**RcTGA1 and glucosinolate biosynthesis pathway involvement in the defence of rose against the necrotrophic fungus Botrytis cinerea**

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**Abstract**

**Background:** Rose is an important economic crop in horticulture. However, its field growth and postharvest quality are negatively affected by grey mould disease caused by Botrytis cinerea. However, it is unclear how rose plants defend themselves against this fungal pathogen. Here, we used transcriptomic, metabolomic and VIGS analyses to explore the mechanism of resistance to *Botrytis cinerea*.

**Result:** In this study, a protein activity analysis revealed a significant increase in defence enzyme activities in infected plants. RNA-Seq of plants infected for 0 h, 36 h, 60 h and 72 h produced a total of 54 GB of clean reads. Among these reads, 3990, 5995 and 8683 differentially expressed genes (DEGs) were found in CK vs. T36, CK vs. T60 and CK vs. T72, respectively. Gene annotation and cluster analysis of the DEGs revealed a variety of defence responses to *Botrytis cinerea* infection, including resistance (R) proteins, MAPK cascade reactions, plant hormone signal transduction pathways, plant-pathogen interaction pathways, Ca2+ and disease resistance-related genes. qPCR verification showed the reliability of the transcriptome data. The PTRV2-RcTGA1-infected plant material showed improved susceptibility of rose to *Botrytis cinerea*. A total of 635 metabolites were detected in all samples, which could be divided into 29 groups. Metabonomic data showed that a total of 59, 78 and 74 DEMs were obtained for T36, T60 and T72 (*Botrytis cinerea* inoculated rose flowers at 36 h; T60: *Botrytis cinerea* inoculated rose flowers at 60 h; T72: *Botrytis cinerea* inoculated rose flowers at 72 h) compared to CK, respectively. A variety of secondary metabolites are related to biological disease resistance, including tannins, amino acids and derivatives, and alkaloids, among others; they were significantly increased and enriched in phenylpropanoid biosynthesis, glucosinolates and other disease resistance pathways. This study provides a theoretical basis for breeding new cultivars that are resistant to *Botrytis cinerea*.

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Background

Rose is one of the four cut flowers in the world with high ornamental and economic value. Black spot, aphid, and grey mould are three major diseases affecting rose production along with insect pests. Grey mould mainly occurs in the late flowering or postharvest period of roses. This will cause an approximate 15% ~ 40% reduction in production, and postharvest corruption will lead to serious economic losses to the cut rose market.

Grey mould of rose is caused by Botrytis cinerea, which belongs to Sclerotinia [1, 2]. Botrytis c. is a typical necrotrophic pathogenic fungus that infects host cells by regulating the programmed death pathway [3]. It usually infects the plant tissue at an early stage and stays for a long time. When the environment is suitable or the host physiological changes, it will burst out suddenly, leading to deterioration and decay of plant tissue. Chemical methods are mainly used to control Botrytis c., such as imidazole, dicarboximides, and anilinopyrimidines. However, due to the long-term use of chemical drugs, Botrytis c. has developed tolerance to them. The worse the effect, the larger the drug dosage. It not only increases the economic cost but also aggravates environmental pollution. Therefore, it is urgent to find an effective method to enhance the resistance of Chinese rose to fungi. With the development of biotechnology, whole-genome sequencing of many plant species has been completed and has been applied to study plant growth, environmental interactions, and metabolism, among others. In recent years, transcriptomics and metabolomics have been used to explore the defense mechanisms of plant diseases and insect pests, such as tea, maize, and sesame [4–6], providing a new idea and method to explore the mechanism of resistance of roses to Botrytis c. infection.

Plants have complex defense mechanisms against Botrytis c., including the MAPK cascade reaction, plant hormone signal transduction pathway, and cAMP signal pathway [7]. Silencing the MAPK-WRKY transcription factor gene increased the susceptibility of tobacco to Botrytis c., indicating that the MAPK cascade was involved in the immune response of tobacco to fungi [8]. MPK3 and MPK6 can enhance the resistance of Arabidopsis to Botrytis c. by inducing the expression of the glip1 gene [9]. Lu et al. identified bZIP transcription factors from the strawberry whole genome and found that the expression of FvbZIP46 was significantly increased by Botrytis c. Transient overexpression and gene silencing of FvbZIP46 indicated that FvbZIP46 was involved in strawberry defense against Botrytis c. [10]. Plant hormone signalling and transcription factors play an important role in plant resistance to Botrytis c. infection. Hu et al. found that exogenous application of N-decanoyl-homoserine lactone enhanced tomato resistance to Botrytis c. by activating JA biosynthesis and signal transduction in tomato [11]. Liu et al. found that BR signal transduction pathway-related genes were upregulated after infection with Botrytis c., and exogenous BR enhanced its defense response to Botrytis c., indicating that BR was involved in the resistance of rose to Botrytis c. [12]. Exogenous 2,4-Epibrassinolide improves the resistance of grapes to Botrytis c. by inhibiting spore germination of Botrytis c. and enhancing the sensitization mechanism of grape fruit [13, 14]. In addition, some microRNAs are also involved in plant defence against Botrytis c. Nie et al. found that overexpressed miR825 and miR825 * improved the susceptibility of Arabidopsis to Botrytis c. In contrast, silencing miR825 and miR825 * enhanced the resistance of Arabidopsis to Botrytis c., indicating that mir825 and mir825 * play a negative regulatory role in Arabidopsis defence against Botrytis c. [15].

To explore the defence mechanism of rose against Botrytis c., Old Blush was used as the experimental material, and the whole genome was published in 2018 (https://lipm-browsers.toulouse.inra.fr/pub/RchiOBHmV2/). In this study, we measured the antioxidant enzyme activities of plants infected with Botrytis c. We then combined transcriptomic and metabolomic analyses to compare the gene expression and metabolomic profiles between CK and different infection time points (T36, T60, T72). These data provide significant information for the future breeding of rose resistance to grey mould.

Results

Effects of Botrytis c. on the phenotypic and biochemistry of old blush

We observed and recorded the phenotypic changes in Old Blush to investigate the effect of Botrytis c. infection on Old Blush growth. After 24 h of inoculation, the
petals developed typical disease spots, which continued to grow larger to form apparent necrotic symptoms after 72 h of inoculation. In contrast, this pathogenic response was not observed in control petals treated with the mock control (Fig. 1a). During long-term evolution, plants have developed sophisticated biochemical and physiological mechanisms to adapt and resist various pathogenic bacteria. PPO, Glu and CHT play an important role in plant resistance to microbial infection. Therefore, enzymatic assays were performed to examine whether PPO, Glu and CHT were responsive to *Botrytis c.* infection. The results showed that the activities of these enzymes significantly differed among the treatments. The CHT activities of the 36 h, 60 h, 72 h petals increased by 26.13, 51.35 and 55.86%, respectively; the PPO activities at 36 h, 60 h, 72 h increased by 5.72, 41.87 and 59.66%, respectively; the GLU activities at 36 h, 60 h, 72 h increased by 78.65, 86.86 and 88.92%, respectively (Fig. 1b). These data indicate that the defence system in Old Blush is responsive to *Botrytis c.* infection.

**Overview of Transcriptomic analysis**

To understand the difference in gene expression between *Botrytis c.*-infected petals and mock petals, we next conducted an RNA-Seq analysis. A total of approximately 54 GB clean reads were generated from twelve biological samples, including nine infected and three control samples. The average Q20 value of the raw reads

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**Fig. 1** Changes in phenotype (a) after Old Blush was inoculated with sterile distilled water (SDW, mock) or *Botrytis c.* (b) defense enzyme activities. All statistical analyses were performed using Student's t-test; *, \( P < 0.05 \); **, \( P < 0.01 \); hpi hours post-inoculation
was 95.88%, indicating high-quality reads. Approximately 82% of the reads were mapped to the reference genome sequences obtained by Trinity splicing. To obtain a comprehensive view of the gene expression profile associated with the response of Old Blush to Botrytis c. infection, we used DESeq2 to identify DEGs. Based on the filtering parameters of FDR < 0.05 and |log2FC| > 1, the expressions of 3990 (2349 upregulated, 1641 downregulated), 5995 (3621 upregulated, 2374 downregulated) and 8683 (4164 upregulated, 4518 downregulated) genes were found to differ significantly in CK-VS-T36, CK-VS-T60 and CK-VS-T72, respectively. In addition, 529, 6803 and 6244 DEGs were identified in the comparison of T36 vs. T60, T60 vs. T72 and T36 vs. T72 (Fig. 2).

To understand the functions of the DEGs associated with Botrytis c. infection, those DEGs were annotated using GOseq. This annotation resulted in three major categories: biological processes, cellular components, and molecular functions. Most of the DEGs were enriched in the ‘response to external stimulus’, ‘response to stimulus’, ‘response to external biotic stimulus’, ‘response to stress’, ‘secondary metabolic process’, ‘flavonoid metabolic process’ and other functional categories (Fig. 3a, b, c). The results showed that Botrytis c. infection activated rose resistance.

To better understand the main pathways activated by Botrytis c. infection, we conducted a Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis of the DEGs. Between the CK and T36 libraries, 728 DEGs were assigned to 127 KEGG pathways. Between the CK and T60 libraries, 1102 DEGs were assigned to 130 KEGG pathways. Between the CK and T72 libraries, 1607 DEGs were assigned to 134 KEGG pathways. Among these DEGs, genes involved in ‘metabolic pathways’ were the most abundant, followed by genes involved in ‘biosynthesis of secondary metabolites’. There were some important disease resistance pathways involved in the ‘MAPK signalling pathway – plant’, ‘phenylpropanoid biosynthesis’, ‘plant hormone signal transduction’, and ‘glutathione metabolism’ (Fig. 4a, b, c). This result indicated that a series of resistance pathways were activated in rose after infection with Botrytis c.

To understand the dynamics of differential gene expression by Botrytis c. infection, 12,842 DEG clustering analyses of expression patterns in different periods were performed by the Short Time-series Expression Miner. Through this analysis, the expression dynamics of all the DEGs at four periods could be clustered into 20 expression patterns, among which 10,334 DEGs showed 7 significant clustering patterns (P < 0.05). Compared with 0 h, the whole infection process included three downregulated expression patterns (modes 0, 9, 2) and four upregulated expression patterns (modes 19, 16, 10, 17). Among them, the number of downregulated genes in modes 9 and 0 was greater than that in mode 3. The number of upregulated genes in modes 17 and 19 was greater than that in modes 16 and 10 (Fig. 5a).

The downregulated genes with the most highly represented pathways were ‘metabolic pathways’ and ‘biosynthesis of secondary metabolites’. This also includes ‘starch and sucrose metabolism’, ‘porphyrin and chlorophyll metabolism’, ‘photosynthesis’, ‘glyoxylate and dicarboxylate metabolism’, ‘fatty acid biosynthesis’ and other plant development processes (Fig. 5c). These metabolic pathways are mainly involved in primary metabolism in plants, indicating that the expression of genes involved in primary metabolism decreased and the process of primary metabolism weakened after infection with Botrytis c.

In contrast, the upregulated genes were the most highly represented pathways in addition to ‘metabolic pathways’, ‘biosynthesis of secondary metabolites’, ‘phenylpropanoid biosynthesis’, 'plant pathogen interaction',

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**Fig. 2** Venn map of differentially expressed genes. (CK) non-infected rose flowers, (T36) Botrytis c. inoculated rose flowers at 36 h, (T60) Botrytis c. inoculated rose flowers at 60 h, (T72) Botrytis c. inoculated rose flowers at 72 h.
Fig. 3 GO analysis based on DEGs in CK VS T36 (a), CK VS T60 (b) and CK VS T72 (c). (CK) non-infected rose flowers, (T36) *Botrytis c.* inoculated rose flowers at 36 h, (T60) *Botrytis c.* inoculated rose flowers at 60 h, (T72) *Botrytis c.* inoculated rose flowers at 72 h
Fig. 4 KEGG analysis based on DEGs in CK VS T36 (a), CK VS T60 (b) and CK VS T72 (c). (CK) non-infected rose flowers, (T36) Botrytis c. inoculated rose flowers at 36 h, (T60) Botrytis c. inoculated rose flowers at 60 h, (T72) Botrytis c. inoculated rose flowers at 72 h.
Fig. 5 Analysis of DEGs. a Cluster analysis of DEGs; (b) KEGG enrichment Bubble Diagrams of profile 17 and profile 19 upregulated DEGs. c KEGG enrichment Bubble Diagrams of profile 0 and profile 9 downregulated DEGs.
‘MAPK signalling pathway-plant’, ‘plant hormone signal transduction’, ‘alpha-linolenic acid metabolism’, ‘glutathione metabolism’ and ‘monoterpenoid biosynthesis’ (Fig. 5b). These metabolic pathways are mainly involved in the secondary metabolism of plants, indicating that the expression of genes involved in secondary metabolite synthesis is increased to enhance the resistance of rose to Botrytis c. Taken together, the statistical results above indicate that rose starts its own defence mechanism in response to Botrytis c. stress by balancing primary and secondary metabolism.

Heatmaps of DEG subclusters were developed to better understand the key DEGs associated with the resistance of Old Blush to Botrytis c. The resulting heatmaps showed DEGs involved in plant-pathogen interactions. Based on their functional annotation, these genes included 2 PAMP-triggered immunity (PTI) genes, 3 R genes, 1 reactive oxygen species (ROS) metabolic pathway gene, 4 calmodulin genes, 3 calcium-binding protein genes, 6 mitogen-activated protein kinase (MAPK) signalling pathway genes, and 5 DEGs involved in the SA response pathway, including 1 NPR gene, 2 PRs, and 2 transcription factor TGA genes. In addition, 21 defence enzyme genes, including one PAL, 8 GLUs, 7 CHTs, 2 PPOs, 2 GPXs and one SOD unigene, are shown in the heatmap (Fig. 6). This indicated the rose response to Botrytis c. by activating the expression of signal transduction pathway genes, transcription factors and disease resistance genes.

Validation of candidate DEGs with qPCR analysis
To validate the reliability of DEGs obtained from RNA-Seq analyses, the expression levels of 12 candidate genes were analysed using qPCR. These genes included 3 TGAs (RcTGA3/6/7), 4 defence enzyme genes (RcPAL1, RcPYL, RcCHI, and RcEP3), 2 plant hormone signalling pathway genes (RcEIN3 and RcBKII), one pathogenesis-related gene (RcHEL), and 2 MAPK signalling pathway genes (RcMKK9 and RcSNRK2). The correlation coefficients (r) between the RNA-Seq and qPCR results were

![Fig. 6](image-url)
Fig. 7 (See legend on next page.)
Silencing of RcTGA1 enhances susceptibility to Botrytis c.

TGA1-TGA7 can interact with NPR1, a key regulator of the salicylic acid signalling pathway, to regulate plant disease resistance in Arabidopsis thaliana. Based on NCBI BLASTx, we identified and isolated the RcTGA1 (gene ID: XM024330029.1) gene located in the SA resistance pathway. To detect the expression pattern of RcTGA1 in response to Botrytis c., we inoculated Botrytis c. on the flowers of Old Blush and detected the expression characteristics of RcTGA1 in flowers using qPCR (Fig. 8d). The results showed that the expression level of RcTGA1 increased significantly after 4 h of infection. It continued to grow within 72 h after infection with Botrytis c. This result suggested that RcTGA1 might be involved in the response of rose to Botrytis c.

It has been previously demonstrated that TRV-mediated VIGS can be used in rose as a tool for the functional analysis of genes involved in flower development and the determination of petal colour and fragrance [16, 17]. To determine whether the increased expression of genes of the SA response pathway is associated with the tolerance of Old Blush to Botrytis c., we cloned fragments of RcTGA1 (an upregulated DEG encoding a transcription cofactor related to the SA signalling pathway) that were 250 bp in length into the TRV vector and inoculated rose petal discs with these fragments via vacuum infiltration. At one-week post-infiltration, we subsequently challenged TRV-RcTGA1- and TRV-inoculated rose petal discs with Botrytis c. The TRV-RcTGA1-inoculated plants showed severely compromised resistance, as evidenced by significantly increased lesion size (Fig. 8a, b). We further used qPCR to detect the silencing efficiency in the petals. The results showed the most significant decrease in RcTGA1 expression upon infiltration with TRV-RcTGA1 (Fig. 8c). These results indicated that RcTGA1 is essential for Botrytis c. resistance in rose.

Overview of Metabonomic analysis

To understand the changes in metabolites and the possible defence mechanisms of Old Blush infected by Botrytis c., metabolite profiling analysis of Old Blush flower samples (CK, T36, T60, T72) was performed. A total of 635 metabolites were detected in all samples, which could be divided into 29 groups (Table S1). Principal component analysis (PCA) showed that the repeatability of different treatments was good. Orthogonal partial least squares discriminant analysis (OPLS-DA) showed that the results of OPLS-DA analysis could be used for subsequent model tests and differential metabolite analysis. Between the CK and T36 treatments, 373 were up-regulated, and 262 were downregulated. Between the CK and T60 treatments, 397 were upregulated, and 238 were downregulated. Between the CK and T72 treatments, 406 were upregulated, and 229 were downregulated. The levels of piperidine, γ-aminobutyric acid, L-histidine, 5-O-p-coumaroylquinic acid, chlorogenic acid, terminal acid, 2-hydroxyoleanolic acid, 2α-hydroxyursolic acid, and 3,4-digalloylshikimic acid increased significantly with the extension of infection time. This indicated that these secondary metabolites were involved in the resistance response of rose to Botrytis c.

To identify the main pathways that Old Blush uses to respond to Botrytis c., we mapped the differentially expressed metabolites to KEGG biological pathways. Thirty-two significantly differentially expressed metabolites between the CK and T36 treatments were assigned to 50 KEGG pathways, including ‘aminoacyl-tRNA biosynthesis’, ‘valine, leucine and isoleucine biosynthesis’, and ‘biosynthesis of amino acids’. Thirty-eight significantly differentially expressed metabolites between the CK and T60 treatments were assigned to 61 KEGG pathways, including ‘aminoacyl-tRNA biosynthesis’, ‘2-oxocarboxylic acid metabolism’, and ‘glucosinolate biosynthesis’. Thirty-seven significantly differentially expressed metabolites between the CK and T72 treatments were assigned to 65 KEGG pathways, including ‘biosynthesis of antibiotics’, ‘valine, leucine and isoleucine biosynthesis’, ‘alanine, aspartate’ and ‘glutamate metabolism’ (Table S2). The results showed that the metabolic pathways related to disease resistance were significantly enriched, indicating that the defence mechanism of rose was activated after infection by Botrytis c.

Glucosinolate biosynthesis

The glucosinolate biosynthesis pathway is involved in the defence responses of many plants to pathogens. In our research, the key metabolites of the glucosinolate biosynthesis pathway, including L-valine, L-isoleucine, and L-leucine, exhibited different levels in different treatments. Compared with that in the CK, their level in T36 increased by factors of 1.78, 1.09, and 1.08, respectively; in T60, they increased by factors of 2, 2.3, and 2.4,
respectively, and in T72, they increased by factors of 1.6, 2.0, and 2.0, respectively. These results indicate that the glucosinolate biosynthesis pathway may be positively involved in the interaction between rose and *Botrytis c.*

**Co-joint analysis**

The co-joint KEGG enrichment analysis showed 85 co-mapped pathways. There were 46, 56, and 60 co-mapping pathways between CK-VS-T36, CK-VS-T60, and CK-VS-T72 and their metabolites. Interestingly, of these co-
mapped pathways, ‘metabolic pathways’, ‘biosynthesis of secondary metabolites’, ‘phenylpropanoid biosynthesis’, ‘flavonoid biosynthesis’, ‘tropane, piperidine and pyridine alkaloid biosynthesis’, ‘tyrosine metabolism’, ‘pentose and glucoronate interconversions’, ‘cyanoamino acid metabolism’, and ‘ascorbate and aldarate metabolism’ were their common pathways of significant enrichment (Fig. 9a, b, c). This indicates that rose responds to pathogen infection by coordinating the primary and secondary metabolic pathways. Based on the O2PLS model, the combined analysis of transcriptomics and metabolomics data showed that the model was reliable (R2 > 0.85). The Pearson correlation coefficients showed that the differential expression patterns of DEGs and metabolites were consistent. The correlations between the top 250 DEGs and their metabolites were further selected and are represented as a heat map (Fig. s1).

Discussion
RNA-seq study for disease resistance
As an important economic horticultural plant, rose infected with Botrytis c. during its growth or after harvest will cause great economic losses to farmers [18]. To understand the mechanism of Old Blush tolerance to Botrytis c., RNA-Seq and metabolomics were employed to analyses DEGs and DEMs in Old Blush at different time points after inoculation with Botrytis c. In our study, a large number of DEGs and DEMs were putatively related to the response of Old Blush to necrotrophic Botrytis c. Numerous DEGs are involved in various defense responses to Botrytis c., including redox-related genes, Ca2+, MAPK signalling pathway-related genes and plant hormone signal transduction pathway-related genes. All of these transcriptomic data indicate that multiple processes in Old Blush are associated with plant defence against pathogens, which is in line with the fact that the plants have evolved a complex defence mechanism [19–22]. When infected by pathogens, the plant recognizes the PAMP secreted by Botrytis c. through the pattern recognition receptor on the cell surface and triggers a series of cellular reactions, including ROS production, changes in cytoplasmic ion flux, calcium-dependent protein and MAPK cascade activation. In the process of preventing pathogen infection, plants encode the R protein to activate effector-triggered immunity and produce a hypersensitive reaction (HR) at the infected site, which promotes cell death [23]. In our study, FLS2 and ROS pathway genes were significantly upregulated after Botrytis c. infection. This result revealed that Botrytis c. infection also triggered effector-triggered immunity of Old Blush.

Resistance of JA and SA signalling pathways to Botrytis c.
Plant hormones, such as salicylic acid and jasmonic acid, are involved in the plant defence response to plant diseases and insect pests [24, 25]. Ren et al. silenced the key JA synthesis gene AOS and JA coreceptor gene COI1, which increased the susceptibility of Old Blush to Botrytis c., indicating that the JA pathway plays an important role in resisting Botrytis c. to rose [12]. Liu et al. analysed transcriptome data and found that BR hormone signal transduction pathway-related genes were upregulated after rose inoculation. Exogenous BR enhanced the resistance of rose petals to Botrytis c. In this study, Old Blush was inoculated with Botrytis c., a typical necrotrophic pathogen. The analysis of DEGs revealed the upregulation of JA signalling pathway genes, including RcJAZ, RcLOXs, and RcCOI1. DEM analysis showed that the content of α-linolenic acid* related to JA synthesis also increased significantly, which was consistent with the results of Ren. SA activates NPR1 expression causing Arabidopsis to gain systemic resistance and improve its disease resistance. SA application to tomato leaves significantly increased the expression level of the SA marker gene PR1 (pathogenesis-related protein 1) and enhanced its resistance to Botrytis c. In our study, the expression of the SA pathway genes RcTGA1s and PR1s was upregulated in Old Blush after inoculation with Botrytis c. This result suggested that SA may be involved in the resistance of Old Blush to Botrytis c. in the early stage.

Functional verification of RcTGA1
Tianyi et al. [26] found that overexpression of MdTGA2.1 can complement the SA-sensitive phenotype of TGA2/5/6 in Arabidopsis thaliana. Van Verk et al. [27] showed that TGA2.2 could specifically interact with NtWRKY12 and regulate the expression of PR-1a in vivo and in vitro. All of these studies indicate that TGAs are broad-spectrum resistance genes in the salicylic acid resistance pathway. In our study, the expression of RcTGA1 continued to increase significantly after infection with Botrytis c. RcTGA1 was silenced in rose petals and then inoculated with Botrytis c., and the lesion diameter on the petal disc was twice that of the control. Taken together, RcTGA1 positively regulates rose tolerance to Botrytis c.

Metabolomics study for disease resistance
Secondary metabolites determine the colour, smell and taste of plants. They are widely involved in plant growth, development, defence or other physiological processes. When plants are infected with pathogens, secondary metabolites are produced to participate in plant disease resistance [28]. Studies have shown that sugar metabolism, a primary metabolic process, affects plant susceptibility and plays a key role in innate defence pathways. The analysis of DEMs showed that compared with CK, the
Fig. 9 Joint KEGG enrichment p-value histogram in CK VS T36 (a), CK VS T60 (b) and CK VS T72 (c). (CK) non-infected rose flowers, (T36) Botrytis c. inoculated rose flowers at 36 h, (T60) Botrytis c. inoculated rose flowers at 60 h, (T72) Botrytis c. inoculated rose flowers at 72 h.
levels of saccharides and alcohols (D-threones, D-glucosamine, inositol, and D-glucose-6-phosphate) significantly decreased, which was consistent with the transcriptome data showing that the expression of genes related to the starch and sugar metabolism pathway was significantly downregulated. It is also known that alkaloids, triterpenoids, tannins and phenolic acids are involved in plant disease resistance. In this study, the content of secondary metabolites related to disease resistance increased, while the content of metabolites used for growth and development decreased. This result indicates that when infected by *Botrytis c.*, the roses increased their contents of disease-resistant metabolites and maintained their own growth and development simultaneously.

**Glucosinolate metabolism**

Glucosinolates have broad antibacterial activity. Research has shown that the glucosinolate metabolism pathway is necessary for Arabidopsis immune pathogens [29]. Stotz et al. [30] showed that the content of glucosinolates in Arabidopsis increased and that the expression of glucosinolate synthesis-related genes was activated by *Sclerotinia sclerotiorum* infection. In our study, when rose was infected by pathogens, the expression of related genes and the content of metabolites increased significantly. Thus, the glucosinolate metabolic pathway may be involved in the resistance of Old Blush to *Botrytis c.*

**Regulation mechanism of resistance to *Botrytis c***

Based on our work, we propose a hypothetical model to explain the resistance of Old Blush to *Botrytis c.* In this model, the plants recognize PAMPs secreted by fungi through their innate recognition receptors, causing a series of cellular responses. At the same time, the plant R protein activates ETI to induce plant hypersensitivity, which leads to cell death and prevents further infection. In addition, the expression of JA synthesis- and signal transduction pathway-related genes *RcJAZ* and *RcLOXs* is activated. The expression of SA-induced disease resistance-related genes, including *RcTGAs* and *RcPR1s*, also increased. The levels of precursors of JA synthesis alpha-linolenic acid and glucosinolate metabolic pathway metabolites, such as L-leucine and L-valine, increased significantly (Fig. 10). Taken together, we speculate that the MAPK-plant signalling pathway, CDPK gene, JA biosynthesis, SA resistance pathway and glucosinolate metabolism pathway are involved in the defence of rose against *Botrytis c.*

**Conclusion**

In conclusion, 54 GB of clean reads were generated through RNA-Seq. A total of 3990, 5995 and 8683 DEGs were obtained in the comparisons of T36h, T60h and T72h, respectively, compared to CK. Functional annotation and cluster analysis of the DEGs showed that a variety of defence responses mediated by R proteins, ROS signalling, Ca2+ signalling, MAPK signalling, and SA signalling were activated in the Old Blush response to *Botrytis c.* *RcTGAI* positively regulates rose resistance to *Botrytis c.* Metabonomic data showed that a total of 59, 78 and 74 DEMs were obtained in T36h, T60h and T72h compared to CK, respectively. A variety of secondary metabolites are related to biological disease resistance, including tannins, amino acids and derivatives, and alkaloids, and they were significantly increased and enriched in phenylpropanoid biosynthesis, glucosinolates and other disease resistance pathways.

**Methods**

**Plant growth and plant infection**

*R. chinensis* Old Blush was grown at the rose germplasm garden of the Flower Research Institute, Yunnan Agriculture Academic Science, Kunming, China. The *Botrytis c.* inoculum was produced by growing strain B05.10 on solid medium (potato dextrose agar; 46 g L−1, dH2O, pH = 5.6) at 22 °C for 10 ~ 14 days. Spore inoculum was prepared by harvesting spores in water, filtering through glass wool to remove the hyphae, and suspending the filtrate in potato dextrose broth (PDB; 24 g per L, dH2O) at 105 conidia mL−1 [31]. Two μL drops of *Botrytis c.* inoculum or PDB (mock) were dropped onto each petal disk. Infected and control disks were individually sampled in a randomized manner from each of the three trays at 36 h, 60 h, and 72 h with three biological repeats for both infected and control treatments at each time point. Petals were immediately frozen in liquid nitrogen at the time of harvesting and stored at −80 °C.

**Measurement of antioxidant enzyme activities**

Frozen flower samples were used to determine the activity of the defense enzymes, including polyphenol oxidase (PPO), chitinase (CHT), and glucan endo-1,3-beta-glucosidase (GLU), and RNA-Seq was conducted. SA and JA contents were determined using an enzyme-linked immune sorbent assay (ELISA) in a facility at the Hanling BiO Company (Kunming, China). This company’s service helped to bind those enzymes (bovine serum albumin) and then produced the corresponding antibodies.

**RNA extraction, library construction and sequencing**

Total RNA was extracted using a Trizol reagent kit (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s protocol. RNA quality was assessed using an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, USA) according to the manufacturer’s protocol. cDNA library construction and sequencing were performed by the BiO Service Platform of Kunming. A total of 39,905, 59,951 and 86,833 clean reads were obtained in the comparisons of T36h, T60h and T72h, respectively, compared to CK. Functional annotation and cluster analysis of the DEGs showed that a variety of defence responses mediated by R proteins, ROS signalling, Ca2+ signalling, MAPK signalling, and SA signalling were activated in the Old Blush response to *Botrytis c.*
Alto, CA, USA) and checked using RNase-free agarose gel electrophoresis. After total RNA was extracted, eukaryotic mRNA was enriched by oligo(dT) beads, while prokaryotic mRNA was enriched by removing rRNA with a Ribo-Zero™ Magnetic Kit (Epicentre, Madison, WI, USA). Then, the enriched mRNA was fragmented into short fragments using a fragmentation buffer and reverse transcribed into cDNA with random primers. Second-strand cDNA was synthesized by DNA polymerase I, RNase H, dNTPs and buffer. Then, the cDNA fragments were purified using a Qia Quick PCR extraction kit (Qiagen, Venlo, The Netherlands), end repaired, poly(A) added, and ligated to Illumina sequence adapters. The ligation products were size-selected by agarose gel electrophoresis, PCR-amplified, and sequenced using an Illumina HiSeq2500 [32].

Transcriptomic data analysis
To obtain high-quality clean reads, we removed the adaptor-containing sequences, poly-N, and low-quality reads. The remaining clean reads were further used in the assembly and gene abundance calculation. Then, clean reads were mapped to the reference genome using the HISAT2 tool [33]. For each transcription region, an FPKM (fragment per kilobase of transcript per million mapped reads) value was calculated to quantify its expression abundance and variations using StringTie software [34, 35].

Differential expression analyses among the four treatments (CK vs. T36/60/72, T36 vs. T60, T36 vs. T72, and T60 vs. T72 with three biological replicates per treatment) were conducted using DESeq2 software [36]. Genes/transcripts with a false discovery rate (FDR) below 0.05 and absolute fold change of ≥2 was considered differentially expressed genes/transcripts.

GO enrichment analysis provided all GO terms that were significantly enriched in the DEGs compared to the genome background and filtered the DEGs that correspond to biological functions. KEGG [37] is the major public pathway-related database. Pathway enrichment analysis identified significantly enriched metabolic pathways or signal transduction pathways in the DEGs compared with the whole genome background.

Quantitative real-time PCR validation
qPCR was used to validate the RNA-seq data for 12 different genes. Specific primers were designed using Premier 5 software (Premier Biosoft, Palo Alto, CA, USA). The RNA samples were used to synthesize cDNA, and a Step OnePlus Real-Time Fluorescent Quantitative PCR system (Trans Start® Green qPCR Super Mix) was used to monitor the amount of DNA. Assays of each gene were repeated three times. Quantification was evaluated using the $2^{-\Delta\Delta C_T}$ method.

![Fig. 10 Hypothetical model of the mechanism of Old Blush tolerance to Botrytis c](image)
Functional verification of RcTGA1
To obtain the TRV-RcTGA1 expression vector, a 250-bp fragment from the ORF of RcTGA1 was cloned into the TRV vector PTRV2 and then electroporated into Agrobacterium strain GV3011. To establish VIGS in rose petals, detached petals were obtained from the outermost whorls of rose flowers at stage 2 of flower opening. Then, a 12-mm disc was punched from the centre of each petal. The petal discs were vacuum infiltrated with Agrobacterium carrying TRV constructs as described by Zhang and Thomma [38]. VIGS was repeated at least three times using at least 16 discs in each experiment. After Botrytis c. inoculation, Student’s t-test was conducted. All primers are listed in Supplemental Table S1.

Extraction and quantification of metabolites
Metabolites were extracted from petals with five replicates per treatment. The compounds extracted were analysed using an LC-ESI-MS/MS system (UPLC, Shim-pack UFLC SHIMADZU CBM30A, http://www.shimadzu.com.cn/; MS/MS (Applied Biosystems 6500 QTRAP)). LIT and triple quadrupole (QQQ) scans were acquired on a triple quadrupole-linear ion trap mass spectrometer (Q TRAP) [39], an AB Sciex QTRAP6500 System, equipped with an ESI-Turbo Ion-Spray interface, operating in positive ion mode and controlled by Analyst 1.6.1 software (AB Sciex). The operation parameters were as follows: ESI source temperature 500 °C; ion spray voltage (IS) 5500 V; curtain gas (CUR) 25 psi; and collision-activated dissociation (CAD). QQQ scans were acquired as MRM experiments with optimized decluttering potential (DP) and collision energy (CE) for each individual MRM transition [40]. The m/z range was set between 50 and 1000.

Metabolites were identified by searching internal databases and public databases (Mass Bank, KN AP Sac K, HMDB, Mo to DB, and METLIN) and comparing the m/z values, RT, and fragmentation patterns with the standards [41].

Metabolomic data analysis
Those with a P value for the T test of < 0.05 and VIP ≥ 1 were considered differential metabolites between those groups. We constructed metabolic pathways based on the information in the KEGG database.

Combined metabolomic and transcriptomic analysis
To reveal the regulatory and influencing mechanism between gene expression and metabolites, we analysed three models based on gene expression and metabolite abundance. The correlation between the top 250 differentially expressed genes and their metabolites was used to draw a heatmap.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12870-021-02973-z.

Additional file 1: Figure S1. Heatmap of the top 250 DEGs and their metabolites.
Additional file 2: Table S1. qPCR primer information.
Additional file 3: Table S2. Number of metabolites.

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Authors’ contributions
Xiangxin Qu, Kaixue Tang and Penghua Gao are the experimental designers and executors of this study; Hao Zhang, Huijun Yan, Qiang Wang, Hongying Jian and Bo Yan participate in the experimental guidance, and Penghua Gao participate in the data processing and paper writing. All authors reviewed the manuscript and approved the final manuscript.

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Availability of data and materials
The Sequence dataset used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Declarations
Ethics approval and consent to participate
Not applicable.
Consent for publication
Not applicable.
Competing interests
The authors declare that they have no competing interests.

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