Incompatible pollen abortion and late-acting self-incompatibility in *Schima superba*

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In angiosperms, self-incompatibility (SI) is a common and widespread mechanism for plant prevention of inbreeding, and late-acting self-incompatibility (LSI) may be ancestral in the group. In this work, we studied *Schima superba*, a species in Theaceae that is a commercially important timer and fire-resistant tree, and revealed its LSI mechanism. Hormones, enzymes, transcriptomes, and proteins were compared between self-pollination (SP) and outcross pollination (OP) in the styles and ovaries from 0 to 120 h after pollination. The self-pollen tubes grew to the bottom of the style and entered the ovary within 48 h but failed to penetrate the ovule. Meanwhile, the hormone and peroxidase levels dramatically changed. Transcriptome and proteome analyses explored the molecular mechanisms of LSI and candidate genes related to LSI in *S. superba*. Overall, 586.71 million reads were obtained, and 79,642 (39.08%) unigenes were annotated. KEGG and GO analysis showed that there were 4531 differentially expressed genes (DEGs) and 82 differentially expressed proteins (DEPs) at 48 h in self- (SP) versus outcross pollination (OP). Among these, 160 DEGs and 33 DEPs were involved in pollen–pistil interactions. “Pollen–pistil interaction,” “signal recognition,” and “component of membrane” were downregulated in SP, whereas “cell wall and membrane biosynthetic process,” and “oxidoreductase activity” were upregulated. The DEGs involved with S-RNases and SCF during SP suggested that the LSI occurred at 48 h in the ovary and that the LSI in *S. superba* was under gametophyte control. Calcium ion increase and release, mitochondrial function loss, and ROS disruption further aggravated PCD progress and cell death. The LSI of *S. superba*, which happened 48 h after pollination, was a key time point. The incompatibility PT ceased growth in the ovary because of S-RNase recognition and PCD in this organ. This study highlights the LSI molecular mechanism in *S. superba* and provides a reference to other species in Theaceae.

### Abbreviations

| Abbreviation | Description                  |
|--------------|------------------------------|
| LSI          | Late-acting self-incompatibility |
| PCD          | Programmed cell death        |
| HR           | Hypersensitive response       |
| SI           | Self-incompatibility          |
| SSI          | Sporophytic SI                |
| GSI          | Gametophytic SI               |
| OSI          | Ovarian SI                    |
| SP           | Self-pollen                   |
| OP           | Outcross-pollen               |
| DEGs         | Differentially expressed genes|
| DEPs         | Differentially expressed proteins|
| PT           | Pollen tube                   |
| IPT          | Incompatible PT               |
| FAA          | Formalin-Aceto-Alcohol        |
| Lpt          | PT length                     |
| Ls           | Total style length            |
| KEGG         | Kyoto Encyclopedia of Genes and Genomes |
| GO           | Gene ontology                 |

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In angiosperms, self-incompatibility (SI) is a common and widespread mechanism to reduce the risk of inbreeding depression, and it is usually under genetic control\(^1\). SI systems can be either sporophytic SI (SSI) or gametophytic (GSI)\(^2\). SSI always occurs in the stigma, and the pollen-specific S-locus cysteine-rich protein (SCR) and stigma-specific S-receptor kinase (SRK) both control pollen tube (PT) growth; the SSI pattern's novel species is in *Brassicaceae*\(^3,4\). In GSI systems, the S-RNase (the female determinant) and S-locus F-box protein (SLF/SFB, the male determinant) interact to mediate the self-pollen tube rejection response, and this pattern is most common in the styles of *Plantaginaceae*, *Solanaceae*, and *Rosaceae*\(^7-9\). In *Papaveraceae*, another kind of GSI is caused by the Ca\(^{2+}\) signaling cascade, leading to apoptosis and the termination of the pollen tube (PT) growth\(^10\). Additionally, ovarian SI (OSI) or late-acting SI (LSI) occurs when the PT ceases growth in the ovary\(^11\). LSI, an ancestrally conserved mechanism of SI, exists widely among *Narcissus papyraceus*, *Clivia gardenia*, *Aloe maculate*, *Theobroma cacao*, and *Thryptomene calycina*\(^12-14\). Since abortion could occur before or after zygote formation, in addition to a much more complicated mechanism in postzygotic SI, the molecular mechanisms of LSI are poorly understood\(^14-16\). LSI can be controlled by the SSI, GSI, or both\(^3\). Studies have shown that some growing molecular signals involved in PT attraction to the synergid cells and ovules affect self-PT growth differentially in LSI species\(^17-20\). For example, SCR/s-locus protein 11 (SP11), which is a synergid-derived cue (LURE) belonging to the supergene family CRPs in *Brassica* pollen, comprises a male signaling ligand determinant\(^21\). Growth of the self-pollination tube is halted in the style, whereas SCF and S-RNase genes expressed in *Camellia sinensis* show that LSI is under gametophytic control\(^22\). Sage predicted that the LSI system's genetic basis is always gametophytic\(^11\). Diallel crosses with sibling progeny arrays suggest that multiple loci with multiple alleles control the LSI response\(^13,24\).

*Schima superba* is a woody plant in the tribe Schimeae in *Theaceae*, which is well-known for the genus *Camellia* in the Theeae tribe. These two tribes have phylogenetic similarities\(^25,26\). Currently, research on SI has made great progress, and several metabolic pathways are regulated by multiple genes in *C. sinensis*\(^22,27\). *Schima superba* is widely distributed in southern China, and its function is mainly for biological fireproofing and valuable timber\(^28,29\). Its fluorescence occurs from the middle of May to the middle of July, and its flowers are small to moderate (length corolla = 31.57 mm) (Fig. 1). In our preliminary study, during outcross pollination (OP), the PTs grew rapidly, reached the bottom of style at 36 h and then penetrate the ovules at 48 h (Fig. 2A,C), however, during self-pollination (SP), the PTs grew slowly from 2 to 36 h after pollination, and then they entered in the ovary, passed the ovule without touching at 48 h (Fig. 2B,D)\(^30\). The fruit abortion rates were over 88% in the SP treatment, and seed yield was only 0.2–0.4%. Therefore, we deduced that *S. superba* is an LSI species. To clarify the LSI molecular mechanism of *S. superba*, we compared hormone levels, such as IAA, ABA, and ZT, between SP and OP in styles and ovaries from 0 to 120 h after pollination. Through physiological results, combined with early anatomical section observation, we further confirmed the accurate time point of PT growth inhibition. Then, at this time point, we associated the transcriptome with proteome analysis to reveal the gene expression pattern, especially to determine whether SSI (S-locus protein), or GSI (SCF/S-RNase), or their combination control PT growth. The results explain the LSI mechanism in *S. superba* and refresh knowledge in *Theaceae*. 

| Acronym | Definition |
|---------|------------|
| TFs     | Transcription factors |
| POD     | Peroxidase |
| CAT     | Catalase |
| SOD     | Superoxide dismutase |
| NBT     | Nitro blue tetrazolium |
| IAA     | Indole-3-acetic acid |
| ABA     | Abscisic acid |
| ZT      | Zeatin |
| HPLC    | High-performance liquid chromatography |
| CC      | Cellular component |
| BP      | Biological process |
| MF      | Molecular function |
| COX     | Cytochrome c oxidase |
| LRR-RLKs| Leucine rich repeat receptor like kinase |
| RALFs   | Rapid alkalization factors |
| SRK     | S-locus receptor kinase |
| SCR     | S-locus cysteine-rich protein |
| SCF     | SKPl-Cullin1-Rbx1-F-box |
| S-RNase | S-locus-encoded ribonuclease |
| MAPK    | Mitogen activated protein kinase |
| cytc    | Cytochrome c |
| IMS     | Intermembrane space |
| Bcl     | B-cell lymphoma |
| DHAR    | Dehydroascorbate reductase |
| GSH     | Ascorbate–glutathione |
| GST     | Glutathione S-transferase |
| GSSG    | Oxidized glutathione |
| APAF    | Apoptotic protease activating factor |
Results

Hormone and enzyme quantification in SP versus OP. To further confirm the PT growth inhibition time point, a quantitative analysis of hormone levels in SP and OP was revealed by HPLC–MS/MS. IAA and ZT biosynthesis were decreased in SP from 24 to 120 h (Fig. 3A,B); however, ABA increased after 60 h in SP (Fig. 3C). The changes in the hormone level showed that 48 h after pollination was a special turning point, in
which IAA, ZT, and ABA were downregulated by 52, 36 and 38%, respectively, in SP ovaries compared to OP ovaries.

Changes in the antioxidant system were important for determining the time point of self-pollinated PT growth inhibition. CAT, POD, and SOD activity increased from 24 to 84 h in both SP and OP (Fig. 4A,B). Although the POD activity was high from 48 to 84 h in SP versus OP, the CAT and SOD activity was low, and the mean value was decreased by 2.3% and 2.7% in SP versus OP (Fig. 4).

**DEGs in self- and outcrossed ovaries.**  Illumina sequencing, de novo assembly, and functional annotation. To identify genes changed during PT growth in the ovary 48 h after OP and SP treatment in *S. superba*, six libraries (three replicates of OP and SP ovaries 48 h after pollination), were sequenced with the Illumina HiSeq 2000 platform. A total of 586.71 million bases of raw reads and 569 million clean reads (Q30 > 92.86%) were generated. In total, 85.36 Gb of nucleotides, 203,767 unigenes, and 79,642 (39.08%) annotated unigenes were generated (Table 1, Supplementary Table S1).

A fold change threshold of ≥ 2.00 and *p* value ≤ 0.05 were used to identify differentially expressed genes (DEGs). There were 4531 DEGs identified in the SP versus OP ovaries, of which 2078 were upregulated and 2453 were downregulated (Supplementary Table S2 and Supplementary Figure S1). The gene ontology (GO) annotation assigned these DEGs to 2453 GO terms and classified them into cellular components (CC: 14%), molecular functions (MF: 24%), and biological processes (BP: 62%) (Supplementary Table S3). Furthermore, 62 putative transcription factor families were identified, and DEGs containing AP2-EREBP were the most abundant (9), followed by bHLH (8), MYB (8), and ABI3VP1 (6) (Supplementary Figure S2). In total, 338 DEGs were categorized into 84 KEGG pathways, 11 of which were significantly enriched (corrected *P* value ≤ 0.05): phenylpropanoid biosynthesis (5.8%), plant–pathogen interaction (5.5%), Pentose and glucuronate interconversions (5.5%), plant...
hormone signal transduction (3.6%), Carotenoid biosynthesis (7.1%), Fatty acid elongation (7.0%), DNA replication (4.8%), Oxidative phosphorylation (2.9%), Limonene and pinene degradation (7.5%), ether lipid metabolism (4.9%), and monoterpenoid biosynthesis (9.1%) (Fig. 5). In addition, “ubiquitin-mediated proteolysis,” “PCD,” and “calcium ion signaling” pathways related to SI were identified as upregulated in SP.

Specifically or preferentially expressed genes involved in SP and OP. To identify differentially expressed genes associated with LSI in S. superba, we compared up- and downregulated DEGs in GO enrichment. Of the 4531 DEGs that were annotated into 2453 GO terms, the upregulated categories were related to “cell wall and membrane biosynthetic process” (13), “cuticle development” (1), and “glycolipid metabolic process” (7) in BP, “host intracellular organelle and membrane” (10) in CC, and “oxidoreductase activity” (16), “ribonuclease T2 activity” (3), and “binding” (21) in MF. However, “pollen–pistil interaction” (9), “recognition of pollen” (9), “response to stress” (8), and “glucose metabolic process” (7) in BP, “signal recognition” (2) and “component of membrane” (32) in CC, “transferase activity” (28), “cellulose synthase activity” (7), and “pheromone activity” (3) in MF were downregulated in SP (S48 vs O48) (Supplementary Table 3 and Supplementary Figure S3). A total of 35 DEGs were annotated in the top 10 KEGG pathways and the top 15 of GO enrichment analysis and were related to ADP binding (5, K13459|K02133), cell wall (7, K13457|K01051), pollination (1, K15397), oxidoreductase activity (13, K09755|K00430|K09843|K00517|K02256), auxin (5, K14488), oligosaccharide (1, K01213), and phosphors signal (3, K14491|K14492|K14500) (Table 2). Furthermore, sesquiterpenoid and triterpenoid biosynthesis (oxidoreductase activity, c119780_g1), glycan biosynthesis (membrane lipid biosynthetic process, c89162_g1), and some immunity-related pathways were upregulated, and some pathways related to amino acid metabolism (Cysteine, c113534_g5|c109638_g2|c123367_g5; Histidine, c109070_g5|c109070_g2; Tyrosine, c118577_g1) and zeatin biosynthesis (c116756_g1) were downregulated in relation to LSI in S. superba.

Of 62 TFs expressed specifically in the self-pollinated ovaries of S. superba, one MYB (c123606_g4) was significantly upregulated; GRAS(c64954_g2), BBR/BPC (c102379_g1), HSF (c117518_g3), and MBF1 (c104077_g2) were specifically upregulated, and FAR1(c107861_g4|c59999_g2|c112693_g3), C2H2 (c148204_g1|c96967_g1), HB (c104944_g1), zf-HD (c95393_g1|c87920_g1), G2-like (c29149_g1), PLATZ (c118044_g7), RWP-RK (c116010_g1), TCP (c104077_g2), and TRAF (c110218_g1) were specifically down regulated in SP. Among these, FAR1, C2H2, GRAS, and TCP were closely related to pollen development.

At 48 h after pollination, growth–related hormones, such as auxin, cytokinin, and ABA31, were significantly downregulated, and after this time, the ABA levels increased and exceeded the normal level (Fig. 3 A–C). At 48 h after pollination, the plant hormone signal transduction genes and proteins underwent corresponding changes;

**Figure 5.** Statistic of pathway enrichment in KEGG of the DEG of SP versus OP.
RT-PCR validation. To determine the transcriptome’s reliability, 23 DEGs associated with “plant–pathogen interaction” (c126261_g1), “pyruvate metabolism” (c114981_g1), and “phenylpropanoid biosynthesis” (c72974_g2), among others, were selected (Supplementary Table S5). These selected DEGs with high repetitiveness verified the accuracy of the sequencing data (Supplementary Figure S4).

DEPs in self- and outcrossed ovaries. To further quantify gene expression in SP, protein expression pattern analysis was performed in SP versus OP ovaries at 48 h after pollination. DEPs were identified by a fold change ≥ 2.0 and ≤ 0.5, and p value ≤ 0.05 in S48 versus O48 (Fig. 6A, Supplementary Figure S5, and Supplementary Table S6). In total, 82 DEPs were identified, and more downregulated proteins (60) were observed in S48 (Fig. 6A). These DEPs were assigned to molecular functions (193 GO terms), cellular components (142 GO terms) and biological processes (786 GO terms), and there were 47, 94, and 229 significant terms, respectively (Fig. 6B, Supplementary Table S7). Most of these DEPs had ion binding, transferase, hydrolase, lyase, and oxidoreductase activity, and they were involved in the development, oxidation–reduction, organic substance metabolic, and cellular metabolic processes.

The categories were significantly upregulated in “oxidative phosphorylation” (COX6B-1|SS204; PDI|SS2022,SS7608; mitochondrial-like ATP synthase subunit d|SS2022, SS2015), and downregulated in “protein glutathionylation” (DHR2|SS9106; GSTL2|SS206), “glutamine biosynthesis” (GLNA|SS7412, SS8421), “aspartate family amino acid catabolic process” (MGL|SS8413; At5g55070|SS6511), and “oxidation–reduction process” (PER12|SS2509, SS2613; At1g75280|SS8315; GLX1|SS3305, SS4306, SS4307; At1g60710|SS8318; SDH1-1|SS8808; PDH2|SS4407; At5g55070|SS6511; At1g53240|SS8319; At3g2090|SS7706; CMTMC|SS8214) (Supplementary Table S7).

In total, 82 DEPs were identified using KEGG analysis and functional enrichment, and 30 pathways were enriched, 10 of which were significantly enriched in metabolism and biosynthesis (Supplementary Table S7).

Association analysis between DEGs and DEPs. Association analysis were performed to clarify the consistency of DEG and DEP expression patterns. DEPs did not have a significant correlation with DEGs at 48 h in the ovaries of SP versus OP (y = 0.047x + 0.1854, R² = 0.0436, P = 0.179, Fig. 7A). Only c113206_g1| SS8413 and c104041_g1 | SS2022 were co-detected from the DEP–DEG association analysis (Fig. 7B). Forty-one proteins and genes annotated in 3 GO functional enrichments were aggregated into four clusters, and only cluster 3 was upregulated in cells (Supplementary Figure S6A). A total of 24 proteins and genes categorized into 7 KEGG pathways were also aggregated into four clusters (Supplementary Figure S6B). These pathways were mainly in oxidation–reduction and basal metabolism.

Discussion
SI is an important mechanism that protects flowering plants, allowing them to overcome inbreeding depression and providing a high level of heterozygosity35. Pollen tubes grow from the stigma to the ovaries, and penetrate the ovules. This forms a zygote and successfully promotes fruit and seed formation. A series of complex signaling controls then occurs. Incompatible pollen can be killed at any stage of this process. Based on our previous40 and present studies, the morphological, anatomical, and gene expression patterns have been comprehensively explored to unravel the complexity of LSI in S. superba. The time point of 48 h, as implicated in this study, was important for PT elongation to the ovary and penetration of the ovule. The transcriptome and proteome data showed that DEGs and DEPs were more enriched in the “oxidation–reduction process,” “ribonuclease T2 activity,” “cell wall or membrane biosynthetic progress,” and “transmembrane transport” in ovaries of SP relative to ovaries of OP. For example, auxin-related DEGs (7), zeatin-related DEG (1), and cytokinin-related DEPs (2) were significantly downregulated 48 h after SP (Supplementary Table S4).

RT-PCR validation. To determine the transcriptome’s reliability, 23 DEGs associated with “plant–pathogen interaction” (c126261_g1), “pyruvate metabolism” (c114981_g1), and “phenylpropanoid biosynthesis” (c72974_g2), among others, were selected (Supplementary Table S5). These selected DEGs with high repetitiveness verified the accuracy of the sequencing data (Supplementary Figure S4).

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of OP, while “recognition of pollen,” “plant hormone signal transduction,” and “Glycolysis” were downregulated (Supplementary Figure S2 and Supplementary Table S8).

Incompatible PT growth and its physiological changes. Researchers have already partly revealed the SI mechanism in some species in Theaceae15,22,26,27,33. Seth27 and Chen33 found that the PTs of SP and OP of C. sinensis entered the ovary at 48 h; however, the IPT recognition system was expressed in the style. Zhang22 showed that PT growth in the SP of C. sinensis was hindered at 24 h in the style. The PT of SP and OP of C. oleif-

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**Figure 6.** Volcano plot of DEPs (A) and GO term classification (B). A, Red plots indicate upregulation, and green plots indicate downregulation.
era entered the ovary at 60 h, but the PT of SP failed to penetrate the ovule. Therefore, 24 h for the style's SI or the ovary's LSI were two features in Theaceae. In our previous research, we found that PT growth stopped and that the IPT recognition system occurred in the ovary 48 h after pollination (Fig. 2)30. From the phylogenetic position of Theaceae and the fossil of Schima, it was shown that the Schima possessed n = 18 symmetric karyotypes, had 5, not 3 locules, had no center axis, and confirmed that the Schimeae (≡ Gordonieae) belonged to an ancient taxon and that this branch should be an early node on the phylogenetic tree in Theaceae24,25,29,34,35. We found the ovary's LSI in S. superba was not effective; during self-pollination, 11.2% still developed fruit30, and C. sinensis had only 1.1%27. This may be for an IPT abortion in the ovary, and this distance may not have been enough to encounter more lethal factors than that from the style to the ovule.

Reactive oxygen species (ROS) are the reactive products of oxygen that have the potential to damage living cells, and they play a key role in diverse development stages, such as self-incompatibility during pollination to induce PCD in incompatible pollen36. In plants, enzymatic and non-enzymatic antioxidants (proline, carotenoids, alpha-tocopherol, glutathione, ascorbic acid, flavonoids, and carotenoids) act as ROS detoxicants. In our results, GST-related proteins, such as the lambda GSTs (GSTL| SS2206) and dehydroascorbate reductases (DHARs| SS9106), which belong to the outlying minor GST classes, lactoylglutathione lyase (GLX1|SS3305, SS4306, SS4307), glutathione (GSH| SS7412| SS8421| c112627_g1| c86003_g1| c90752_g1| c97760_g1), and SOD were all downregulated at 48 h in the ovary of SP. The reduction of detoxification factor levels in SP induces ROS accumulation and damage that can be lethal35–38.

PT guidance and IPT recognition. PT guidance is divided into two processes: pre-ovular and ovular guidance, both of which occur in the ovary37. Receptor-like kinases (RLKs) mediate signaling pathways to control ovular guidance38,39. In our results, 16 G-type lectin S-receptor-like serine/threonine-protein kinases, which included 5 RLK1 (c126066_g2| c100047_g1| c100047_g3| c103929_g1| c126606_g1) and one RKS1 (c106281_g1), in addition to one PT reception gene at rapid alkalinization factor (RALF), c86998_g1, were significantly downregulated at 48 h in the SP ovary.

S-RNase, which is exclusively expressed in pistil, is associated with female determinants in GSI and implicated in genetically identical pollen and the rejection of self-pollen, and it would trigger mitochondrial collapse and IPT's PCD40. In our study, 16 G-type lectin S-receptor-like serine/threonine-protein kinases, which included 5 RLK1 (c126066_g2| c100047_g1| c100047_g3| c103929_g1| c126606_g1) and one RKS1 (c106281_g1), in addition to one PT reception gene at rapid alkalinization factor (RALF), c86998_g1, were significantly downregulated at 48 h in the SP ovary.

The male determinant that interacts with S-RNase degradation is the S-locus region, which contains the novel F-box protein named SLF/SFBs42. The SLF/SFBs interacted with cognate S-RNase and prevented non-self S-RNase catalytic activity. When non-self S-RNase enters the PT, it forms an SCF complex and targets the S-RNase for ubiquitination and degradation; PT then continues growth. Self-S-RNase binds to the recognition domain, not the active domain, resulting in dysfunction of SCF for polyubiquitination of self-S-RNases. Then, self-S-RNases are

Figure 7. Association analysis between DEGs and DEPs. (A) Correlation scatter plot between DEGs and DEPs. The purple dots indicate the none significant difference protein, and the green dots indicate the significant difference protein. (B) Venn diagram showing the overlaps of RNA information obtained by transcriptome and protein information identified by the proteome.
released to trigger subsequent PT growth inhibition events. In our results, 6 SFB putative genes were identified: F-box protein SKIP23-like (c88773_g1|c41648_g1) and F-box LRR-repeat protein (c120445_g3|c103038_g1|c51202_g1|c110213_g1). Phylogenetic analyses of _S. superba_ SFB-like genes together with Rosaceae, Solanaceae and Plantaginaceae SFB and SKP genes, shown in Supplementary Figure S9, revealed that the c51202_g1 gene clusters with the _Prunus_ SFB gene. Interestingly, c51202_g1 had 5 polymorphic SNP loci in the F-box domain, tightly linked to the RHV region (second intron) of c110815_g1 (Fig. 8). However, this gene was expressed in all tissues analyzed here (Supplementary Figure S10), whereas the S-pollen gene(s) were mainly expressed in anthers/pollen only, such as c41648_g1 and c110213_g1; thus, the c51202_g1 detected in _S. superba_, similar to _Prunus_ species, may function as a general inhibitor (GI), as in _Prunus_, and c41648_g1 and c110213_g1 may function as SLP. Further experiments should be conducted to clarify these proteins’ interactions and verify this presumption. qRT-PCR showed that c110815_g1 was mainly expressed in the ovary and increased in SP ovaries from 24 to 72 h, and c51202_g1, c41648_g1, and c110213_g1 were significantly upregulated at 48 h in the ovary of SP (Fig. 9). We presumed that these candidate genes were active in the self-S-RNase’s cytotoxicity at 48 h in the ovary, and LSI in _S. superba_ could be gametophyte controlled.

**Programmed cell death in IPT.** Apoptosis or PCD is a highly conserved mechanism that removes unwanted cells in eukaryotes. At 48 h in the ovary’s LSI in _S. superba_, the IPTs ceased growth and died. As mentioned above, the S-RNase induces apoptosis or PCD by direct degradation of ribosome RNA or indirect phosphorylation of the protein, destabilization of cytoskeletons, and release of cytochrome c (cyt c). Indirect progress related to Ca²⁺ signaling is transduced by protein phosphorylation, and this signal transduction mechanism relates to mitogen-activated protein kinase (MAPK) cascades. The Ca²⁺ concentration in the cytosol can also trigger downstream sensors to form stimulus-specific information. In _S. superba_, one MAPK gene c120765_g5 was significantly upregulated, and the “Ca²⁺ channel activity,” “calcium ion transmembrane transporter activity,” and “calcium ion transport” pathway-related genes and proteins were also upregulated.

The important prophase characteristic of PCD inhibits mitochondrial outer membrane permeabilization (MOMP), and the mitochondria system loses its function. Then, some soluble proteins, such as cyt c, diffuse from the intermembrane space (IMS) to cytosol. GO analysis of DEGs showed that 30 GO class 14 DEGs related to mitochondria, such as inner mitochondrial membrane protein complex, mitochondrial membrane organization, and mitochondrial proton-transporting ATP synthase complex, were downregulated, and none of these were upregulated at 48 h in the ovary of SP (Supplementary Table S9). This demonstrated that, at 48 h in the ovary of SP, the mitochondrial membrane system was dysfunctional, and PCD had already started. In addition, during apoptosis, Bcl-2 family proteins regulate soluble proteins (e.g., cyt c), which are released from mitochondria. The Bcl-2-associated athanogene-like protein gene c122719_g2, which contained the BAG domain, was 2.5-fold more highly expressed at 48 h in the ovary of SP (Supplementary Table S10). Kang reported that _AtBAG6_, which contains the BAG domain, was overexpressed in Arabidopsis, and promoted mitochondrial fusion, causing cyt c release into the cytosol and inducing PCD.

Cyt c leakage from the mitochondria into the cytosol is an initial marker for PCD. In our study, the concentrations of cytochrome oxidase COX (SS204) and ATP synthase (SS2015) were 9.7- and 5.8-fold higher, respectively, at 48 h in the ovary of SP. COX is the enzyme that catalyzes the oxidation of reduced cyt c by molecular oxygen. A higher COX concentration at 48 h in the ovary of SP indicates that the plants have already started the self-protection mechanism due to higher levels of cyt c. If the COX concentration exceeds the critical
value, the activity is limited by either the cyt c or ascorbate concentration, and PCD occurs. These results are consistent with the results of the disordered oxidation–reduction system shown previously.

**Conclusions**

In this study, the analysis of PT growth characteristics, internal physiology changes, transcriptome, and proteome of SP and OP ovaries revealed the LSI mechanism in *S. superba* and identified several candidate genes. The time point 48 h after pollination in the ovary's LSI in *S. superba* was different from 24 h after pollination in style's SI in *C. sinensis*. High expression levels of 2 S-RNase genes and 3 SLF-related genes suggest that LSI in *S. superba* is under gametophyte control. Self S-RNase induced PCD of IPT could occur by indirect transduction of Ca++ signaling and release of cyt c. Mitochondrial fusion and function loss caused the PCD; in addition, oxidation–reduction system disorder promoted cell death. These results revealed the LSI molecular mechanism in *S. superba* and provided a reference for other plants in the Theaceae family.

**Methods**

**Plant materials and pollination treatment.** This study was conducted by the Forestry Genetic and Breeding Lab at the Research Institute of Subtropical Forestry, CAF, State Forestry Administration, China (RISF-CAF). The State Forestry Administration is responsible for national parks and other protected areas. No specific permission was required for these locations/activities, as they were based on a non-destructive collection of plant material. The species is not endangered or protected, and the locations are not privately owned or protected by law.

Two cultivars, 'JO59' and 'YX1', grown in Lanxi nursery, Zhejiang province, China, were used in this study. These cultivars were identified, bred, and preserved by our institution. The scions were collected from selected trees of the natural forest of *S. superba* from Jianou and Youxi in Fujian; they were then grafted on local stock and maintained in Lanxi until 2013. The collection of plant material complied with institutional, national, and international guidelines. Field studies were conducted in accordance with local legislation.

In May 2019, at 9:00–11:00 am on sunny days, we performed pollination [self-pollination (SP) was 'JO59' × 'JO59', outcross-pollination (OP) was 'JO59' × 'YX1']. The styles and ovaries were collected at different intervals (2, 4, 8, 12, 16, 24, 36, 48, 60, 72, 84, 96, and 120 h after pollination).

Then, the styles and ovaries of each sample with three replications were frozen in liquid nitrogen and stored at −80 °C for enzyme activity, hormone, amino acid, protein, and RNA-seq analysis.

**Enzyme assays.** Three defense-related enzymes, peroxidase (POD), catalase (CAT), and superoxide dismutase (SOD), were analyzed using SP and OP ovaries from 24 to 120 h. POD activity was determined using the guaiacol–catalase activity method. SOD activity was measured using inhibition in the photoreduction of

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**Figure 9.** qRT-PCR expression analysis of 3 SFB genes (A–C) and 2 S-RNase (D,E) using GAPDH as the internal control from 24 to 72 h in the style and ovary of SP and OP. **p values (0.05).**
Hormone extraction and determination. Using HPLC–MS/MS, the levels of auxin (IAA), abscisic acid (ABA), zeatin (ZT) were determined by Zoonbio Biotechnology Co., Ltd. (Nanjing, China). The SP and OP ovaries (0.1 g fresh weight), with three biological replicates from 24 to 120 h, were selected to extract the hormone, and HPLC–MS/MS analysis was performed (Agilent 1290). HPLC analysis was carried out using a poroshell 120 SB-C18 column (2.1 mm × 150 mm; 2.7 μm). The parameters were set as follows: spray voltage, + 4500 V, atmospheric temperature, 400 °C, the pressure of the air curtain, nebulizer, and aux gas were 15, 65, and 70 psi, respectively. SAS v8.0 (SAS Institute, Cary, NC, USA) was used to analyze the data.

Transcriptome analysis. The total RNA from the S48 and O48 ovaries was extracted using an RNAprep pure Plant Kit (Dingguo, Beijing, China). RNA quality and quantity were verified using 1% agarose gels and Agilent 2100 (Agilent Technologies, CA, USA). The mRNA was enriched and purified by magnetic beads with oligo T (dT), cleaved and synthesized first-strand cDNA using random hexamers. The double-strand cDNA libraries were then purified using the AMPure XP system (Beckman Coulter, Beverly, USA). Transcriptome sequencing was performed using the Illumina HiSeq™ 2000 sequencing platform in Novogene (Tianjin, China). The adapter and low-quality sequences (Q < 20 or less than 35 bp) were removed in raw reads. Then the clean reads were assembled using Trinity assembler. The functions of the unigenes were annotated using Blastx searches with e-value < 1e-5 against the protein databases. Blast2GO software was used for gene Ontology (GO) term analysis. The plant Transcription Factor Database (PlnTFDB) (version 3.0) was used to determine the transcription factors (TFs). FPKM was used to estimate the expression of each gene, and the DESeq R package 1.10.1 was used to estimate the differential expression between the two treatments. DEGs were determined with an adjusted p value < 0.05 determined by DESeq. The R-seq data were uploaded to the sequence read archive (accession no. SUB6596208).

Protein extraction and expression analysis. With some modification, protein extraction was performed following Isaacson. Then, 10 μg samples were run on 12% SDS-PAGE gel and visualized by CBB stain. Gel images with 300 dots per inch were scanned using an image scanner (GE Healthcare, USA). A total of 450 μl solution with 1500 μg protein sample was used to finish isoelectric focusing with the following parameters: 50uA per strip, hydration at 50 V for 8 h, 100 V for 1 h, 200 V for 1 h, 500 V for 1 h, 1000 V for 1 h, 10,000 V for 1 h, 20,000 V for 1 h, 100,000 V for 1 h, 5000 V for 2 h, temperature, 20 °C. The Ettan-DALT-Six system was run for 45 min at 100 V and then at 300 V for 6–8 h. Using PDquest 8.0 software, all gel images were processed in three steps: spot detection, volume quantification, and matching. The differential protein spots were selected using two thresholds (p ≤ 0.05, fold change ≥ 2 or ≤ 0.5). An ABI 5800 MALDI-TOF/TOF Plus mass spectrometer (Applied Biosystems, Foster City, USA) was used for peptide MS and MS/MS detection, and CalMix5 was used to calibrate the instrument (ABI5800 Calibration Mixture). GPS Explorer V3.6 software (Applied Biosystems, USA) with default parameters was used for data integration, and the proteins were identified using the Mascot V2.3 search engine (Matrix Science Ltd., London, U.K.).

Transcriptome and proteome association analysis. The corresponding transcripts were identified using the gene ID of the proteome, and the corresponding relationship between the protein and the transcript was determined. A series of association analyses were then performed. Correlation analysis was conducted on the fold change (taken as log2) of genes (proteins) identified by transcriptome and proteome analyses in the two omics studies. The drawing process was implemented using Python. The different multiples of the corresponding transcriptome genes were determined in the two omics studies. The difference multiples were taken as log2 and then plotted using the ComplexHeatmap of the R package. GO annotation of the transcriptome and proteome omics stydies were then performed using two thresholds (p ≤ 0.05, fold change ≥ 2 or ≤ 0.5). An ABI 5800 MALDI-TOF/TOF Plus mass spectrometer (Applied Biosystems, Foster City, USA) was used for peptide MS and MS/MS detection, and CalMix5 was used to calibrate the instrument (ABI5800 Calibration Mixture). GPS Explorer V3.6 software (Applied Biosystems, USA) with default parameters was used for data integration, and the proteins were identified using the Mascot V2.3 search engine (Matrix Science Ltd., London, U.K.).

RT-PCR. Using specific primers designed by Beacon Designer 7.90 (Premier Biosoft International, Palo Alto, CA, Supplementary Table S5), the 22 expressed transcripts were validated. The candidate gene expression patterns were tested in two ways: (1) tests in organs to determine whether they were expressed in whole plants (leaves) or pollen (fresh and obtained from stamens before pollination), style, or ovary; (2) candidate genes were also detected in the style and ovary (contained the pollen tubes) of SP and OP at 0, 24, 48, and 72 h to clarify the trend in the gene expression pattern and which time point was important. To synthesize cDNA, 100 ng of total RNA in the ovaries was used (PrimeScript™ RT reagent Kit, Takara, Dalian, China). Then, RT-PCR was performed on the ABI QuantiStudio 7 Flex Real-Time PCR System (Applied Biosystems) using a SYBR® Fast qPCR Mix (Takara, Dalian, China). The program was as follows: 95 °C for 30 s, 40 cycles at 95 °C for 15 s, and 60 °C for 30 s. The reference gene was set as GAPDH. Three biological and technical replicates were performed and the gene expression relative quantitation was calculated using the 2−ΔΔCt method.

Ethics approval and consent to participate. All field studies were performed in accordance with the local legislation in China and complied with the convention on trade in endangered species.
Data availability
The RNA-seq and proteome datasets in this article are available in supplementary files. All raw data were submitted to the NCBI Sequence Read Archive: BioProject ID PRJNA625576.

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