine the transfection efficiency. All transfection assays were performed at least in triplicate and repeated three times.

5′- and 3′-RACE
A RACE experiment was performed on total RNA samples using a SMARTer RACE 5′/3′ Kit (TaKaRa) according to the manufacturer’s protocol. Briefly, 1-μg RNA was treated with 5′-CDS Primer A or 3′-CDS Primer A at 72°C for 3 min and cooled to 42°C for 2 min. For the 5′-RACE cDNA synthesis reaction, 1 μL of the SMARTer II Oligonucleotide was added to the 5′-CDS Primer A treated RNA sample. The denatured RNAs were reverse transcribed with 8 μL of Master Mix at 42°C for 90 min and at 70°C for 10 min. The cDNAs were amplified by PCR. The PCR products were detected by agarose gel electrophoresis and purified using a PCR purification kit (MACHERY-NAGEL Inc.). The PCR products were sequenced, and sequencing results were compared with the genomic sequences.

DNA methylation analysis
Leaves of three plants per line were selected for DNA isolation. DNA was extracted from 4-week-old leaves using a DNeasy Plant Mini Kit (Qiagen), and DNA (200 ng) was used for bisulfite conversion using an EpiTect Bisulfite kit (Qiagen). The converted DNAs were used for PCR to evaluate the methylation status of ARG1, CARG, and Actin genes with specific primer sets. The primers are shown in Supplemental Data Set S8. The amplified products were gel purified (Gel Extraction kit; MACHERY-NAGEL Inc.), ligated into the pGEM-T Easy Vector (Promega), and transformed into Escherichia coli. The plasmid DNAs were isolated and sequenced using the T7 or M13 forward primers.

Small RNA-seq analysis
We applied an informatics pipeline to filter plant siRNAs and miRNAs from the complete set of small RNAs. A total of 228,228,937 distinct small RNA reads from 12 sorghum libraries with Cs or mock-inoculated plants were analyzed using the pipeline. As a first step, the adaptors and low-quality reads were removed using FASTX-Toolkit (Gordon and Hannon, 2010). The next step involved removing structural RNAs such as tRNAs and rRNAs. The third step involved selecting the RNA read sizes between 18 nt and 28 nt. The fourth step was to remove low-abundance small RNAs (retaining only those with less than 10 transcripts per million in at least one of 12 libraries), Cs genome reads, as well as highly repetitive small RNAs (those with more than 20
hits to the genome). A total of 121,338 distinct small RNAs were retained. Finally, mirDeep-P (Yang and Li, 2011) was employed to detect predicted miRNAs. To identify consistent miRNAs, all small RNA libraries were separately processed based on the above method. miRNAs were considered to be candidate miRNAs if they could be detected in three libraries with the same treatment in SC283 or TAM428. To further verify our predicted miRNAs, highly similar homologs in miRBase V21 were identified using Segemel (Hoffmann et al., 2009). miRNAs that passed all filter processing steps were identified as novel miRNAs. All small RNA-seq reads were aligned against the BTx623 reference genome (PhytozomeV10: Sbicolor_313_v3.1) using Tophat2 (version 2.1.1) software (Kim et al., 2013).

**Statistical tests**

Statistical parameters are presented in the figures and figure legends. Analyses of gene expression, fungal biomass, and ChIP-qPCR data were performed using at least three replicates, and data are shown as mean ± standard deviation (so). Statistically significant differences were determined by least significant difference, Student’s t-test, and Turkey’s honest significant difference test. Significance was considered to be *P* < 0.05, **P** < 0.01. The statistical analysis was performed using JMP 9.0, JMP 16, and GraphPad Prism 6.0 software. Summaries of statistical analyses are provided in Supplemental File S5.

**Accession numbers**

The RNA-seq and microRNA-seq data have been submitted to NCBI https://www.ncbi.nlm.nih.gov/bioproject/PRJNA667277 under accession number PRJNA667277. The nucleotide sequences of the CARG–ARG1 loci were sequenced or downloaded from NCBI. Additional data related to this paper may be requested from the authors.

**Supplemental data**

The following materials are available in the online version of this article.

**Supplemental Figure S1.** SNP-index and Δ (SNP-index) plots for 10 chromosomes of bulked DNA from resistant and susceptible recombinant inbred sorghum lines.

**Supplemental Figure S2.** Location and sequences of MITEs in the CARG–ARG1 locus.

**Supplemental Figure S3.** ARG1 and CARG gene expression in resistant and susceptible genotypes.

**Supplemental Figure S4.** Amino acid sequence alignment of ARG1 protein from different resistant and susceptible genotypes.

**Supplemental Figure S5.** Disease responses of sorghum genotypes carrying different ARG1 alleles.

**Supplemental Figure S6.** Disease responses of F2 plants after drop inoculation with Cs spores.

**Supplemental Figure S7.** Sequence alignments of ARG1 alleles from different genotypes.

**Supplemental Figure S8.** The 3′-UTR of the CARG NAT gene produces small RNAs.

**Supplemental Figure S9.** Small RNAs derived from the 3′-UTR of the Natural Antisense RNA transcript gene in susceptible genotype show similarity to sorghum miR6225.

**Supplemental Figure S10.** DNA methylation analysis of ARG1 exons in SC283 and TAM428.

**Supplemental Figure S11.** H3K4, H3K9, and H3K36 di- and tri-methylation at the sorghum Actin gene in anthracnose resistant and susceptible lines.

**Supplemental Figure S12.** ARG1 confers resistance to the fungal diseases target spot and rust.

**Supplemental Figure S13.** Phylogenetic analysis of ARG1 with 85 other CC containing NLR receptors.

**Supplemental Figure S14.** Phylogenetic trees of NLR genes from Sorghum.

**Supplemental Data Set S1.** Sorghum natural variants used in the study.

**Supplemental Data Set S2.** Responses of sorghum cultivars to different isolates of Cs.

**Supplemental Data Set S3.** Phenotyping of RILs and parents for Cs resistance.

**Supplemental Data Set S4.** Summary of Illumina Hiseq 2500 sequencing for SC283 x TAM428 RILs and eight sorghum cultivars.

**Supplemental Data Set S5.** Genotyping of RILs using 12 InDel markers. The resistant allele (SC283) is marked 1; the susceptible allele (TAM428) is marked 2. Rec. is recombination.

**Supplemental Data Set S6.** Sequence variants within the 5′-UTR, exon, and 3′-UTR of predicted 29 genes in the 10–11 Mb on chromosome 7 between SC283 and TAM428.

**Supplemental Data Set S7.** List of domain hits by NCBI conserved domains, SMART, and InterPro analyses.

**Supplemental Data Set S8.** List of primers used in this study.

**Supplemental File S1.** Genomic structures and sequences of CARG–ARG1 genes in TAM428 and SC283.

**Supplemental File S2.** Text file of protein sequences used for the phylogenetic analysis shown in Supplemental Figure S14.

**Supplemental File S3.** Text file of protein sequences used for the phylogenetic analysis shown in Figure 9.

**Supplemental File S4.** Alignment of ARG1 sequences from different genotypes.

**Supplemental File S5.** Summary of statistical analyses.

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