Genome-wide identification and characterization of long non-coding RNAs conferring resistance to Colletotrichum gloeosporioides in walnut (Juglans regia)

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Abstract

Background: Walnut anthracnose caused by *Colletotrichum gloeosporioides* (Penz.) Penz. and Sacc. is an important walnut production problem in China. Although the long non-coding RNAs (lncRNAs) are important for plant disease resistance, the molecular mechanisms underlying resistance to *C. gloeosporioides* in walnut remain poorly understood.

Results: The anthracnose-resistant F26 fruits from the B26 clone and the anthracnose-susceptible F423 fruits from the 4-23 clone of walnut were used as the test materials. Specifically, we performed a comparative transcriptome analysis of F26 and F423 fruit bracts to identify differentially expressed LncRNAs (DELs) at five time-points (tissues at 0 hpi, pathological tissues at 24 hpi, 48 hpi, 72 hpi, and distal uninoculated tissues at 120 hpi). Compared with F423, a total of 14525 DELs were identified, including 10645 upregulated lncRNAs and 3846 downregulated lncRNAs in F26. The number of upregulated lncRNAs in F26 compared to in F423 was significantly higher at the early stages of *C. gloeosporioides* infection. A total of 5 modules related to disease resistance were screened by WGCNA and the target genes of lncRNAs were obtained. Bioinformatic analysis showed that the target genes of upregulated lncRNAs were enriched in immune-related processes during the infection of *C. gloeosporioides*, such as activation of innate immune response, defense response to bacterium, incompatible interaction and immune system process, and enriched in plant hormone signal transduction, phenylpropanoid biosynthesis and other pathways. And 124 known target genes for 96 hub lncRNAs were predicted, including 10 known resistance genes. The expression of 5 lncRNAs and 5 target genes was confirmed by qPCR, which was consistent with the RNA-seq data.

Conclusions: The results of this study provide the basis for future functional characterizations of lncRNAs regarding the *C. gloeosporioides* resistance of walnut fruit bracts.

Background

Walnut (*Juglans regia* L.) is a diploid tree species (2n = 32), with approximately 667 Mb per 1C genome and an N50 size of 464955 (based on a genome size of 606 Mbp) [1]. It is an ecologically important ‘woody oil’ tree species worldwide [2], and its kernel is a rich source of nutrients with health benefits for humans [3]. The peptides extracted from walnut seeds have antioxidant and anticancer activities and have the protective effects on the oxidative damage induced by H2O2 [4]. Recent advances in biotechnology and genomics show potential to accelerate walnut breeding, such as gamma-irradiated pollen inducing haploid walnut plants [5], constructing the novel Axiom J. regia 700K SNP array [6], and combining different assemblies to obtain the optimal version [7]. Walnut anthracnose caused by *Colletotrichum gloeosporioides* (Penz.) Penz. and Sacc can cause leaf scorch or defoliation and fruit gangrene, which is currently the disastrous disease in walnut production [8]. Due to the long incubation period of anthracnose, the concentrated onset time, and the strong outbreak, the use of chemical fungicides is still the main method of disease control [9]. The *C. gloeosporioides* lifestyle transitions associated with the infection of the host include the following three stages: attachment, biotrophy, and
necrotrophy [10]. The pathogen of *C. gloeosporioides* in walnut overwinters in the diseased part with mycelium, and begins to move when the temperature reaches 11-15°C in the following spring [11]. Specifically, the formation of adherent cells is critical for fungal development during the *C. gloeosporioides* infection [12]. In a previous study, LAC2 was revealed to contribute to the formation of adherent cells to enhance the pathogenicity of *C. gloeosporioides* [13]. However, it is unclear how walnuts recognize and resist infections by *C. gloeosporioides*, and the regulatory network of hub and peripheral genes underlying the resistance of walnuts to *C. gloeosporioides* remains uncharacterized. Therefore, elucidating the molecular basis of this resistance mechanism is imperative for the breeding of walnut resistant to *C. gloeosporioides* [8, 14, 15].

Long non-coding RNA (lncRNA) is a type of RNA comprising 200–1,000,000 nt and structural characteristics similar to those of mRNA, but it does not encode a protein [16]. The lncRNAs were initially considered to be the transcription ‘noise’ of protein-coding genes, and were often ignored in transcriptome analyses [17]. However, the continuous development of sequencing technologies and transcriptome analyses has revealed that many lncRNAs in *Arabidopsis thaliana* [18], *Triticum aestivum* [19], *Zea mays* [20], and other plant species are related to stress responses, morphological development, and fruit maturation. For example, a heat-responsive lncRNA (TCONS_00048391) is an eTM for bra-miR164a and may be a competing endogenous RNA (ceRNA) for the target gene NAC1 (Bra030820), with effects on bra-miR164a expression in Chinese cabbage (*Brassica rapa* ssp. *chinensis*) [21]. Qin et al confirmed that the DROUGHT INDUCED IncRNA regulates plant responses to abiotic stress by modulating the expression of a series of stress-responsive genes [22]. In *A. thaliana*, two IncRNAs, COOLAIR and COLDAIR, are associated with FLOWERING LOCUS C and play an crucial role in vernalization [23,24].

Many recent studies have proved that IncRNAs are important for plant–pathogen interactions. A role for nine hub IncRNAs and 12 target genes in the resistance of *Paulownia tomentosa* to witches’broom was uncovered via a high-throughput sequencing experiment, and their functions were analyzed with an RNA-lncRNA co-expression network model [25]. In tomato (*Solanum lycopersicum*), the IncRNA16397-GRX21 regulatory network reportedly decreases the reactive oxygen species content and cell membrane damage to enhance the resistance to *P. infestans* [26]. Moreover, the involvement of the WRKY1-IncRNA 33732-RBOH module in regulating H$_2$O$_2$ accumulation and resistance to *P. infestans* was determined based on a comparative transcriptome analysis [27]. In cotton (*Gossypium* spp.), a functional analysis demonstrated that a lack of two hub IncRNAs, GhIncNAT-ANX2 and GhIncNAT-RLP7, enhances seedling resistance to *Verticillum dahliae* and *Botrytis cinerea*, possibly because of the associated upregulated expression of LOX1 and LOX2 [28]. In wheat (*Triticum aestivum* L.), IncRNAs have a tissue-dependent expression pattern that can respond to powdery mildew infections and heat stress [29]. Additionally, four kinds of IncRNAs have important effects on *Puccinia striiformis* infections [30]. However, there are no reports regarding the role of IncRNAs in the walnut fruit resistance to anthracnose.

In this study, Illumina HiSeq 4000 sequencing was used to analyze the disease-resistant (F26) and susceptible (F423) fruit bracts at different *C. gloeosporioides* infection stages. The number and characteristics of IncRNAs were analyzed. Additionally, the hub IncRNAs related to disease resistance
were screened and functionally analyzed to predict the role of IncRNAs in walnut fruit bract resistance to anthracnose. To the best of our knowledge, this is the first report on walnut IncRNAs and their biological functions related to fruit bract resistance to *C. gloeosporioides*. Our data may be a useful resource for clarifying the regulatory functions of IncRNAs influencing walnut fruit resistance to *C. gloeosporioides*.

**Results**

**Symptoms and physiological changes of walnut fruit infected by *C. gloeosporioides***
The resistant (F26) and susceptible (F423) fruit bracts were infected by *C. gloeosporioides*, the fruit bracts of F423 showed obvious symptoms at 48 hpi; the disease-resistant fruit F26 at 72 hpi. The susceptible samples showed obvious *C. gloeosporioides* conidial at 120 hpi (Fig 1a). During the infection, the activities of some enzymes and the content of hormones also changed correspondingly. Compared to the F423, the activities of chitinase, ROS-scavenging enzymes (catalase, CAT and superoxide dismutase, SOD) and the content of H$_2$O$_2$ in F26 were higher (Fig 1b-e). The content of salicylic acid (SA) and jasmonic acid (JA) in F26 was significantly higher than that in F423, and reached a peak at 72hpi after infection (Fig 1f, g).

**Whole genome identification of IncRNAs expressed in walnut fruit bracts***

To identify IncRNAs expressed in walnut fruits in response to *C. gloeosporioides*, we constructed 20 cDNA libraries from the anthracnose-resistant and the anthracnose-susceptible walnut fruits at the following five infection stages: tissue at 0 hpi (hours post inoculation), infected tissue at 24, 48, and 72 hpi, and distal uninoculated tissue at 120 hpi (Additional file 1: Table S1). The libraries were sequenced with an Illumina HiSeq 4000 platform. A total of 265.4 Gb clean data were obtained, with an average of 13.27 Gb per library. Approximately 69.7% of the clean reads in all libraries were mapped to the walnut reference genome (Additional file 2: Table S2). The aligned transcripts were assembled, combined, and screened with the FEELnc software to obtain 22,336 IncRNAs (length ≥ 200 nt, ORF coverage < 50%, and potential coding score < 0.5), including 18,403 unknown IncRNAs (23.97%) and 3,933 known IncRNAs (5.12%) (Fig 2).

**Characterization of walnut fruit bract IncRNAs***

A total of 58,369 mRNAs and 22,336 IncRNAs were obtained for the walnut fruit bracts (all samples combined) (Additional file 3: Table S3; Additional file 4: Table S4). The IncRNAs were characterized according to their locations relative to the partner RNA. A total of 40,429 (67.57%) IncRNAs were located in intergenic regions (i.e., only 32.43% genic IncRNAs). Additionally, 19,767 (48.89%) and 7,302 (37.63%) of the intergenic IncRNAs and genic IncRNAs were located in the antisense strand, respectively (Fig 3a) (Additional file 5: Table S5). Most IncRNAs contained two or three exons, which differentiated them from mRNAs (Fig 3c). Moreover, there was considerable diversity in the distribution of mRNA and IncRNA lengths (Fig 3b). Furthermore, the expression level of most IncRNAs was significantly lower than that of mRNAs (Fig 3d).
Differentially expressed lncRNAs at various infection stages

The lncRNAs that were differentially expressed between the disease-susceptible F423 fruits and the disease-resistant F26 fruits at different *C. gloeosporioides* infection stages were analyzed. Compared with F423, a total of 14,525 DELs were identified, including 10,645 up-regulated lncRNAs and 3,846 down-regulated lncRNAs in F26. The number of upregulated and downregulated lncRNAs in the various comparisons were respectively as follows: 7,668 and 1,386 in the F26_0hpi vs F423_0hpi comparison; 6,910 and 1,165 in the F26_24hpi vs F423_24hpi comparison; 1,721 and 1,593 in the F26_48hpi vs F423_48hpi comparison; 898 and 1,133 in the F26_72 hpi vs F423_72 hpi comparison; and 4,711 and 550 in the F26_120 hpi vs F423_120 hpi comparison (Fig 4a, b) (Additional file 6: Table S6). Additionally, compared with F423, a total of 34,007 differentially expressed mRNAs were identified, including 15,247 upregulated mRNAs and 13,198 downregulated mRNAs in F26. The number of upregulated and downregulated mRNAs in the various comparisons were respectively as follows: 6,836 and 4,622 in the F26_0 hpi vs F423_0 hpi comparison; 6,392 and 3,955 in the F26_24 hpi vs F423_24 hpi comparison; 3,454 and 4,347 in the F26_48 hpi vs F423_48 hpi comparison; 2,709 and 3,113 in the F26_72 hpi vs F423_72 hpi comparison; and 4,976 and 3,563 in the F26_120 hpi vs F423_120 hpi comparison (Fig 4c, d) (Additional file 7: Table S7). These results revealed the similarities in the expression of lncRNAs and mRNAs. And the number of upregulated lncRNAs and mRNAs in F26 compared to in F423 was significantly higher at the early stages of *C. gloeosporioides* infection.

Identification of co-expressed lncRNA modules

To identify the hub lncRNAs and predict their potential target genes in trans-regulatory relationships, a weighted gene co-expression network analysis (WGCNA) was used to generate a correlation matrix of the expression levels of 10,645 upregulated lncRNAs and 15,247 upregulated mRNAs. A total of 19 expression modules were screened (Fig 5a) (Additional file 8: Table S8). The relationships between modules and the resistance traits of the walnut fruit bracts were analyzed and four significantly correlated modules (|r| ≥ 0.8) were identified. The MEviolet module was correlated with F26_0hpi (r = 0.95, p = 9e−11), which contains 406 lncRNAs and 1350 mRNAs. The MElightyellow module was correlated with F26_24hpi (r = 0.86, p = 1e−06), which contains 165 lncRNAs and 892 mRNAs. The MBrown2 module was correlated with F26_48hpi (r = 0.82, p = 8e−0.86), which contains 128 lncRNAs and 224 mRNAs. The MEwhite module was correlated with F26_72hpi (r = 0.81, p = 1e−05), which contains 111 lncRNAs and 378 mRNAs (Fig 5c). Regarding F26_120 hpi, the rand p value for the MEorange module was 0.73 and 3e−0.4, respectively. The highest r value (0.77) for F423 was calculated for the MEdarkseagreen module and F423_48hpi (Fig 5b). And the MEorange module contains 76 lncRNAs and 227 mRNAs (Fig 5c). These results suggested that lncRNAs are closely related to the disease resistance of walnut fruit bracts.

Enrichment analysis of genes co-expressed with lncRNAs

The GO and KEGG pathway databases were used to analyze the genes co-expressed with lncRNAs in each significant module and MEorange module. In the MEviolet module, a total of 208 GO terms were
assigned, including 106, 8 and 94 GO terms in “biological process”, “cellular component” and “molecular functions”, respectively (Additional file 9: Table_S9). Among these enriched GO terms, most of them were related to biosynthesis and gene expression regulation, and the ones related to plant immunity were “response to stimulus” (GO:0050896) (187 genes) and “cellular response to stimulus” (GO:0051716) (114 genes) (Fig 6a). In total, 104 enriched KEGG pathways were identified, of which 30 pathways were significantly enriched in this module (Additional file 10: Table_S10). The top 30 significantly enriched pathways for target genes are mentioned in Fig 7a. “Plant hormone signal transduction” (ko04075) (22 genes), “Fatty acid metabolism” (ko01212) (15 genes), “Fatty acid elongation” (ko00062) (12 genes), “Ribosome” (ko03010) (12 genes), and “Spliceosome” (ko03040) (11 genes) were the most significant KEGG pathways.

In the MElightyellow module, a total of 164 GO terms were assigned, including 79, 16 and 69 GO terms in “biological process”, “cellular component” and “molecular functions”, respectively (Additional file 9: Table_S9). Among them, GO terms related to plant immunity included “activation of innate immune response” (GO: 0002218) (4 genes), “activation of immune response” (GO: 0002253) (4 genes), and “induced systemic resistance, jasmonic acid mediated signaling pathway” (GO: 0009864) (3 genes) (Fig 6b). In total, 93 enriched KEGG pathways were identified, of which 30 pathways were significantly enriched in this module (Additional file 10: Table_S10). The top 30 significantly enriched pathways for target genes are mentioned in Fig 7b. “Starch and sucrose metabolism” (ko00500) (14 genes), “Plant hormone signal transduction” (ko04075) (13 genes), “Phenylpropanoid biosynthesis” (ko00940) (11 genes), “Biosynthesis of amino acids” (ko01230) (10 genes), and “DNA replication” (ko03030) (8 genes) were the most significant KEGG pathways.

In the MEbrown2 module, a total of 126 GO terms were assigned, including 89, 5 and 32 GO terms in “biological process”, “cellular component” and “molecular functions”, respectively (Additional file 9: Table_S9). In addition to the terms related to biological metabolism and gene expression regulation, the items related to plant immunity “response to endogenous stimulus” (GO:0009719) (15 genes), “cellular response to endogenous stimulus” (GO:0071495) (13 genes) and “cellular response to hormone stimulus” (GO:0032870) (12 genes) were also enriched significantly (Fig 6c). In total, 38 enriched KEGG pathways were identified, of which 30 pathways were significantly enriched in this module (Additional file 10: Table_S10). The top 30 significantly enriched pathways for target genes are mentioned in Fig 7c. “Cyanoamino acid metabolism” (ko00460) (3 genes), “Plant hormone signal transduction” (ko04075) (6 genes), “Nitrogen metabolism” (ko00910) (2 genes), “Terpenoid backbone biosynthesis” (ko00900) (2 genes) were the most significant KEGG pathways.

In the MEwhite module, a total of 142 GO terms were assigned, including 95, 4 and 43 GO terms in “biological process”, “cellular component” and “molecular functions”, respectively (Additional file 9: Table_S9). Among the biological process category, the significantly over represented GO terms were “response to stimulus” (GO: 0050896) (67 genes), followed by “response to stress” (GO: 0006950) (51 genes) and “defense response” (GO: 0006952) (43 genes), which were all related to plant immunity. In addition, other terms related to plant immunity were also enriched, such as “immune system process”
(GO:0002376) (14 genes), “response to biotic stimulus” (GO:0009607) (14 genes) and “innate immune response” (GO:0045087) (13 genes), etc (Fig 6d). In total, 54 enriched KEGG pathways were identified, of which 30 pathways were significantly enriched in this module (Additional file 10: Table_S10). The top 30 significantly enriched pathways for target genes are mentioned in Fig 7d. “Carbon metabolism” (ko01200) (5 genes), “Cysteine and methionine metabolism” (ko00270) (4 genes), “Amino sugar and nucleotide sugar metabolism” (ko00520) (4 genes) were the most significant KEGG pathways.

In the MEorange module, a total of 128 GO terms were assigned, including 87, 8 and 33 GO terms in “biological process”, “cellular component” and “molecular functions”, respectively (Additional file 9: Table_S9). Among the biological process category, “response to organic substance” (GO: 0010033) (14 genes), “response to endogenous stimulus” (GO: 0009719) (13 genes), and “response to external stimulus” (GO: 0009605) (10 genes) etc., associated with plant immunity were significantly enriched (Fig 6e). In total, 32 enriched KEGG pathways were identified, of which 30 pathways were significantly enriched in this module (Additional file 10: Table_S10). The top 30 significantly enriched pathways for target genes are mentioned in Fig 7e. “Plant hormone signal transduction” (ko04075) (4 genes), “Thiamine metabolism” (ko00730) (3 genes), “Starch and sucrose metabolism” (ko00500) (3 genes) and “Fatty acid degradation” (ko00071) (2 genes) were the most significant KEGG pathways.

### Network analysis of hub IncRNAs

The hub IncRNAs are important for regulating the whole network. Therefore, we screened the 96 hub IncRNAs and 124 known target genes according to their weight value and connectivity in five modules (Additional file 11: Table_S11). In the MEviolet module, the 25 known target genes for 15 hub IncRNAs were found to be involved in multiple functions (Fig 7a), such as probable galacturonosyl transferase 10 and ultraviolet-B receptor UVR8-like. In addition, target genes encoding receptor-like serine / threonine-protein kinase NCRK (XM_018958556.1) and eukaryotic translation initiation factor 5A-2-like (XM_018994862.1) are known resistance genes (Fig 8a). In the MElightyellow module, 16 hub IncRNAs were generated and their 22 known target genes were involved in many functions (Fig 8b). And the target genes encoding G-type lectin S-receptor-like serine/threonine-protein kinase LECRK1 (XM_018950446.1), probably inactive leucine-rich repeat receptor-like protein kinase At2g25790 (XM_018989953.1) and TMV resistance protein N-like (XM_018961957.1) were known resistance genes (Fig 8b). In the MEbrown2 module, 24 hub IncRNAs and their 15 known target genes were generated (Fig 8c), the target gene encoding probable LRR receptor-like serine / threonine-protein kinase At3g47570 (XM_018962714.1) was known resistance gene (Fig 8c). In the MEwhite module, 23 hub IncRNAs were generated and their 38 known target genes were involved in many functions (Fig 8d). The target genes encoding putative disease resistance protein At1g50180 (XM_018965430.1), probable LRR receptor-like serine/threonine-protein kinase At1g63430 (XM_018973294.1) and L-type lectin-domain containing receptor kinase IV.2-like (XM_018954279.1) were known resistance genes (Fig 8d). In the MEorange module, 18 hub IncRNAs were generated and their 24 known target genes were involved in many functions (Fig 8e). And the target gene encoding the inactive LRR receptor-like serine / threonine-protein kinase BIR2 (XM_018967526.1) was known resistance gene (Fig 8e). All disease resistance genes in walnut are listed in Additional file 12:
Table_S12. These results suggested that lncRNAs may participate in the resistance of walnut bracts to *C. gloeosporioides* by acting on their target genes. Based on the enrichment results of KEGG, we predicted the possible pathway of hub lncRNAs (Additional file 13: Table_S13). Most of the hub lncRNAs and its target genes in the five modules are enriched in the pathways of material metabolism and biosynthesis. In the white module, IncRNA MSTRG.94840.7 and MSTRG.103441.8 were enriched in “plant pathogen interactions” and “plant hormone signal transduction” pathways, which may be related to plant immunity.

**Validation of hub IncRNAs and target genes**

We randomly selected 5 hub IncRNAs and 5 target genes for qRT-PCR analysis with the aim to validate the expression profiles between F26 and F423 obtained by RNA-Seq. The list of hub IncRNAs specific primers used for qRT-PCR analysis is listed in Additional file 14: Table_S14. The hub lncRNAs selected for qRT-PCR confirmation were MSTRG.13585.8, MSTRG.152205.1, MSTRG.11713.16, MSTRG.112028.8, and MSTRG.62751.2, the target genes were related to probable galacturonosyl transferase 10 (LOC109014322), G-type lectin S-receptor-like serine / threonine-protein kinase LECRK1 (LOC108979712), NHL repeat-containing (LOC108987880), probable LRR receptor-like serine/threonine-protein kinase At3g47570 (LOC108989177), and putative disease resistance protein At1g50180 (LOC108991254). The qRT-PCR analysis showed that the expression of MSTRG13585 and LOC109014322 peaked at 0hpi, MSTRG11713, MSTRG152205, LOC108979712 and LOC108987880 at 24hpi, MSTRG112028 and LOC108989177 at 48hpi, MSTRG62751 and LOC108991254 at 72hpi (Fig 9), which were consistent with the RNA-seq data (Additional file 15: Table_S15), with similar trends observed for the hub lncRNAs and their target genes.

**Discussion**

In previous studies, lncRNAs were identified and analyzed in various biological processes important for seed development [43], photomorphogenesis [44], fruit development [45,46], and biotic and abiotic stress responses [22, 47]. Additionally, there has been substantial research on the role of lncRNAs in plant–pathogen interactions. In *A. thaliana*, lncRNAs reportedly enhance the resistance to *Pseudomonas syringae pv. tomato* DC3000 by promoting *PR1* expression [48]. In tomato, IncRNA23468 functions as a ceRNA that modulates NBS-LRR gene expression by mimicking the target of miR482b, thereby increasing the resistance to *P. infestans* [49]. Walnut anthracnose has been responsible for the premature fruit drop and yield losses that have adversely affected walnut production in China [13]. In this study, we investigated the role of lncRNAs in the resistance of walnut fruit bracts to anthracnose based on sequence analyses. Walnut anthracnose is caused by *C. gloeosporioides*, which completes its infection process as a hemibiotroph [10,50]. First, conidia germinate to generate appressoria, which produce invasion pegs that initiate the infection into susceptible plants. The primary mycelium produced in plant cells exists as a biotroph, after which the secondary mycelium produced in the infected site switches to necrotrophic growth [51,52]. We previously determined that the *C. gloeosporioides* life cycle in walnut tissue involves attachment at 24hpi, biotrophy at 48hpi, and necrotrophy at 72hpi (data unpublished). In this study, RNA-seq was performed to build the IncRNA and mRNA profiles of the walnut fruit bract tissue.
at 0 hpi, infected tissue at 24, 48, and 72 hpi, and distal uninoculated tissue at 120 hpi. A total of 58,369 mRNAs and 22,336 IncRNAs were identified, including 3,933 known IncRNAs and 18,403 unknown IncRNAs. Consistent with the results of similar studies on other organisms, the identified putative IncRNA had fewer exons, shorter transcripts, and lower expression levels than protein-coding genes [53,54].

The release of walnut reference genome [1], enabled the study of walnut genetics at a genome-wide scale. Based on the reference genome, the whole-genome resequencing [55], the development of high-density genotyping tools [56], and the genetic dissection of important agronomical traits in walnut [57] have been completed. The development of bioinformatic analysis technology has enabled researchers to reveal that IncRNA functions and characteristics are far more complex than previously thought [16]. A recent comparative transcriptome analysis between wild-type and WRKY1-overexpressing tomato plants revealed 199 IncRNAs (DELs) and indicated that many of the IncRNA target genes that are likely affected by WRKY1 and associated with the resistance of tomato to P. infestans are involved in the response to biotic stimulus (GO:0009607) and plant-pathogen interaction (KO4626) [26]. In another recent study, 4,594 putative IncRNAs were identified in comprehensive dynamic IncRNA expression networks under heat stress conditions. Co-expression networks revealing the interactions among the differentially expressed IncRNAs, mRNAs, and microRNAs indicated that several phytohormone pathways are associated with heat tolerance, including salicylic acid and brassinosteroid pathways [21]. In the current study, we obtained 10,645 upregulated IncRNAs and 15,247 upregulated mRNAs among the five comparisons (F26_0hpi vs F423_0hpi, F26_24 hpi vs F423_24 hpi, F26_48 hpi vs F423_48 hpi, F26_72 hpi vs F423_72 hpi, and F26_120 hpi vs F423_120 hpi). The number of up-regulated IncRNAs and mRNAs in the F26 vs F423 was significantly higher at the early stages of C. gloeosporioides infection.

The functions of IncRNAs cannot currently be inferred from their sequence or structure, but IncRNAs can function in trans mode to target gene loci distant from where the IncRNAs are transcribed [58]. In F26, a total of 5 modules related to disease resistance were obtained by WGCNA during the infection of C. gloeosporioides. Many target genes of IncRNAs in these modules are enriched in plant immune related items and pathways, such as “activation of innate immune response”, “activation of immune response” in MElightyellow module, “defense response to bacterium, incompatible interaction” in MEbrown2 module, “defense response” and “immune system process” in MEwhite module. These results suggest that these genes may play important roles in the process of resistance to C. gloeosporioides of walnut fruit bracts. Phytohormones are known to be important in the regulation of defense responses in plants [59-61]. Plants can exhibit systemic acquired resistance through the salicylic acid (SA) / jasmonic acid (JA)-mediated signaling network [62-65]. In our study, a total of 32 genes were identified in the significantly enriched KEGG pathway “Plant hormone signal transduction”. Meanwhile, there are 3 and 5 genes enriched in “jasmonic acid mediated signaling pathway” and “response to jasmonic acid” respectively. We also showed that some genes were enriched in “auxin-activated signaling pathway” and “cellular response to auxin stimulus” at 24 hpi. Therefore, auxin may play a role in the resistance of walnut bracts to C. gloeosporioides. In addition, our result showed that the phenylpropanoid biosynthesis was one of the most significantly enriched pathways in the process of resistance to C. gloeosporioides of walnut fruit bracts. In this pathway, phenylalanine ammonium lyase (PAL) is the key regulatory enzyme in

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altering the biosynthesis and accumulation of flavonoids and lignin [66]. Lignin plays a structural role in the secondary cell walls formation [67], and flavonoids mediate plants against UV radiation and act as a visual signal for attracting pollinators [68,69]. In Caragana korshinskii, C. korshinskii adjusts its phenylpropanoid biosynthesis process to water-deficit environments and activates PAL by drought stress [70].

During long-term evolutionary interactions with plants, several pathogens successfully cause effector-triggered susceptibility response (ETS) by producing a number of effectors. Simultaneously, plants have evolved R genes that recognize these effectors and function through highly specific interactions between effectors and their corresponding nucleotide-binding site and leucine-rich repeat (NB-LRR) class receptors [71]. In tomato, IncRNA23468 reportedly increases the expression of the NBS-LRR target genes (encoding R proteins), resulting in enhanced resistance to Pinestans [49]. In the current study, we detected 10 R genes among the target genes of 96 hub IncRNAs. During the infection of C. gloeosporioides on the walnut fruit bracts, the results of RNA-seq showed that the expression of R genes (XM_018950446.1, XM_018989953.1 and XM_018961957.1 in MElightyellow module, XM_018962714.1 in MEbrown2 module, XM_018965430.1, XM_018973294.1 and XM_018954279.1 in MElwhite module) were up-regulated at 24hpi, 48hpi and 72hpi respectively, and expression of the highly connected IncRNAs (MSTRG.11713.16, MSTRG.146621.3 and MSTRG.136680.2 in MElightyellow module, MSTRG.123346.3 in MEbrown2 module, MSTRG.94840.7, MSTRG.18285.3 and MSTRG.45846.2 in MElwhite module) had the same trends (Additional file 14 Table_S14). These findings imply that IncRNAs may help mediate the disease resistance of walnut fruit bracts through the target R genes. The specific interaction between IncRNAs and R gene needs further verification. The expression levels of five hub IncRNAs (MSTRG13585, MSTRG11713, MSTRG152205, MSTRG112028, and MSTRG62751) and their target genes were further confirmed by qPCR, the results of which were consistent with the RNA-seq data. The data presented here provides researchers with the biological basis for future investigations of the mechanism underlying the disease resistance of walnut fruit bracts.

Conclusions

In this study we generated the expression profile of IncRNA in anthracnose-resistant F26 and anthracnose-susceptible F423 at five times. Compared with F423, a total of 14525 DELs were identified, including 10645 upregulated IncRNAs and 3846 downregulated IncRNAs in F26. Bioinformatic analysis showed that the target genes of upregulated IncRNAs were enriched in immune-related processes, plant hormone signal transduction, phenylpropanoid biosynthesis and other pathways during the infection of C. gloeosporioides. Hub IncRNAs with high connectivity to disease resistant genes were predicted. These results contribute to our understanding of the potential mechanism by which IncRNAs involved in C. gloeosporioides resistance and will facilitate the functional verification of the IncRNA in the future.

Methods

Plant materials and fungal isolates
The scions of walnut seedling tree B26 was provided by walnut specialized farmers’ cooperative of Dongliugang village, Baishi Town, Wenshang County, Shandong Province, China (35°46′56.2″N, 116°40′30.8″E). The 4-23 walnut tree was from F1 progeny of an intraspecific cross between walnut cultivar ‘Yuan Lin’ (susceptible to anthracnose)× ‘Qing Lin’ (resistant to anthracnose) which was carried out by ourselves in 2002. The plant materials were conserved by patch budding onto walnut seedling rootstock at the Forestry Experimental Station of Shandong Agricultural University, Tai’an, Shandong Province, China (36°10′ 19.2″N, 117°09′ 1.3″E) in late May 2009. In 2015-2017, we evaluated the anthracnose resistance of each plant for three consecutive years followed by previously described [8,14], and it was found that B26 clone was highly resistant to anthracnose in fruit bract, and the 4-23 clone was highly susceptible to anthracnose in fruit bract. The fruits of B26 clone (i.e., F26) and 4-23 clone (i.e., F423) were used as experimental materials. The voucher specimen of F26 and F423 had been deposited to our lab but not to any publicly available herbarium. We didn't use wild plants in this study and according to national and local legislation, no specific permission was required to collect these plants. C.gloeosporioides m9 isolates (GenBank ID: GU597322) used in this study were maintained by our group.

**Fungal pathogen inoculation of walnut fruits**

*Colletotrichum gloeosporioides* was cultivated on potato dextrose agar medium for 5–7 days at 28°C. To prepare conidial suspensions, the colonies were washed with sterile distilled water containing 0.05% (v/v) Tween 80, passed through a filter (40–100 μm pores), quantified with a hemocytometer, and diluted with sterile distilled water to 105–106 conidia/ml [0.001% (v/v) Tween 80 final concentration]. Healthy fruits from the east-, south-, and west-facing parts of each tree were collected in mid-June and disinfected with 0.6% sodium hypochlorite and rinsed with sterile water. The punch inoculation of the detached walnut fruits was completed as previously described [8]. Based on anatomical changes to the infected walnut fruit bract, samples of the inoculation site were collected at 0, 24, 48, and 72 hpi, 0 hpi as a control. Additionally, distal uninoculated tissue was collected at 120 hpi. Take two independent samples as biological replicates at each infection time (Additional file 1: Table S1). All samples were flash-frozen in liquid nitrogen and stored at −80°C until analyzed.

**Determination of physiological and biochemical data**

The activity of CAT, Chitinase, SOD and contents of H₂O₂, salicylic acid and jasmonic acid at five time points of F26 and F423 were determined according to the instructions on the kit. Each sample was repeated three times. The CAT, Chitinase and SOD activity levels were measured and performed according to kit instructions (Solarbio® cat. No. BC0820) and detected by TU-1901 UV Spectrophotometer (Beijing Purkinje General Instrument Co., Ltd., Beijing). The content of H₂O₂, salicylic acid and jasmonic were detected by the Solebao kit (Solarbio® cat. No. BC3595) with microdetermination.

**RNA extraction, library construction, and sequencing**

Total RNA was extracted from F423 and F26 samples with the Thermo Gene JET Plant RNA Purification Mini Kit (Thermo Fisher Scientific Inc., USA). The purity and concentration of the extracted RNA were...
determined with the NanoDrop2000 spectrophotometer (Thermo Fisher Scientific Inc.) (OD_{260/280} \geq 1.8, OD_{260/230} \geq 1.5, and concentration > 40 ng/µl). The RNA integrity was assessed by agarose gel electrophoresis. Ribosomal RNA was removed with the Ribo-Zero™ Magnetic Kit (Epicentre) and the remaining RNA (polyA+ and polyA−) was recovered. The RNA was randomly fragmented to approximately 200-bp sequences in Fragmentation Buffer (Thermo Fisher Scientific Inc.) and then used as the template to synthesize first-strand cDNA with random hexamers. The second cDNA strand was synthesized with dNTPs, RNaseH, and DNA polymerase I. The overhanging ends were filled in with T4 DNA polymerase and Klenow DNA polymerase to generate blunt ends, after which the A base was added to the 3’ end and the fragment was ligated to a linker. The AMPureXP beads were used for selecting fragments. The second cDNA strand containing U was degraded with the USER enzyme, after which a sequencing library was obtained by PCR amplification. A total of 20 sequencing libraries were constructed. The Qubit 2.0 DNA Broad Range Assay (Invitrogen, USA) was used for a preliminary quantification. The sequencing library inserts were detected with the Agilent 2100 Bioanalyzer. Finally, the effective library concentrations (> 2 nM) were accurately quantified by qPCR. Paired-end sequencing (2 × 150 bp) was completed in KeGene Science & Technology Co. Ltd. (Shandong, China) with an Illumina HiSeq 4000 platform.

Read mapping and transcriptome assembly

The quality of the raw sequencing data was checked with FASTQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Adapters and low-quality tags in the raw data were eliminated. Ribosomal RNA data were also removed. The remaining clean reads for the 20 cDNA libraries were combined and mapped to the *J. regia* genome sequence (https://www.ncbi.nlm.nih.gov/genome/?term=Juglans%20regia) with the HISAT program (version 0.11.5) (parameter setting: -rna-strandness RF) [31]. To construct transcriptomes, the mapped reads were assembled with StringTie (version 1.3.1) [32]. After combining the StringTie results for each sample with StringTie-Merge, the read counts were calculated for transcripts with bedtools (version 2.27.1) (bedtools.readthedocs.org /en/latest/#) [33].

Identification of lncRNAs

To obtain the potential long non-coding RNAs, based on all the assembled transcripts, we have firstly excluded the known transcripts according to the class code “=”. Then the remaining transcripts were used to remove the potential protein coding transcripts, miRNA-like, and other transcript types via blasting against the database of Rfam, Refseq, Uniprot, miRbase, and Pfam. Finally, the remaining transcripts were employed for coding potential prediction by using FEELnc tool. First, the FEELnc filter was used to remove short transcripts (default 200nt) and assess single-exon transcripts [34]. The FEELnc codpot predictors were used to calculate a coding potential score. The assembled sequences were used for reconstructing the transcriptome. Finally, RNAs longer than 200nt and derived from \( \geq 2 \) exons, with an ORF coverage < 50% and a potential coding score < 0.5 were designated as lncRNAs [35].

Classification of lncRNAs
The lncRNAs were analyzed regarding their corresponding positions in the reference genome and the positional relationships between lncRNAs and partner RNAs based on 10,000–100,000 fragments. The lncRNAs were then divided into genic lncRNAs (overlapping partner RNAs) and intergenic lncRNAs (lincRNAs). The genic lncRNAs were further divided as overlapping, containing, or nested subtypes. Intergenic lncRNAs were divided as divergent, convergent, and same strand subtypes.

**Analysis of differential expression patterns**

Genes differentially expressed between the disease-resistant and susceptible fruits at five infection stages were analyzed with DESeq2 (version 1.22.1) [36]. After assessing the significance of any differences, the genes with a $p$ value $\leq 0.05$ and a $|\log_2 \text{foldchange}| \geq 1$ were designated as differentially expressed genes.

**Quantitative real-time PCR**

Total RNA samples extracted from walnut fruits at individual infection stages were analyzed by qPCR. Briefly, first-strand cDNA was obtained with the TransScript One-Step gDNA Removal and cDNA Synthesis SuperMix for qPCR (Transgen, China). The IncRNA expression level was quantified with the TransStart Tip Green qPCR SuperMix (Transgen) and the CFX Connect Real-Time System (Bio-Rad). The qPCR program was as follows: 95 °C for 30 s; 40 cycles of 95 °C for 10 s and 60 °C for 30 s. For a melting curve analysis, the temperature was increased from 70 °C to 95 °C (0.5 °C/5 s). All samples were analyzed in triplicate. The 18S rRNA gene was used as a housekeeping gene. The cycle threshold ($Ct$) $2^{\Delta \Delta Ct}$ method (Software IQ5 2.0) [37] was used for the relative quantification of mRNAs. The primers used for RT-qPCR were designed with Beacon Designer 7 software and were synthesized by Sangon Biotech (Shanghai, China; Supplementary Table S14).

**Prediction of IncRNA functions based on co-expression**

Co-expression modules were generated with the WGCNA package (version 1.67) as previously described [38] (http://lab.genetics.ucla.edu/horvath/Coexpression Network/). The IncRNAs and mRNAs that were not detected in at least one infection stage were not considered. In this analysis, the soft thresholding power was set to 12, after which the adjacency function was used to construct the adjacency matrix. A topological overlap measure map was constructed based on the adjacency matrix to calculate the similarity matrix of the IncRNA and mRNA expression between different nodes. The IncRNAs and mRNAs were hierarchically clustered based on the algorithm. To generate a number of clusters, modules were defined after eliminating or combining branches. The co-expression module dynamic shear tree parameters were determined as described by Gerttula [39]. The minimum number of genes was set to 30, the split sensitivity (deep Split) was set to 2, and the other settings were software default parameters. The module was related to the trait, and the correlation matrix between the module and the trait was calculated. The module with the highest correlation coefficient and the smallest $p$ value was designated as the module most relevant to the trait. In this study, a significantly correlated module was identified based on a correlation coefficient ($r$) $\geq 0.8$ [35] and $p < 0.05$. The co-expression networks of IncRNAs and
hub IncRNAs in highly correlated modules were generated with the Cytoscape software (version 3.7.1) [40].

**Functional enrichment analysis**

The genes targeted by IncRNAs were functionally annotated based on the GO and KEGG pathway (http://www.genome.jp/kegg/) databases. The KOBAS program (version 2.0) was used to determine the significantly enriched KEGG pathways among the target genes [41]. According to the operation requirements of KOBAS 2.0, All data files were written with a parser. The gene-term mapping can be retrieved by parsing the raw data files for each pathway. The gene annotation and gene-ID relations were retrieved from KEGG Genes and BioMart. We mapped the genes in all databases to KEGG GENES and KEGG ORTHOLOGY (KO). The gene-pathway and is stored in our backend SQL relational database. The FASTA protein sequence files were preprocessed for BLAST [42].

**Abbreviations**

- LncRNAs: long non-coding RNAs
- DELs: different expressed IncRNAs
- WGCNA: Weighted Gene Coexpression Network Analysis
- hpi: hours post inoculation
- eTM: endogenous target mimics
- ceRNA: competing endogenous RNA
- GO: Gene Ontology
- KEGG: Kyoto Encyclopedia of Genes and Genomes
- ETS: effector-triggered susceptibility response
- NB-LRR: nucleotide-bindingsite and leucine-rich repeat
- ROS: reactive oxygen species
- SA: salicylic acid
- JA: jasmonic acid

**Declarations**

**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and materials**

The data sets are included within the article and its Additional files. The raw sequencing data were deposited in NCBI Sequence Read Archive under the accession number GSE147083 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE147083).

**Competing interests**

The authors declare that they have no conflicts of interest.

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**Authors’ contributions**

KQY conceived the idea and revised the manuscript. SF and HF conducted meta-analysis, drew figures and drafted the manuscript. XL collected the experimental materials. QW and YD helped in drawing figures and drafting the manuscript. All authors listed have made direct and substantial efforts for improving the manuscript and approved the final version.

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