

# Loss of cold tolerance is conferred by absence of the *WRKY34* promoter fragment during tomato evolution

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Natural evolution has resulted in reduced cold tolerance in cultivated tomato (*Solanum lycopersicum*). Herein, we perform a combined analysis of ATAC-Seq and RNA-Seq in cold-sensitive cultivated tomato and cold-tolerant wild tomato (*S. habrochaites*). We identify that *WRKY34* has the most significant association with differential chromatin accessibility and expression patterns under cold stress. We find that a 60 bp InDel in the *WRKY34* promoter causes differences in its transcription and cold tolerance among 376 tomato accessions. This 60 bp fragment contains a GATA *cis*-regulatory element that binds to SWIBs and GATA29, which synergistically suppress *WRKY34* expression under cold stress. Moreover, *WRKY34* interferes with the CBF cold response pathway through regulating transcription and protein levels. Our findings emphasize the importance of polymorphisms in *cis*-regulatory regions and their effects on chromatin structure and gene expression during crop evolution.

The divergence of gene function, primarily driven by mutation, gene duplication, and gene loss, is fundamental to evolutionary processes<sup>1,2</sup>. Such divergence in gene function may be caused by mutations in coding regions that alter protein function. For instance, a 45 bp deletion in the *ZmRRI* coding region prevents its phosphorylation by ZmMPK8, inhibiting its degradation via the 26S proteasome pathway and thereby enhancing maize cold tolerance<sup>3</sup>. Alternatively, divergence in gene function may also result from mutations in *cis*-regulatory regions, which interact intricately to shape expression patterns across different tissues during development<sup>4</sup>. For example, the CRISPR/Cas9 *cis*-regulatory allelic series of tomato *SIWOX9* reveals that different pleiotropic functions can be mapped to specific *cis*-regulatory regions<sup>5</sup>. Evolutionary innovations in transcriptional regulation often result from changes in *cis*-regulatory regions, where abundant sequences directly binding to transcription factors offer significant mutational potential to alter gene expression and phenotypes<sup>6</sup>. For instance, a crucial variation in the W-box motif within the *SIWRKY33*

promoter suppresses its self-transcriptional activity in response to cold stress, thus contributing to the cold sensitivity observed in cultivated tomatoes compared to cold-tolerant wild tomatoes<sup>7</sup>. Nevertheless, our comprehension of the alterations and roles of *cis*-regulatory regions during crop evolution, as well as their regulatory mechanisms, remains limited.

In recent years, our understanding of transcriptional regulation has broadened to include the level of chromatin structure<sup>8</sup>. Chromatin remodeling, which involves the dynamic modification of chromatin structure, plays a crucial role in controlling the accessibility of transcriptional machinery to DNA<sup>9</sup>. In the medical field, chromatin remodeling has been extensively studied for its impact on cell differentiation, organ development, and its involvement in diseases such as cancer<sup>10,11</sup>. For instance, the chromatin remodeler CHD6 promotes colorectal cancer development by regulating TMEM65-mediated mitochondrial dynamics<sup>12</sup>. In botany, chromatin remodeling is increasingly recognized as pivotal in how plants respond to

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environmental cues. Several studies have indicated that environmental stress can alter chromatin structure, thereby influencing transcriptional regulation<sup>13,14</sup>. For example, heat stress triggers genome-wide chromatin accessibility changes in tomato, with HSFA1 binding promoting the formation of promoter-enhancer contacts to drive the expression of heat stress-responsive genes<sup>15</sup>. Similarly, cold stress enhances chromatin accessibility and leads to bivalent histone modifications of active genes in potato<sup>16</sup>. Moreover, a lamin-like protein OsNMC1 in rice modifies chromatin accessibility by interacting with a chromatin remodeler OsSWI3C, thereby regulating numerous genes involved in root growth and drought response<sup>17</sup>. Hence, changes in chromatin structure may serve as the initial step in initiating transcriptional stress responses. Nevertheless, the mechanisms and extent to which environmental stress induces chromatin dynamics remain largely unknown. Additionally, the causal relationship between chromatin dynamics and transcriptional responses under environmental stress requires further elucidation.

The SWIB/MDM2 domain superfamily of proteins comprises a group of proteins characterized by the presence of the SWIB (SWI/SNF complex BCL7/BCL7A interacting domain) and/or MDM2 (Mouse Double Minute 2 homolog) domains<sup>18,19</sup>. These proteins are evolutionarily conserved and exist in various eukaryotes<sup>20,21</sup>. Numerous studies have highlighted the critical role of SWIB/MDM2 domain proteins in diverse cellular processes, particularly in chromatin remodeling and gene expression regulation<sup>20</sup>. For example, the SWP73 protein, a member of the SWIB/MDM2 domain superfamily, is essential for supporting yeast growth at elevated temperatures. Additionally, it plays a pivotal role in repressing seedling growth by modulating chromatin accessibility of genes regulating hypocotyl cell size in *Arabidopsis*<sup>22,23</sup>. Furthermore, a study underscores the importance of SWIB-4, a SWIB domain protein in spinach chloroplasts, which not only structures the nucleoid core but also binds DNA via its histone H1 motif, thus playing a crucial role in the compaction and regulation of chloroplast DNA<sup>24</sup>. Nevertheless, the functions of these proteins in plants, especially their ability for direct DNA binding in the nucleus, remain largely unexplored.

Cold stress poses significant threats to crops, leading to reduced growth, impaired development, and lower yields. Plants have evolved various pathways to withstand cold stress, including the CBF-COR pathway and hormonal pathways<sup>25</sup>. In the CBF-COR pathway, ICE (Inducer of CBF Expression) proteins act as upstream regulators, activating C-repeat binding factors (CBFs), which in turn induce the expression of cold-responsive (COR) genes to enhance cold tolerance<sup>26</sup>. Different subspecies within a species often exhibit distinct cold tolerances due to evolutionary adaptations to their specific environments. For example, a study employing a combination of genetic mapping and gene expression analysis revealed that temperate *japonica* rice varieties have evolved lower expression of *HAN1* gene due to an increase in MYB *cis*-elements within its promoter during domestication. This adaptation enhances chilling tolerance mediated by jasmonic acid (JA), aiding in the adaptation to a temperate climate<sup>27</sup>. Similarly, using a metabolite genome-wide association study (mGWAS), variations in the *ZmICE1* promoter were identified to affect its interaction with ZmMYB39, thereby influencing cold tolerance in maize through the regulation of metabolic reprogramming and *COR* gene expression<sup>28</sup>. Therefore, it is of great importance to utilize multi-omics analysis to identify key genetic loci for cold tolerance in crops. Different wild tomatoes have different levels of cold tolerance, and *Solanum habrochaites* is considered to be one of the most cold tolerant wild tomatoes<sup>29</sup>.

In this study, through the combined analysis of transposase-accessible chromatin sequencing (ATAC-Seq) and transcriptome sequencing (RNA-Seq), we observe that expression of *WRKY34* remains largely unchanged following cold treatment in cold-sensitive cultivated tomato *S. lycopersicum*. Conversely, in cold-tolerant wild tomato

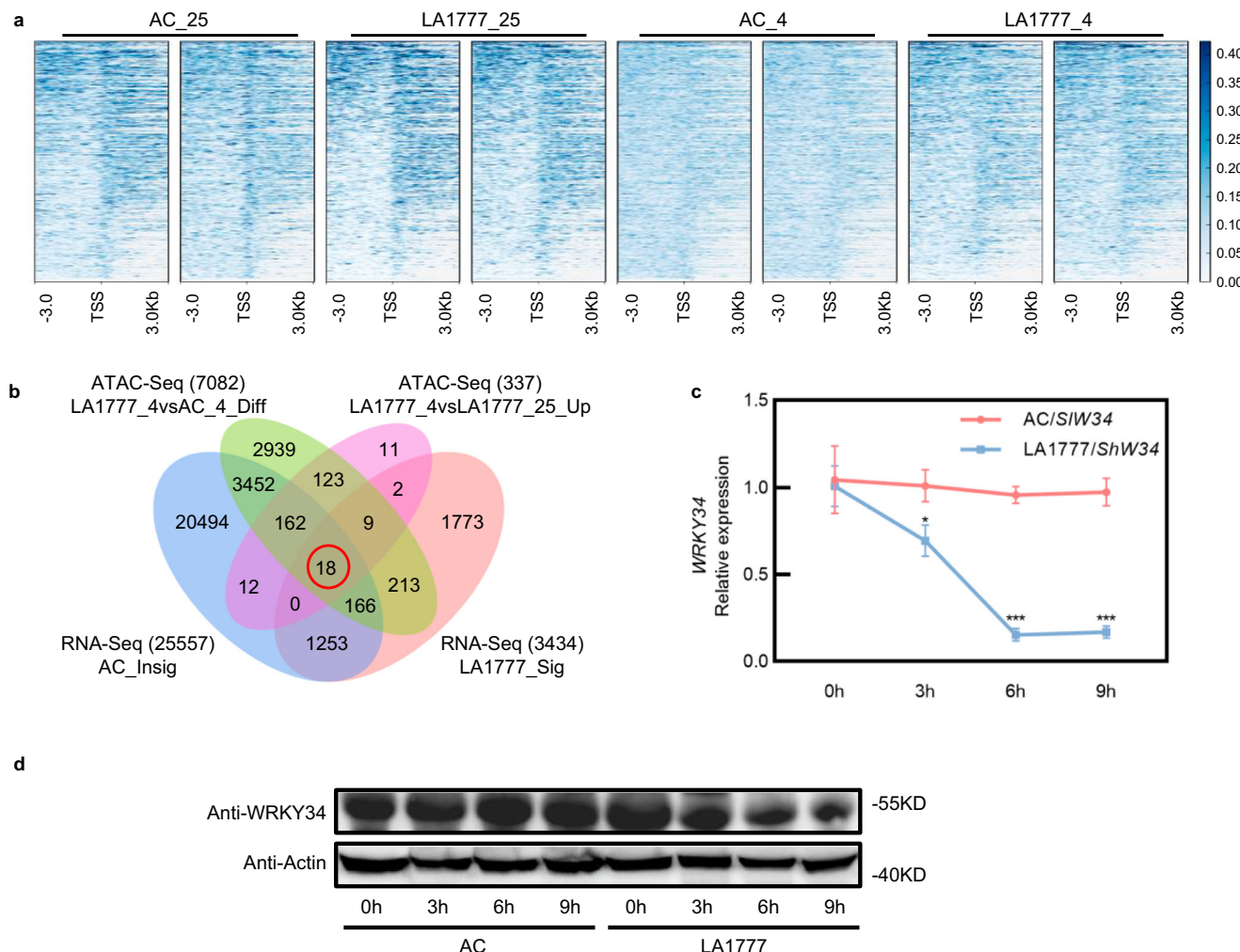
*S. habrochaites*, exposure to cold leads to transcriptional suppression of *WRKY34*, accompanied by chromatin opening. Importantly, we identify a 60 bp InDel in the *WRKY34* promoter that affects its binding to SWIBs and transcription factor GATA29, thereby influencing its chromatin accessibility and expression level under cold stress. Additionally, we demonstrate that SWIBs and GATA29 interact with each other to cooperatively suppress the expression of *WRKY34*. Furthermore, *WRKY34* interferes with the CBF-COR cold response pathway through interaction with CBF1 or direct transcriptional inhibition, thereby negatively regulating cold tolerance. Our study elucidates that polymorphisms in *cis*-regulatory regions leading to differences in chromatin structure and gene expression during crop evolution, providing insights into the natural regulatory mechanism of cold tolerance in tomato.

## Results

### A potential role of *WRKY34* in tomato response to cold stress

Previous research has shown substantial difference in cold tolerance between cultivated tomato and various wild tomato species<sup>7</sup>. To delve deeper into the distinctions in the potential regulatory mechanisms of cold tolerance between wild and cultivated tomatoes, we exposed the cold-sensitive cultivated tomato (*Solanum lycopersicum*) Ailsa Craig (AC) and the cold-tolerant wild tomato (*S. habrochaites*) LA1777 to cold stress for 6 h, followed by ATAC-Seq and RNA-Seq analyses, respectively. ATAC-Seq data analysis revealed that genome-wide chromatin accessibility decreased rather than increased in both wild and cultivated tomatoes after cold treatment (Fig. 1a and Supplementary Fig. 1a). High Spearman correlation coefficients between biological replicates indicate the reliability of ATAC-Seq results (Supplementary Fig. 1b). Genes near peaks exhibiting decreased chromatin accessibility (closing chromatin regions) under cold stress in both cultivated tomato AC and wild tomato LA1777 were predominantly enriched in pathways related to growth and development, such as photosynthesis, starch and sucrose metabolism and amino acid metabolism (Supplementary Fig. 1c). However, under cold stress conditions, the enrichment ratio of chromatin accessibility peaks in wild tomato LA1777 was significantly higher than that in cultivated tomato AC (Supplementary Fig. 1d). Furthermore, genes associated with differential chromatin accessibility peaks (DCAPs) between the two accessions were primarily enriched in gene ontology (GO) terms such as DNA binding, organic cyclic compound binding and heterocyclic compound binding (Supplementary Fig. 1e). Analysis of ATAC-Seq data revealed 7082 genes associated with DCAPs between the two accessions (AC and LA1777) under cold stress conditions (Fig. 1b and Supplementary Data 1), along with 337 genes showing increased chromatin accessibility peaks (opening chromatin regions) in wild tomato LA1777 post cold treatment (Fig. 1b and Supplementary Data 2). Analysis of RNA-Seq data revealed 3434 differentially expressed genes (DEGs) in wild tomato LA1777 under cold stress, whereas 25557 genes in cultivated tomato AC exhibited no significant difference under cold stress (Fig. 1b and Supplementary Data 3, 4).

To further identify candidate genes responsible for cold tolerance in tomato, we integrated the ATAC-Seq and RNA-Seq data as described above and generated a Venn diagram (Fig. 1b). The intersection of the Venn diagram revealed that the expression levels of 18 genes remained unchanged in the cold-sensitive cultivated tomato AC after cold treatment, but showed differential expression in the cold-tolerant wild tomato LA1777 after cold treatment. Interestingly, chromatin accessibility of these 18 genes were significantly different between the two accessions (AC and LA1777) under cold stress, with chromatin opening observed in the cold-tolerant wild tomato LA1777 after cold stress (Fig. 1b and Supplementary Data 5). The expression of these 18 genes was validated through RT-qPCR analysis, confirming the reliability of the RNA-Seq data (Supplementary Fig. 2a). Additionally, RT-qPCR validation was performed on three genes from the RNA-Seq data that



**Fig. 1 | Combined analysis of ATAC-Seq and RNA-Seq, expression patterns and protein accumulation differences of WRKY34 in cultivated and wild tomatoes after cold stress.** **a** Heatmaps of ATAC-Seq read density in all samples (two replicates per sample) within a 3 kb window centered on transcriptional start sites (TSS). Detected accessible regions are showed. Each row represents one peak. The color represents the intensity of chromatin accessibility. **b** Venn diagram illustrating cold response candidate genes identified by combined ATAC-Seq and RNA-Seq analysis. Categories include: ATAC-Seq (7082) LA1777\_4vsAC\_4\_Diff, indicating 7082 differential chromatin accessibility peaks (DCAPs)-associated genes between the two accessions (cultivated tomato AC and wild tomato LA1777) under cold stress conditions identified by ATAC-Seq data; ATAC-Seq (337) LA1777\_4vsLA1777\_25\_Up, indicating 337 increased chromatin accessibility peaks (opening chromatin regions) related genes in wild tomato LA1777 after cold stress identified by ATAC-Seq; RNA-Seq (25557) AC\_Insig, indicating 25557 genes with no significant change in

expression in cultivated tomato AC after cold stress identified by RNA-Seq data; RNA-Seq (3434) LA1777\_Sig, indicating 3434 differentially expressed genes (DEGs) in wild tomato LA1777 under cold stress identified by RNA-Seq data. **c** Expression levels of *SIWRKY34* in *S. lycopersicum* (Ailsa Craig, AC/SIW34) and *ShWRKY34* in *S. habrochaites* (LA1777/ShW34) plants under cold stress. Total RNA was isolated from leaf samples at the indicated times. **d** WRKY34 protein accumulation in AC and LA1777 under cold stress. The Actin protein was used as a loading control. The experiments were repeated three times with similar results (c, d). Values are expressed as means  $\pm$  SD,  $n = 3$  ( $^*P < 0.05$  and  $^{***}P < 0.001$ ; two-tailed Student's *t*-test). AC\_25, cultivated tomato AC under normal temperature (25 °C); AC\_4, cultivated tomato AC under cold stress (4 °C); LA1777\_25, wild tomato LA1777 under normal temperature (25 °C); LA1777\_4, wild tomato LA1777 under cold stress (4 °C). Source data are provided as a Source Data file.

exhibited significant expression changes after cold treatment in AC but no difference in LA1777, as well as three genes that showed no difference in expression after cold treatment in both AC and LA1777 (Supplementary Fig. 2b, c). The expression patterns of these genes validated by RT-qPCR were consistent with those observed in the RNA-Seq data. Subsequently, we analyzed the 18 genes that displayed differential expression patterns and chromatin accessibilities between wild and cultivated tomatoes after cold stress. These genes, including *MAPKKK86*, transcription factor *WRKY34*, phosphatidylinositol transfer protein *SFH5*, and chaperone protein *dnaJ*, among others, play diverse roles in plant responses to cold stress, encompassing signal transduction, gene transcription, DNA repair, metabolism regulation, protein processing, and cell structure maintenance (Supplementary Data 5). The differential accessibility regions of these 18 genes were

primarily located in distal intergenic regions, with four of these genes exhibiting significantly induced expression levels in LA1777 after cold treatment, while the expression of fourteen genes was significantly down-regulated following cold treatment in LA1777 (Supplementary Fig. 3 and Supplementary Data 5).

Chromatin accessibility of genes, particularly in the promoter regions, can directly impact gene transcriptional activity<sup>30</sup>. Previous studies have shown that many crucial regulatory elements, such as binding sites for most transcription factors, are typically situated within 3 kb of gene promoters<sup>31,32</sup>. Consequently, we directed our attention to genes whose differential chromatin accessibility regions were located within the 3 kb promoter region. We identified only one gene, *WRKY34*, that not only exhibited a differential accessibility region within 3 kb of the promoter between two accessions (AC and

LA1777) under cold stress, but also corresponded to the chromatin opening region in LA1777 after cold treatment within 3 kb of the promoter (Supplementary Fig. 3 and Supplementary Data 5). The genome browser view of ATAC-Seq results for *WRKY34* illustrated that the chromatin surrounding *WRKY34* was predominantly closed in both AC and LA1777 plants under normal conditions. However, while chromatin remained closed in AC plants after cold stress, the chromatin within 2–3 kb upstream of the *WRKY34* promoter noticeably opened in LA1777 plants after cold stress (Supplementary Fig. 3). Furthermore, the transcripts of *WRKY34* exhibited minimal change in cold-sensitive cultivated tomato AC, but were significantly down-regulated in cold-tolerant wild tomato LA1777 under cold stress (Fig. 1c and Supplementary Data 5). Additionally, consistent with the transcript levels of *WRKY34*, we observed that *WRKY34* protein accumulation in cultivated tomato AC leaves remained largely unaffected by cold stress, whereas the abundance of *WRKY34* protein in wild tomato LA1777 leaves gradually declined with increasing duration of cold treatment (Fig. 1d). These results suggest that transcription factor *WRKY34* may play a potential role in regulating tomato cold tolerance.

### WRKY34 negatively regulates cold tolerance of tomato

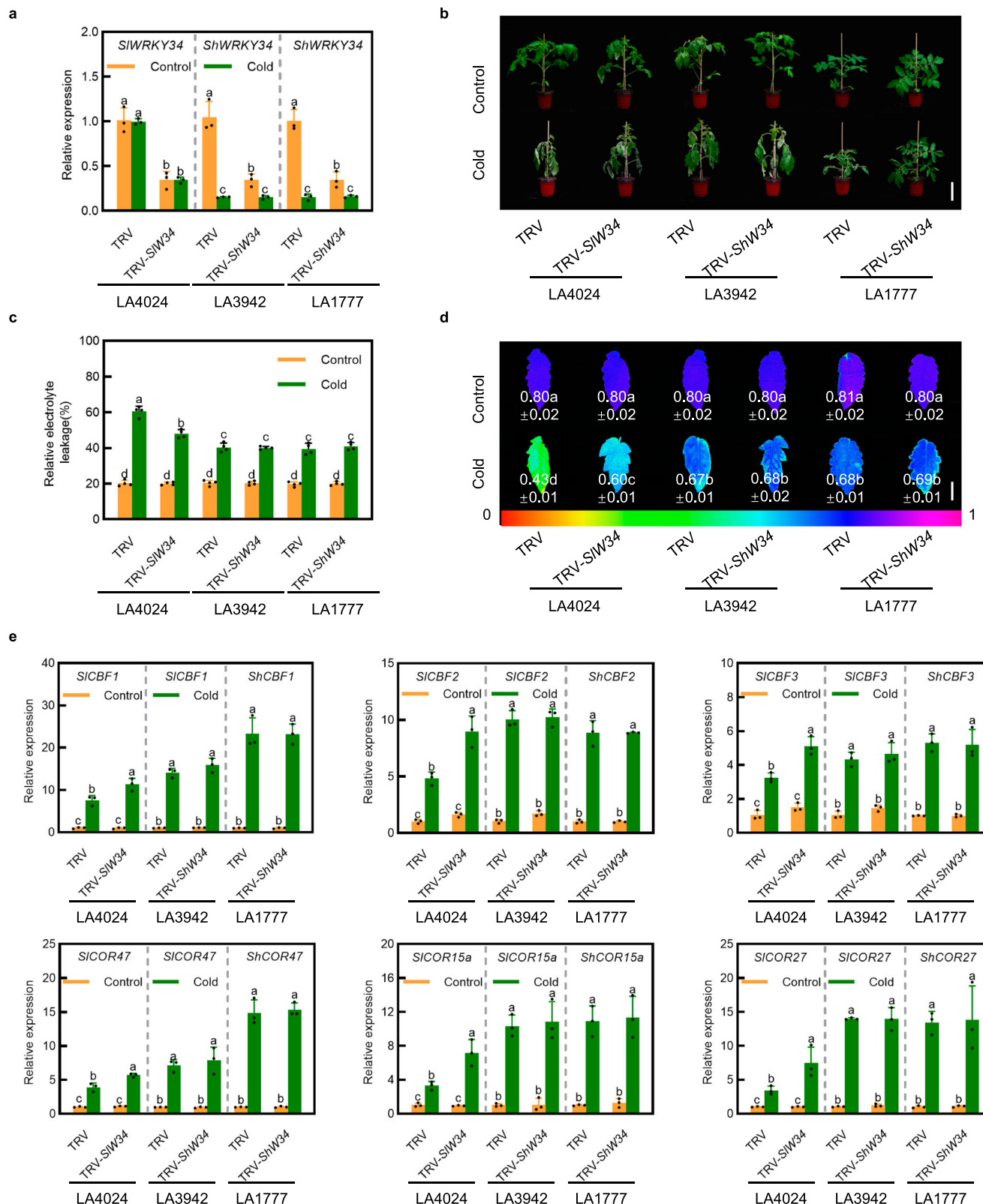
Both cultivated tomato *SIWRKY34* and wild tomato *ShWRKY34* are situated at the end of chromosome 5 in their respective genomes. To investigate the function of *WRKY34* alleles in response to cold stress in wild and cultivated tomatoes, we introduced *S. habrochaites* introgression line LA3942, which carries *ShWRKY34* instead of *SIWRKY34*, along with its recurrent parent *S. lycopersicum* LA4024 and donor parent *S. habrochaites* LA1777 (Supplementary Fig. 4). We silenced *SIWRKY34* (TRV-*SIW34*), *ShWRKY34* (TRV-*ShW34*) in LA4024, LA3942 and LA1777 plants using virus-induced gene silencing (VIGS) technique, respectively. Silencing efficiency exceeded 65%, resulting in significantly reduced *WRKY34* expression levels in silenced seedlings compared to non-silenced seedlings (TRV) (Fig. 2a). After cold stress, *ShWRKY34* expression in TRV control and *ShWRKY34*-silenced seedlings (TRV-*ShW34*) of LA3942 and LA1777 was decreased significantly, while *SIWRKY34* expression in TRV and *SIWRKY34*-silenced seedlings (TRV-*SIW34*) of LA4024 showed no significant difference compared to their respective control plants (Fig. 2a). Under normal conditions, no discernible phenotype change was observed in seedlings after *WRKY34* silencing (Fig. 2b). Notably, TRV seedlings of LA3942 and LA1777, containing the *ShWRKY34* gene, exhibited greater tolerance to cold stress than TRV seedlings of LA4024, which contains the *SIWRKY34* gene. This was evidenced by lower relative electrolyte leakage (REL), higher maximum photochemical efficiency of photosystem II (*Fv/Fm*) and higher survival rate under cold stress (Fig. 2b–d and Supplementary Fig. 5a). Silencing of *ShWRKY34* in LA3942 and LA1777 still maintained strong cold tolerance in tomato seedlings. Importantly, silencing of *SIWRKY34* in LA4024 significantly increased cold tolerance, resulting in lower REL, higher *Fv/Fm* and higher survival rate compared with its TRV under cold stress (Fig. 2b–d and Supplementary Fig. 5a).

Furthermore, we assessed the expression levels of cold responsive genes *CBFs* and *CORs* in the aforementioned tomato plant lines. As shown in Fig. 2e, under cold stress, cold-induced up-regulation of *CBFs* and *CORs* in TRV seedlings of LA3942 and LA1777 was significantly higher than that in TRV seedlings of LA4024. Silencing *ShWRKY34* in LA3942 and LA1777 maintained similar expression levels of these cold responsive genes after cold treatment compared to their respective TRV control plants. Importantly, silencing of *SIWRKY34* in LA4024 significantly increased the expression of these cold responsive genes after cold treatment compared to its TRV control plants (Fig. 2e). These results indicate that *SIWRKY34* does not respond to cold stress and negatively regulates cold tolerance in cultivated tomato, while the expression levels of *ShWRKY34* are decreased in both wild tomato and the *ShWRKY34* introgression line with strong cold tolerance.

### Cold-suppressed *WRKY34* expression is associated with a 60 bp InDel in its promoter region

To further elucidate the role of *WRKY34* in tomato cold tolerance, we compared protein sequences of *WRKY34* from six different tomato species (*S. lycopersicum*; *S. lycopersicum* var. *cerasiforme*; *S. pimpinellifolium*; *S. chilense*; *S. pennellii*; *S. habrochaites*) through amino acid alignments. The comparison revealed that *WRKY34* proteins across different tomato species were similar and conserved. For example, there are only ten amino acid differences among the six different tomato species, with merely four amino acid distinctions between cultivated tomato *S. lycopersicum* and wild tomato *S. habrochaites* (Supplementary Fig. 6a). We subsequently constructed *CaMV* 35S promoter-driven *SIWRKY34* and *ShWRKY34* overexpressing lines (OE), as well as *slwrky34* mutants through CRISPR/Cas9-mediated techniques in LA4024 background. Under normal conditions, the *wrky34* mutants exhibited smaller fruits and fewer seeds per fruit compared to wild-type (WT) (Supplementary Fig. 7). Moreover, tissue-specific expression analysis revealed that tomato *WRKY34* is prominently expressed in roots, followed by flowers and buds, with lower expression levels observed in leaves, and the lowest expression in fruits (Supplementary Fig. 8a). Consistently, *WRKY34* protein accumulation was highest in roots and lowest in fruits (Supplementary Fig. 8b). Interestingly, both overexpression of *SIWRKY34* and *ShWRKY34* compromised seedlings cold tolerance, evidenced by higher REL, lower *Fv/Fm* and lower survival rate compared to WT (Supplementary Figs. 5b and 6b–d). Conversely, *slwrky34* mutants exhibited extreme tolerance to cold stress, displaying lower REL, higher *Fv/Fm* and higher survival rate than WT seedlings (Supplementary Figs. 5b and 6b–d). Additionally, overexpression of *WRKY34* significantly suppressed the expression of cold responsive genes *CBFs* and *CORs* under cold stress, whereas the knockout of *WRKY34* promoted the expression of these genes under cold stress (Supplementary Fig. 6e). Hence, both *SIWRKY34* and *ShWRKY34* negatively regulate tomato cold tolerance.

Next, we concluded that differences in expression patterns of *WRKY34s* under cold stress, rather than their protein function, dictated the disparity in cold tolerance between wild and cultivated tomatoes. To explore the reasons behind the differential expression patterns and chromatin accessibilities of *WRKY34* between wild and cultivated tomatoes under cold stress, we amplified and sequenced the 3000 bp length promoter of *SIWRKY34* and the 3019 bp length promoter of *ShWRKY34* (Supplementary Fig. 9). Evidently, compared to the *SIWRKY34* promoter of cultivated tomato, we identified a 60 bp insertion at -2315 bp upstream of the *ShWRKY34* translation initiation site (ATG) in wild tomato, precisely situated in the opening chromatin regions of the *ShWRKY34* promoter under cold stress in wild tomato LA1777 (Fig. 3a and Supplementary Fig. 9). Two *cis*-elements, W-box and GATA-box, were identified in the 60 bp InDel by PLANTCARE software (Supplementary Fig. 9). To verify whether the 60 bp InDel of the *WRKY34* promoter is accountable for the variation in its expression levels under cold stress, we performed a dual-luciferase (LUC) transcriptional activation assay in tobacco. Tobacco leaves were transformed with constructs containing the *LUC* reporter gene driven by *SIWRKY34* and *ShWRKY34* promoters. Under cold stress, the *SIWRKY34* promoter (*pSIW34*) did not obviously affect LUC activity, whereas the *ShWRKY34* promoter (*pShW34*) significantly inhibited the expression of *LUC* reporter gene. Insertion of the 60 bp InDel at position -2350 in the context of the *SIWRKY34* promoter (*pSIW34<sup>+60bp</sup>*) led to a significant decrease in LUC activity after cold stress (Fig. 3b, c). To determine whether two *cis*-elements are involved in cold-suppressed *WRKY34* expression, we mutated W-box and GATA-box in the 60 bp InDel of the *ShWRKY34* promoter (*pShW34<sup>mW-box</sup>* and *pShW34<sup>mGATA-box</sup>*) and fused them to *LUC* reporter constructs, respectively (Fig. 3b). The results revealed that *pShW34<sup>mGATA-box</sup>* restored cold-suppressed LUC activity, whereas *pShW34<sup>mW-box</sup>* still exhibited significant suppression of LUC activity after cold stress (Fig. 3c), suggesting that the *cis*-element



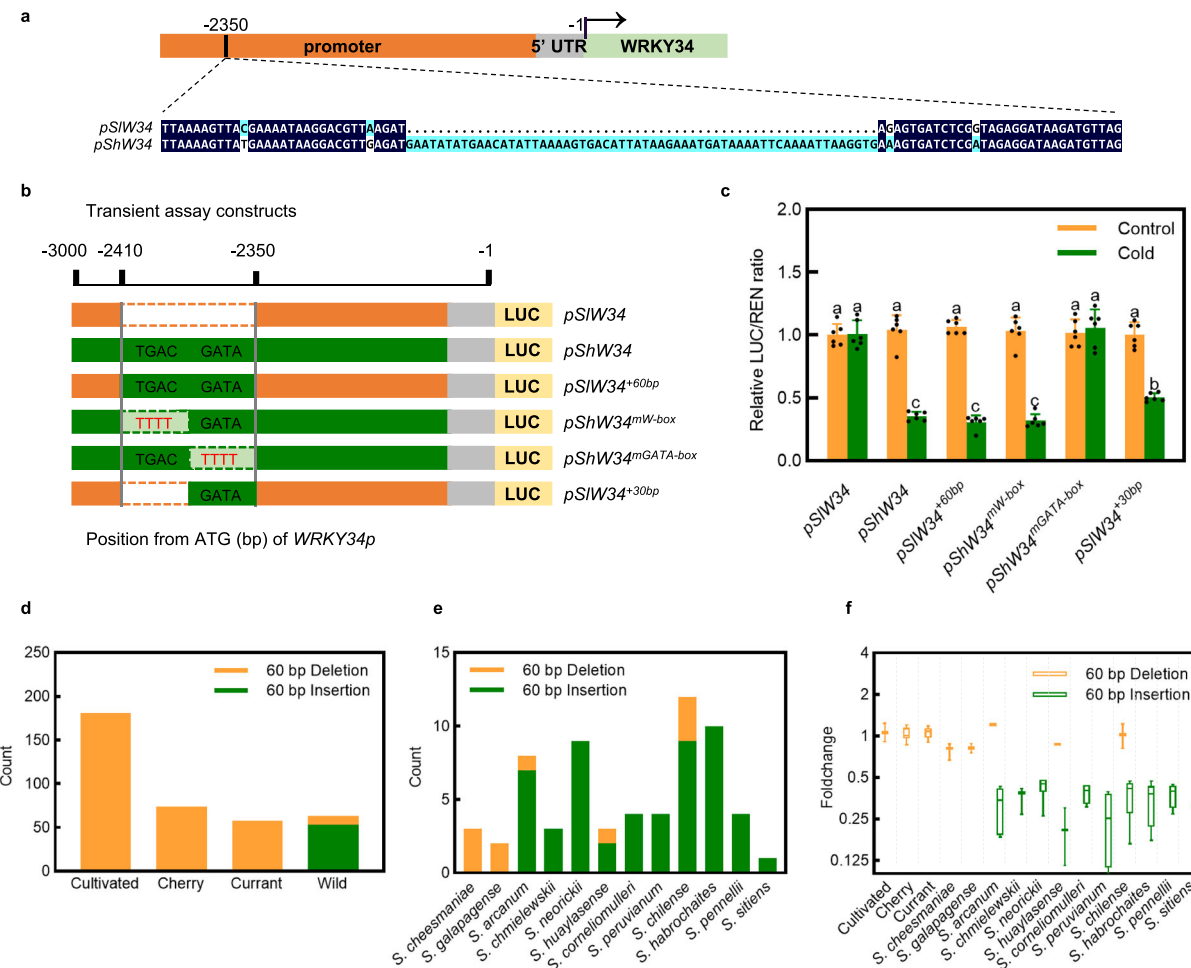
GATA-box within the 60 bp InDel of the *ShWRKY34* promoter plays a crucial role in suppressing *ShWRKY34* expression under cold stress. Furthermore, we constructed a *pSIW34<sup>+30bp</sup>/LUC* fusion vector, wherein only a 30 bp InDel containing the GATA-box was inserted at position -2350 in the context of the *SIWRKY34* promoter, and measured LUC activity. Interestingly, compared with *pShW34*, *pSIW34<sup>+60bp</sup>* and *pShW34<sup>60bp-box</sup>*, the LUC activity of *pSIW34<sup>+30bp</sup>* was only partially reduced

under cold stress, suggesting that both the *cis*-element GATA-box and the entire 60 bp InDel fragment are important for the suppression of *WRKY34* transcription after cold stress (Fig. 3c).

To further explore whether the 60 bp InDel has been influenced and selected by evolution and domestication, we analyzed the variation of the 60 bp InDel in 181 cultivated tomatoes (*S. lycopersicum*), 74 cherry tomatoes (*S. lycopersicum* var. *cerasiforme*), 58 currant

**Fig. 2 | Functional identification of *WRKY34* alleles in response to cold stress in cultivated and wild tomatoes.** **a** Expression levels of *WRKY34* in introgression line LA3942, its background material LA4024, donor parent LA1777 and their *WRKY34*-silenced seedlings under cold stress. Total RNA was isolated from leaf samples after 6 h of cold stress. Data are presented as means  $\pm$  SD of three biological replicates. The relative expression levels of *WRKY34* in TRV of LA4024, LA3942 and LA1777 under normal conditions were set to “1”. **b** Phenotypes of LA3942, LA4024, LA1777 and their *WRKY34*-silenced seedlings at 7 d after cold treatment. Eight plants of each genotype and treatment were tested with similar results. Bar: 10 cm. Relative electrolyte leakage (REL, **c**) and maximum photochemical efficiency of photosystem II (*Fv/Fm*, **d**) in LA4024, LA3942, LA1777 and their *WRKY34*-silenced seedlings at 7 d after cold treatment. The false color code depicted at the bottom of the

images ranges from 0 (black) to 1 (purple). Bar: 2 cm. Data are presented as means of four biological replicates  $\pm$  SD (**c**) or means of eight leaflets from independent plants (**d**). **e** Expression levels of *CBFs* and *CORs* in LA4024, LA3942, LA1777 and their *WRKY34*-silenced seedlings under cold stress. Total RNA was isolated from leaf samples after 6 h of cold stress. The relative expression levels of *CBFs* and *CORs* in TRV of LA4024, LA3942 and LA1777 under normal conditions were set to “1”. At least twice experiments were repeated independently with similar results. Data are presented as means  $\pm$  SD of three biological replicates. Different letters indicate significant differences ( $P < 0.05$ ) according to one-way ANOVA with Duncan’s multiple range test. TRV, non-silenced seedlings; TRV-*SIWRKY34*, *SIWRKY34*-silenced seedlings; TRV-*ShWRKY34*, *ShWRKY34*-silenced seedlings. Source data are provided as a Source Data file.



**Fig. 3 | The 60 bp InDel in the *WRKY34* promoter results in different expression patterns of *WRKY34* under cold stress.** **a** *WRKY34* gene structure in cultivated and wild tomatoes. The gray box represents 5' UTR, the green box represents the coding sequence, the orange box represents the promoter and the black line in the promoter represents the 60 bp InDel region. **b** Schematic diagram of the reporter constructs with the full-length promoters of *WRKY34* or different mutated *WRKY34* promoters fused to the *LUC* reporter gene in the vector pGreenII 0800-LUC, while the internal control *REN* reporter gene was driven by the *CaMV 35S* promoter. *pSIW34*, full-length promoter of *SIWRKY34*; *pShW34*, full-length promoter of *ShWRKY34*; *pSIW34+60bp*, *SIWRKY34* full-length promoter inserts a 60 bp InDel from *ShWRKY34* promoter; *pShW34<sup>mW-box</sup>*, *ShWRKY34* full-length promoter mutants W-box in the 60 bp InDel; *pShW34<sup>mGATA-box</sup>*, *ShWRKY34* full-length promoter mutants GATA-box in the 60 bp InDel; *pSIW34+30bp*, *SIWRKY34* full-length promoter inserts a

30 bp InDel from *ShWRKY34* promoter. **c** LUC relative activities driven by different promoters were determined in transient transgenic tobacco plants 6 h after cold stress. The LUC relative activity driven by *pSIW34* in transient expressed tobacco leaves under normal conditions was set to “1”. Data are presented as means  $\pm$  SD of six biological replicates. The experiment was repeated three times with similar results. Different letters indicate significant differences ( $P < 0.05$ ) according to one-way ANOVA with Duncan’s multiple range test. **d** The 60 bp InDel of 181 cultivated tomatoes, 74 cherry tomatoes, 58 currant tomatoes and 63 wild tomatoes. **e** The 60 bp InDel of 63 different wild tomatoes. **f** The foldchange of *WRKY34* expression levels under cold stress in different tomato varieties. The horizontal line in the middle indicates mean. The experiment used three biological replicates in one experiment and was repeated twice with similar results. Source data are provided as a Source Data file.

tomatoes (*S. pimpinellifolium*) and 63 wild tomatoes including 3 *S. cheesmaniae*, 2 *S. galapagense*, 8 *S. arcanum*, 3 *S. chmielewskii*, 9 *S. neorickii*, 3 *S. huaylasense*, 4 *S. corneliomulleri*, 4 *S. peruvianum*, 12 *S. chilense*, 10 *S. habrochaites*, 4 *S. pennellii* and 1 *S. sitiens* accessions

(Supplementary Data 6). Surprisingly, we observed no 60 bp insertion in *WRKY34* promoters across all cultivated tomatoes, cherry tomatoes, currant tomatoes, and two wild tomato species, *S. cheesmaniae* and *S. galapagense* accessions. However, the 60 bp insertions were prevalent

in 91.4% (53 out of 58) of other wild tomatoes, including all *S. chmielewskii*, *S. neorickii*, *S. corneliomulleri*, *S. peruvianum*, *S. habrochaites*, *S. pennellii*, and *S. sitiens* accessions, as well as 87.5% (7 out of 8) *S. arcanum*, 75% (9 out of 12) *S. chilense*, and 66.7% (2 out of 3) *S. huaylasense* accessions (Fig. 3d, e and Supplementary Data 6). To verify the relationship between the 60 bp InDel and *WRKY34* expression in response to cold stress, we selected a subset of cultivated tomatoes, cherry tomatoes, currant tomatoes and all wild tomatoes to measure the expression levels of *WRKY34* at 6 h after cold treatment. The results showed that the *WRKY34* variants harboring the 60 bp deletion exhibited no significant alteration in expression after cold stress. In contrast, the expression of *WRKY34* variants with the 60 bp insertion demonstrated a notable decrease after cold stress (Fig. 3f and Supplementary Fig. 10). These results suggest that the 60 bp InDel is significantly associated with cold-suppressed *WRKY34* expression in all wild and cultivated tomatoes, and the 60 bp insertion in the *WRKY34* promoter is prevalent in wild tomatoes, yet it has progressively vanished during the extended evolutionary transition to currant tomatoes.

### SWIBs and GATA29 directly bind to the 60 bp InDel fragment of the *WRKY34* promoter

To explore how the 60 bp InDel fragment represses *WRKY34* expression, we employed a yeast one-hybrid (Y1H) screen using the 60 bp InDel region as bait DNA. Prey proteins from a tomato complementary DNA library, fused with the yeast GAL4 transcription activation domain (GAL4 AD), were screened. Out of 297 putative DNA-binding proteins identified, we chose a GATA family transcription factor, SIGATA29 capable of binding to the GATA-box, and two SWIB/MDM2 domain proteins, SISWIBa and SISWIBb, known to influence chromatin opening (Supplementary Data 7), for further analysis.

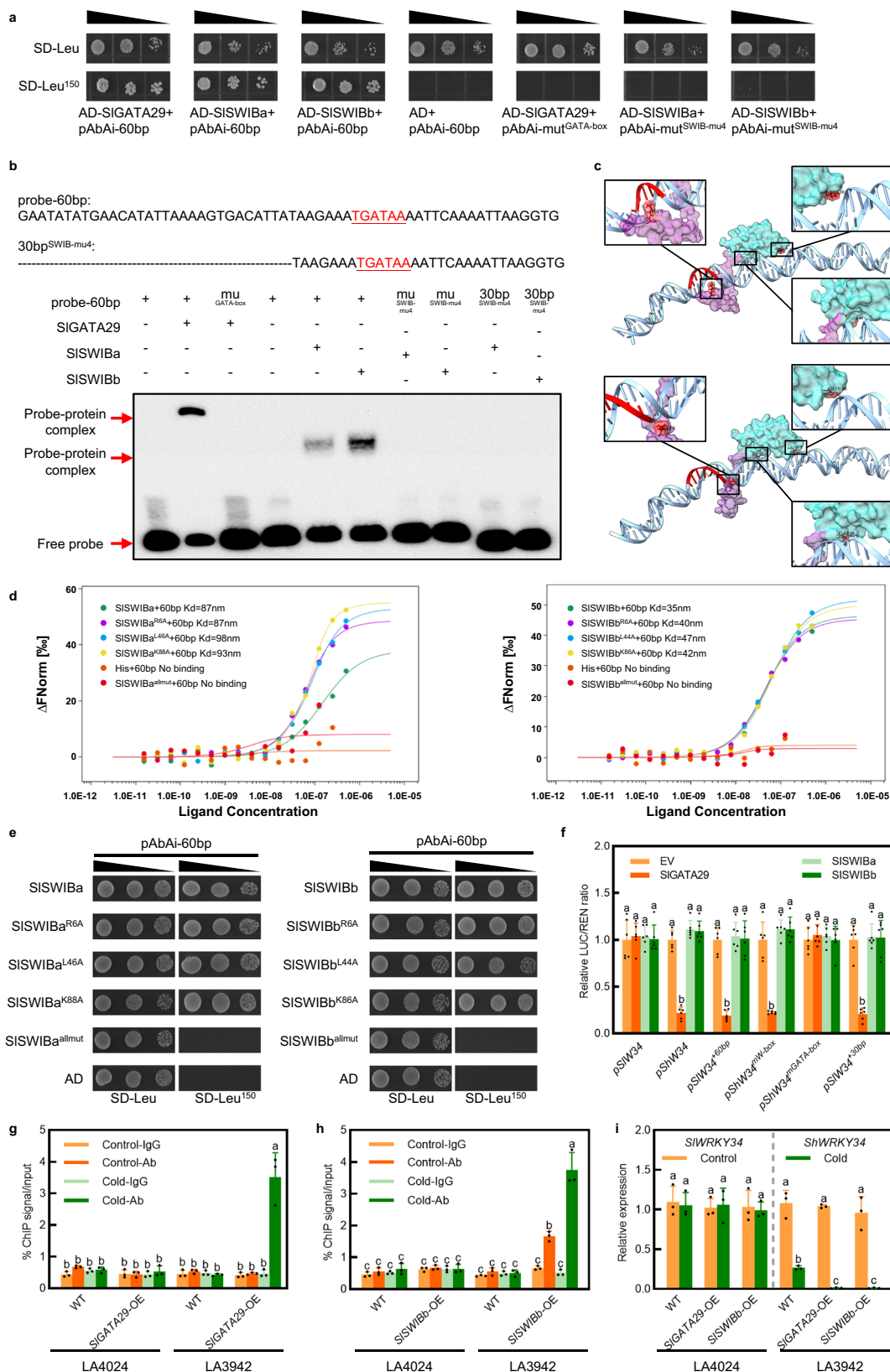
Using full-sequence constructs of these genes, we performed gene-specific Y1H assays to determine their specific binding to the aforementioned 60 bp InDel. Yeast cells containing the 60 bp bait vector and either the pGADT7-SIGATA29 vector or pGADT7-SISWIBa/b vectors grew on the SD-Leu media with 150 ng ml<sup>-1</sup> aureobasidin A (AbA) (SD-Leu<sup>150</sup>). Conversely, transformants lacking SIGATA29 or SISWIBa/b failed to grow on this media (Fig. 4a). To identify the core DNA binding sites of SWIB/MDM2 domain proteins SISWIBa/b, we created six different 60 bp mutation probes with various mutation sites (mu1-mu6) and performed electrophoretic mobility shift assay (EMSA) and microscale thermophoresis (MST). The results revealed mu4 (TGATAA) as the common core DNA binding site of SISWIBa and SISWIBb, consistent with GATA-box; hence, we named it as SWIB-mu4 (Supplementary Fig. 11). Transformants with mutated GATA-box or SWIB-mu4 (mut<sup>GATA-box</sup> or mut<sup>SWIB-mu4</sup>) bait vectors did not grow on SD-Leu<sup>150</sup> media (Fig. 4a), indicating that SIGATA29 and SISWIBa/b could bind to the GATA-box and SWIB-mu4 of the 60 bp InDel in yeast, respectively. EMSA results further confirmed that SIGATA29 and SISWIBa/b bound to the GATA-box and SWIB-mu4 in the 60 bp InDel, respectively (Fig. 4b). However, SISWIBa/b could not bind to a 30 bp probe with a complete SWIB-mu4, indicating that the binding of SISWIBa/b to the 60 bp InDel requires not only the SWIB-mu4 but also the full-length 60 bp DNA fragment (Fig. 4b). Previous studies have primarily considered SWIB/MDM2 domain proteins as regulators of chromatin accessibility through their influence on nucleosomes<sup>33</sup>. To date, there has been no report on the direct binding of SWIB/MDM2 proteins to chromatin DNA. To further identify the key amino acid sites in SWIBs for binding to the 60 bp InDel, we made an accurate prediction of protein-nucleic acid complexes using RoseTTAFoldNA<sup>34</sup>. Predicted results suggested that three conserved amino acids in two SISWIB homologous proteins, including Arg6 in both SISWIBa and SISWIBb, Leu46 in SISWIBa and Leu44 in SISWIBb, and Lys88 in SISWIBa and Lys86 in SISWIBb, might be involved in binding to the 60 bp InDel through hydrogen bonds (Fig. 4c). To verify the key

binding roles of these amino acids, we separately mutated these three amino acids and also mutated all three amino acids together, then tested their binding with the 60 bp InDel. Both MST and Y1H results showed that mutating any single amino acid still allowed SISWIBa/b to bind to the 60 bp InDel, but when all three amino acids were mutated to alanine, they no longer bound to the 60 bp InDel in vitro (Fig. 4d, e). These results further indicate that SWIBs have the ability to directly bind to DNA, and we infer that they would form a helical protein to wrap around DNA, with SWIB-mu4 being the key recognition or binding site.

To investigate the effect of SIGATA29 and SISWIBa/b on downstream *WRKY34* gene expression, we performed dual-LUC assays and found that SIGATA29 suppressed LUC activity when *WRKY34* promoters contained the GATA-box (*pShW34*, *pSIW34<sup>+60bp</sup>*, *pShW34<sup>mW-box</sup>* and *pSIW34<sup>+30bp</sup>*), but had little effect on *WRKY34* promoters without the GATA-box (*pSIW34* and *pShW34<sup>mGATA-box</sup>*). Interestingly, when the reporter gene was co-transfected with *SISWIBa* or *SISWIBb* gene, the LUC activity of all *WRKY34* promoters did not change, suggesting that SISWIBs do not directly regulate *WRKY34* expression (Fig. 4f). To verify the binding of SIGATA29 and SISWIBs to the 60 bp InDel in vivo, we overexpressed SIGATA29 and SISWIBb in cultivated tomato LA4024 and introgression line LA3942, respectively. ChIP-qPCR analysis showed that SIGATA29 and SISWIBb could not bind to the *ShWRKY34* promoter of LA4024 with or without cold treatment, due to the absence of the 60 bp InDel. SIGATA29 could not bind to the *ShWRKY34* promoter of LA3942 under normal conditions, but could bind to its promoter under cold stress (Fig. 4g). At normal temperature, SISWIBb could directly bind to the *ShWRKY34* promoter of LA3942, and the binding was further increased after cold stress (Fig. 4h). Moreover, overexpression of SIGATA29 or SISWIBb in LA3942 background suppressed the expression of *ShWRKY34* under cold stress, while the expression of *SIWRKY34* in LA4024 background did not respond to cold stress (Fig. 4i). Additionally, we compared the protein sequences of GATA29 and SWIBs between cultivated tomato *S. lycopersicum* and wild tomato *S. habrochaites*. The comparison results showed only one amino acid difference between SIGATA29 and ShGATA29 (Supplementary Fig. 12a). Compared with two homologous SISWIBa and SISWIBb, only one ShSWIB was identified in *S. habrochaites* LA1777. The protein sequence homology of SISWIBa and SISWIBb was 68.6%, and the protein sequence homology of ShSWIB and SISWIBb was 99.2% with only one amino acid difference (Supplementary Fig. 12b). Y1H and EMSA results also demonstrated that ShGATA29 and ShSWIB directly bind the 60 bp InDel of *WRKY34* promoter by specifically interacting with GATA-box and SWIB-mu4, respectively. Additionally, the binding of ShSWIB to the 60 bp InDel also requires a full-length 60 bp DNA (Supplementary Fig. 12c, d). Moreover, dual-LUC assays indicated that ShGATA29 suppressed LUC activity when *WRKY34* promoters contained the GATA-box (*pShW34*, *pSIW34<sup>+60bp</sup>*, *pShW34<sup>mW-box</sup>* and *pSIW34<sup>+30bp</sup>*), but had little effect on *WRKY34* promoters lacking the GATA-box (*pSIW34* and *pShW34<sup>mGATA-box</sup>*). However, the LUC activity derived from all *WRKY34* promoters did not exhibit LUC suppression when the reporter was co-transfected with ShSWIB (Supplementary Fig. 12e). These findings suggest that both the transcription factor GATA29 and the SWIB/MDM2 domain protein SWIBs, found in both wild and cultivated tomatoes, can bind to the 60 bp InDel fragment of the *WRKY34* promoter.

### SWIB and GATA29 synergistically suppress *WRKY34* expression through the 60 bp InDel fragment

Chromatin remodeling factors are typically recruited to target genes via specific transcription factors, thereby synergistically regulating the expression of target genes<sup>35</sup>. Interestingly, using yeast two-hybrid (Y2H) assays, we found that SIGATA29 interacted with both SISWIBa and SISWIBb in yeast (Fig. 5a). To further validate this interaction, we performed a glutathione S-transferase (GST) pull-down assay. The



results revealed that SIGATA29-GST successfully pulled down SISWIBa-His or SISWIBb-His, while the negative control GST failed to do so (Fig. 5b). Similarly, a bimolecular fluorescence complementation (BiFC) assay confirmed the interaction of SIGATA29 with SISWIBa and SISWIBb in the nucleus (Fig. 5c). Furthermore, the interaction between SIGATA29 and SISWIBs was verified by co-immunoprecipitation

(Co-IP) assays in tobacco leaves through co-expressing *SIGATA29-HA* and *SISWIBa-GFP* or *SIGATA29-HA* and *SISWIBb-GFP* (Fig. 5d). These results indicate that SIGATA29 interacts with both SISWIBa and SISWIBb in vitro and in vivo.

To investigate the effect of the interaction between SIGATA29 and SISWIBs on downstream *WRKY34* gene expression, we performed

**Fig. 4 | SIGATA29 and SISWIBa/b directly bind to the 60 bp InDel of the WRKY34 promoter and suppress WRKY34 expression under cold stress.** **a** Yeast one-hybrid assay (Y1H) testing the binding of SIGATA29 and SISWIBa/b to pAbAi-60bp in the SD-Leu medium with or without 150 ng ml<sup>-1</sup> AbA. Empty vector containing the AD serves as a negative control. **b** EMSA results showing SIGATA29 and SISWIBa/b directly bind to the 60 bp InDel. Probe sequences are shown (top). mu<sup>GATA-box</sup> and mu<sup>SWIB-mu4</sup>, mutated probes in which the TGATAA motif was changed to AAAAAA. **c** Complex structures of SISWIBa with the 60 bp DNA fragment (upper panel) and SISWIBb with the 60 bp DNA fragment (lower panel). Blue and red indicate the SWIB/MDM2 domain and the predicted binding site, respectively. SWIB-mu4 is highlighted in red. **d** MST assays of the binding of SISWIBa/b and their three amino acid mutations to the 60 bp InDel. Kd, dissociation constant. **e** Y1H testing the binding of SISWIBa/b and their three amino acid mutations to pAbAi-60bp in the SD-Leu medium with or without 150 ng ml<sup>-1</sup> AbA. **f** Transient dual-

luciferase (dual-LUC) assay in tobacco leaves. Empty vector (EV) was included as control. Data are presented as means  $\pm$  SD of six biological replicates. ChIP-qPCR assays showing the binding of SIGATA29 (**g**) and SISWIBb (**h**) to the 60 bp InDel in vivo. Data are presented as means  $\pm$  SD of three biological replicates. **i** Expression levels of WRKY34 in LA4024, LA3942 and SIGATA29 overexpressing (SIGATA29-OE) seedlings and SISWIBb overexpressing (SISWIBb-OE) seedlings in the background of LA4024 or LA3942 under cold stress. Total RNA was isolated from leaf samples after 6 h of cold stress. Data are presented as means  $\pm$  SD of three biological replicates. The relative expression levels of WRKY34 in WT of LA4024 and LA3942 under normal conditions were set to "1". Experiments in (**a, b, d, e**) were repeated twice, and in (**f-i**), three times, all yielding similar results. Different letters above bars indicate significant differences ( $P < 0.05$ ) determined using one-way ANOVA with Duncan's multiple range test. Source data are provided as a Source Data file.

transient dual-LUC assays with different types of *WRKY34* promoters. LUC activity derived from *WRKY34* promoters containing the GATA-box (*pShW34*, *pSIW34<sup>+60bp</sup>*, *pShW34<sup>inW-box</sup>* and *pSIW34<sup>+30bp</sup>*) decreased significantly when co-transfected with *SIGATA29*, and this reduction was further augmented after co-transfection with *SISWIBa* or *SISWIBb*, except for *pSIW34<sup>+30bp</sup>*. Conversely, LUC activity derived from *WRKY34* promoters lacking the GATA-box (*pSIW34* and *pShW34<sup>inGATA-box</sup>*) showed no change when co-transfected with either *SIGATA29* alone or with both *SIGATA29* and *SISWIBs* (Fig. 5e).

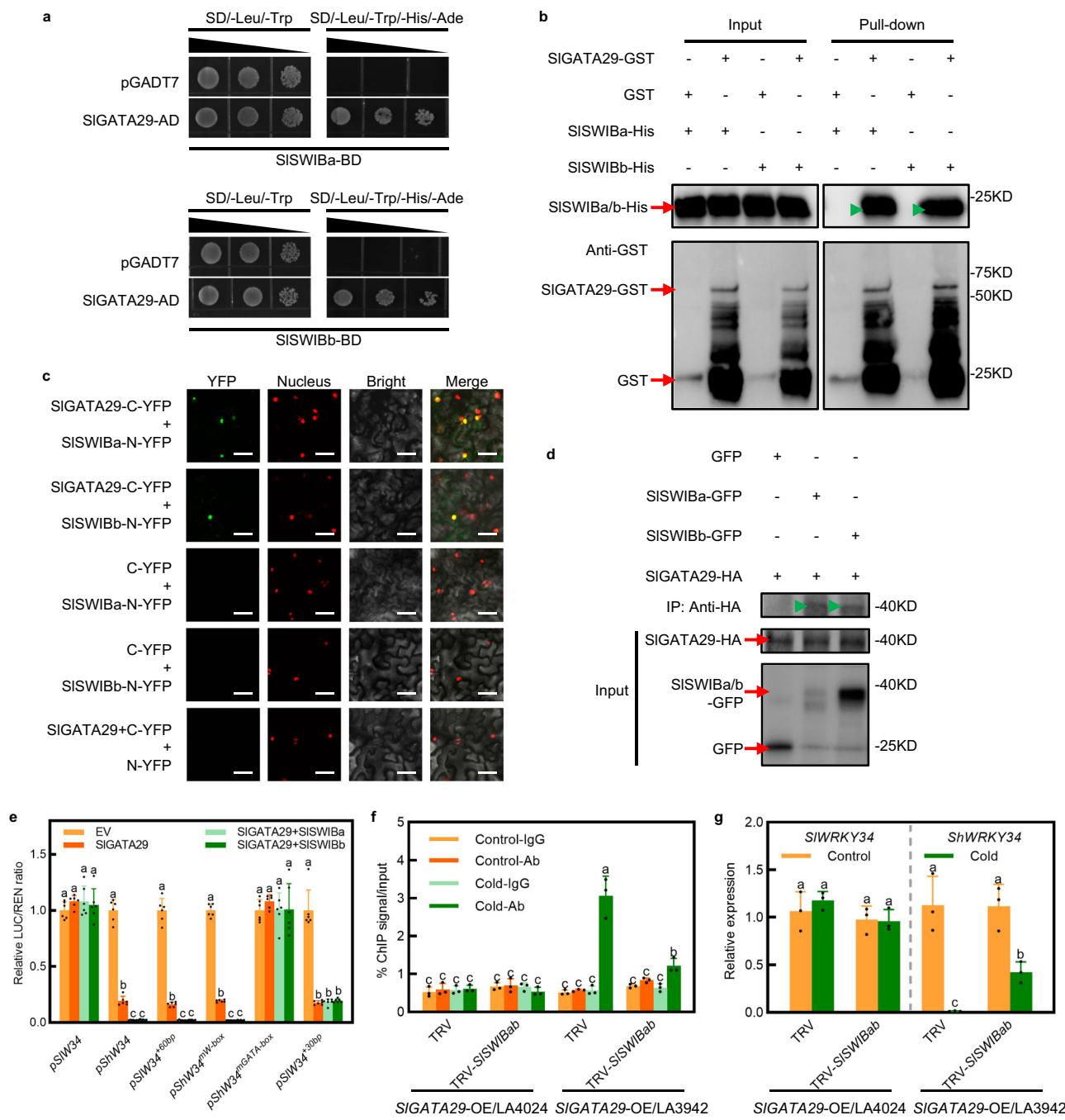
To further examine the synergistic effects of SIGATA29 and SISWIBs, we performed EMSA in vitro. As shown in Supplementary Fig. 13, while SIGATA29 could specifically bind to 60 bp probe containing the GATA-box, the addition of purified SISWIBa or SISWIBb proteins did not significantly affect its binding ability. Therefore, we speculated that the synergistic suppression effect of SIGATA29 and SISWIBa/b on *WRKY34* expression might be linked to chromatin accessibility, and such changes in chromatin accessibility require a eukaryotic environment. As shown in Fig. 5f, in LA4024 where the *ShWRKY34* promoter lacks the 60 bp InDel, there was no binding of SIGATA29 to the *ShWRKY34* promoter in both TRV and *SISWIBa* and *SISWIBb* co-silenced seedlings (TRV-*SISWIBab*) of *SIGATA29* overexpressing lines. Conversely, in LA3942 background, where *ShWRKY34* promoter contains the 60 bp InDel fragment, significant binding of SIGATA29 to the *ShWRKY34* promoter was observed in TRV of *SIGATA29* overexpressing lines. However, when *SISWIBa* and *SISWIBb* were co-silenced, the binding of SIGATA29 to the *ShWRKY34* promoter containing the 60 bp InDel fragment was significantly reduced (Fig. 5f). Correspondingly, the expression of *ShWRKY34* in LA4024 background remained unchanged, whereas the expression of *ShWRKY34* in *SIGATA29* overexpressing lines of LA3942 background was significantly decreased under cold stress. Silencing *SISWIBab* in *SIGATA29* overexpressing lines of LA3942 background significantly alleviated the decrease in *WRKY34* expression under cold stress (Fig. 5g). These results suggest that SISWIBs can enhance the suppression effect of SIGATA29 on *WRKY34* expression, and a complete 60 bp InDel in the *WRKY34* promoter is indispensable for achieving the synergistic suppression effect of SIGATA29 and SISWIBs.

Given the high protein sequence similarity between SIGATA29 and ShGATA29, as well as between SISWIBs and ShSWIB, Y2H also detected the interaction between ShGATA29 and ShSWIB in yeast (Supplementary Fig. 14a). We further verified the interaction of ShGATA29 and ShSWIB by performing pull-down assays in vitro (Supplementary Fig. 14b). BiFC and Co-IP results also confirmed the interaction between ShGATA29 and ShSWIB in vivo (Supplementary Fig. 14c, d). Similarly, we performed dual-LUC assays to investigate the synergistic suppression effect of ShGATA29 and ShSWIB (Supplementary Fig. 14e). The results showed that ShSWIB and ShGATA29 exhibit similar functions with SISWIBs and SIGATA29 in synergistic suppression of *WRKY34* expression.

### Impact of SWIB and GATA29 on chromatin accessibility and cold tolerance via the 60 bp InDel

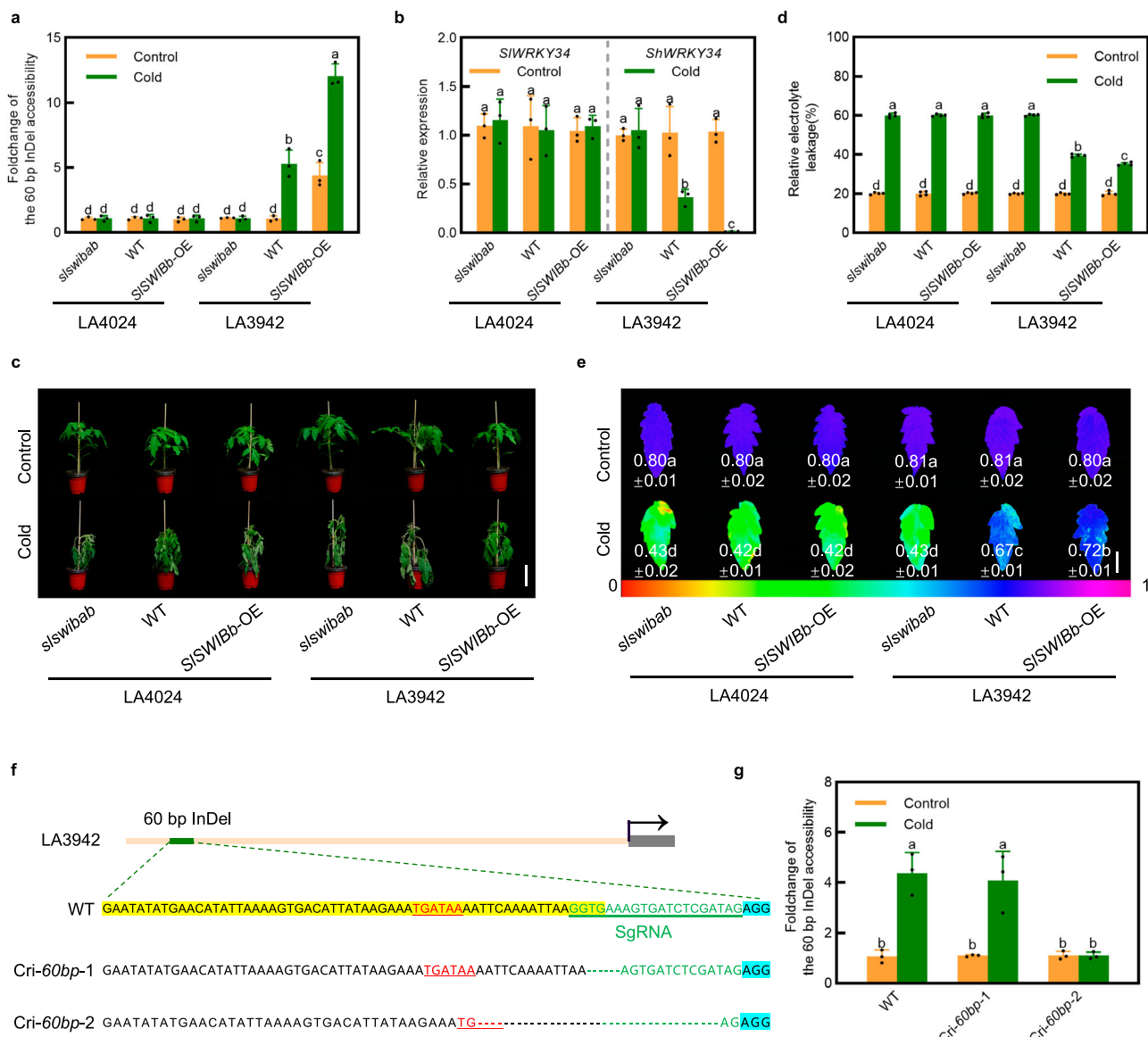
To further elucidate the effects of SWIBs on chromatin accessibility associated with the 60 bp InDel fragment of *WRKY34* promoter, we constructed *SISWIBb* overexpressing lines driven by the *CaMV 35S* promoter (*SISWIBb-OE*) and *sISWIBab* double mutants mediated by CRISPR/Cas9 in the background of LA4024 or LA3942. In an independent ATAC-qPCR assay, the region around the 60 bp InDel fragment of the *WRKY34* promoter exhibited an increase in chromatin accessibility following cold treatment in LA3942 background (Fig. 6a). In contrast, the chromatin accessibility for this region did not change in response to cold stress in LA4024 background. Furthermore, in the *sISWIBab* double mutants of LA3942 background, the chromatin accessibility of this region was also lost in response to cold stress (Fig. 6a). Interestingly, overexpression of *SISWIBb* in LA3942 background significantly increased chromatin accessibility of this region, and this region was more accessible under cold stress in LA3942 *SISWIBb-OE* lines (Fig. 6a). RT-qPCR analysis revealed that *WRKY34* did not respond to cold stress in the background of LA4024, consistent with the results of chromatin accessibility. Meanwhile, *WRKY34* expression in LA3942 seedlings was significantly decreased after cold treatment, while *WRKY34* expression in *sISWIBab* double mutants of LA3942 background remained unchanged regardless of cold treatment (Fig. 6b). Overexpression of *SISWIBb* in LA3942 background did not affect the expression of *WRKY34* under normal conditions. However, the expression of *WRKY34* in *SISWIBb-OE* lines of LA3942 background was significantly suppressed and lower than that in WT LA3942 seedlings after cold stress (Fig. 6b). Consistent with the expression of *WRKY34*, knockout of *SISWIBa* and *SISWIBb* or overexpression of *SISWIBb* in LA4024 background had no effect on the cold tolerance of LA4024, as indicated by similar REL, *Fv/Fm* and survival rate (Fig. 6c-e and Supplementary Fig. 5c). However, *sISWIBab* double mutants of LA3942 exhibited compromised cold tolerance compared to WT, with higher REL, lower *Fv/Fm* and lower survival rate. In contrast, overexpression of *SISWIBb* enhanced the cold tolerance of LA3942 resulting in lower REL, higher *Fv/Fm* and higher survival rate (Fig. 6c-e and Supplementary Fig. 5c).

We also constructed *SIGATA29* overexpressing lines driven by the *CaMV 35S* promoter (*SIGATA29-OE*) and *sIGATA29* mutants mediated by CRISPR/Cas9 in the background of LA4024 or LA3942. As shown in Supplementary Fig. 15a, the expression of *WRKY34* showed no response to cold stress in LA4024 background. In contrast, the expression of *WRKY34* in LA3942 background was significantly suppressed under cold stress, while cold-suppressed *WRKY34* expression was compromised when *SIGATA29* was knocked out in LA3942 background (Supplementary Fig. 15a). Conversely, overexpression of *SIGATA29* further suppressed *WRKY34* expression in LA3942 background under cold stress (Supplementary Fig. 15a). Consistent with the expression of *WRKY34*, *SIGATA29* knockout or overexpression in LA4024 background had no effect on the cold tolerance of LA4024, as indicated by similar REL, *Fv/Fm* and survival rate (Supplementary



**Fig. 5 | SISWIBa/b and SIGATA29 synergistically suppress WRKY34 expression.** **a** Yeast two-hybrid assay (Y2H) testing the interactions of SIGATA29 and SISWIBa/b in yeast. **b** Pull-down assays show that GST-tagged SIGATA29 physically interacts with His-tagged SISWIBa or His-tagged SISWIBb. The green triangle indicates the target band of SISWIBa-His or SISWIBb-His pulled down by SIGATA29-GST. The combination of GST and SISWIBa-His or SISWIBb-His was used as a negative control. **c** Bimolecular fluorescence complementation assay (BiFC) showing the interactions between SIGATA29 and SISWIBa/b in tobacco. Bar: 50  $\mu$ m. **d** Co-IP showing the interactions between SIGATA29 and SISWIBa/b. The green triangle indicates the SIGATA29-HA target band immunoprecipitated with SISWIBa-GFP or SISWIBb-GFP. The combination of GFP and SIGATA29-HA was used as a negative control. **e** Transient dual-luciferase (dual-LUC) assay in tobacco leaves. Empty vector (EV) was included as control. Data are presented as means  $\pm$  SD of six biological replicates. **f** ChIP-qPCR assays show that silencing SISWIBab affects the binding of

SIGATA29 to the 60 bp InDel in vivo. Data are presented as means  $\pm$  SD of three biological replicates. **g** Expression levels of WRKY34 in SIGATA29 overexpressing lines of LA4024 or LA3942 background and their SISWIBab-silenced seedlings under cold stress. Total RNA was isolated from leaf samples after 6 h of cold stress. Data are presented as means  $\pm$  SD of three biological replicates. The relative expression levels of WRKY34 in TRV of SIGATA29 overexpressing lines of LA4024 or LA3942 background under normal conditions were set to "1". In (a-d), experiments were repeated twice with similar results. In (e-g), experiments were repeated three times with similar results. Different letters above bars indicate significant differences ( $P < 0.05$ ) determined using one-way ANOVA with Duncan's multiple range test. TRV, non-silenced seedlings; TRV-SISWIBab, SISWIBa and SISWIBb co-silenced seedlings; SIGATA29-OE/LA4024, SIGATA29 overexpressing lines in LA4024 background; SIGATA29-OE/LA3942, SIGATA29 overexpressing lines in LA3942 background. Source data are provided as a Source Data file.



**Fig. 6 |** *SISWIBa/b* enhances chromatin opening of the *WRKY34* promoter for improved tomato cold tolerance, dependent on the 60 bp InDel. **a** ATAC-qPCR results from testing chromatin accessibility in the 60 bp InDel region of the *WRKY34* promoter under cold stress in *slswbab* double mutants and *SISWIBb* over-expressing (*SISWIBb*-OE) seedlings of LA4024 or LA3942 background. Data are presented as means  $\pm$  SD of three biological replicates. **b** Expression levels of *WRKY34* in *slswbab* double mutants and *SISWIBb*-OE seedlings of LA4024 or LA3942 background under cold stress. Total RNA was isolated from leaf samples after 6 h of cold stress. Data are presented as means  $\pm$  SD of three biological replicates. The relative expression levels of *WRKY34* in WT of LA4024 and LA3942 under normal conditions were set to “1”. **c** Phenotypes of *slswbab* double mutants and *SISWIBb*-OE seedlings in LA4024 or LA3942 background at 7 d after cold treatment. Eight plants of each genotype and treatment were tested with similar results. Bar: 10 cm. **d** Relative electrolyte leakage (REL, **d**) and maximum photochemical efficiency of photosystem II (*Fv/Fm*, **e**) in *slswbab* double mutants and

*SISWIBb*-OE seedlings of LA4024 or LA3942 background at 7 d after cold treatment. The false color code depicted at the bottom of the images ranges from 0 (black) to 1 (purple). Bar: 2 cm. Data are presented as means of four biological replicates  $\pm$  SD (**d**) or means of eight leaflets from independent plants (**e**). **f** CRISPR/Cas9-induced deletions in the 60 bp InDel region of the *WRKY34* promoter of LA3942. Blue-marked area represents PAM, green-marked area indicates SgRNA, yellow-marked area represents the 60 bp InDel, and red-marked area represents the SWIB-mu4/GATA-box. **g** ATAC-qPCR results from testing chromatin accessibility in the 60 bp InDel region of the *WRKY34* promoter in response to cold stress in different types of 60 bp InDel deletion mutants. Data are presented as means  $\pm$  SD of three biological replicates. All experiments were repeated twice with similar results. Different letters above bars indicate significant differences ( $P < 0.05$ ) determined using one-way ANOVA with Duncan’s multiple range test. Source data are provided as a Source Data file.

Figs. 5d and 15b-d). However, *slgata29* mutants of LA3942 exhibited compromised cold tolerance compared to WT, as indicated by higher REL, lower *Fv/Fm* and lower survival rate. In contrast, overexpression of *SIGATA29* enhanced cold tolerance of LA3942, resulting in lower REL, higher *Fv/Fm* and higher survival rate (Supplementary Figs. 5d and 15b-d).

To explore the critical role of the 60 bp InDel in altering chromatin accessibility, we used the CRISPR/Cas9 to delete the 60 bp in the

*ShWRKY34* promoter of LA3942. We obtained two lines (named *Cri-60bp-1* and *Cri-60bp-2*) with different deletions within the 60 bp InDel region (Fig. 6f). Specifically, *Cri-60bp-1* exhibits a deletion of only 6 bp at the 3' end of the 60 bp InDel, whereas *Cri-60bp-2* has a more extensive deletion of 35 bp, which disrupts the crucial SWIB-mu4/GATA-box (Fig. 6f). As expected, ATAC-qPCR assays showed that compared with WT, the chromatin accessibility of the 60 bp InDel region was significantly weakened in *Cri-60bp-2* under cold stress,

while the chromatin accessibility of this region in *Cri-60bp-1* was almost unchanged (Fig. 6g). Moreover, the repression effect of *WRKY34* expression under cold stress was abolished in *Cri-60bp-2*, but not in *Cri-60bp-1* (Supplementary Fig. 16a). Consistently, *Cri-60bp-2* plants were more sensitive to cold stress than WT, exhibiting a significant increase in REL and a decrease in *Fv/Fm* under cold stress (Supplementary Fig. 16b–d). These observations confirm that the 60 bp InDel, regulated by SWIB and GATA29, is a major causal variation underlying the differential expression and chromatin accessibility of *WRKY34* under cold stress.

### WRKY34 disrupts the CBF-COR cold response pathway at both transcript and protein levels

*WRKY34* overexpression or mutation can influence the expression of *CBFs* and *CORs* in LA4024 after cold stress (Supplementary Fig. 6e). To further analyze whether *WRKY34* affects the CBF-COR cold response pathway, we detected the protein interaction between *WRKY34* and *CBFs* or *CORs*. Interestingly, *SIWRKY34* can interact with *SICBF1* but not *SICBF2/3* or the other three *SICORs* with its C terminal domain (Fig. 7a). Moreover, the C terminal domain of *ShWRKY34* also interacted with *ShCBF1* or *SICBF1*, but not with *ShCBF2/3*, *ShCORs*, *SICBF2/3* and *SICORs* (Supplementary Fig. 17). Using GST-pull down assays, we demonstrated that *SIWRKY34*-GST pulled down *SICBF1*-His, while the negative control GST failed to do so (Fig. 7b). BiFC results showed that the full-length *SIWRKY34* and *SICBF1* proteins interacted in the nucleus (Fig. 7c). Co-IP results revealed that *SIWRKY34*-HA associated with *SICBF1*-GFP, but not with free GFP (Fig. 7d). These results confirm the interaction between *SIWRKY34* and *SICBF1*.

Many previous studies have shown that the CBF-COR pathway is central to plant cold tolerance<sup>36</sup>. We hypothesized that *SIWRKY34* interferes with the transcriptional function of *SICBF1* under cold stress by interacting with *SICBF1*, thus weakening the cold tolerance of cultivated tomato. To test this hypothesis, we conducted EMSA and dual-LUC assays. As shown in Supplementary Fig. 18a, several CBF binding elements, known as Dehydration-Responsive Elements (DREs), were identified in the promoters of *SICBF1* and *SICOR47*. *SICBF1* directly bound to DRE elements in *SICBF1* and *SICOR47* promoters in vitro; however, *SICBF1*-bound probe signals decreased progressively with increasing concentrations of *SIWRKY34* purified proteins (Fig. 7e). Additionally, *SICBF1* could transcriptionally activate the expression of itself and *SICOR47*, while *SIWRKY34* co-transfected with *SICBF1* significantly impaired *SICBF1*-induced *SICBF1* and *SICOR47* expression (Fig. 7f). Furthermore, silencing *SICBF1* in *slwrky34* mutants significantly reduced cold tolerance with higher REL and lower *Fv/Fm* under cold stress, compared with *slwrky34* mutants (Fig. 7g–i).

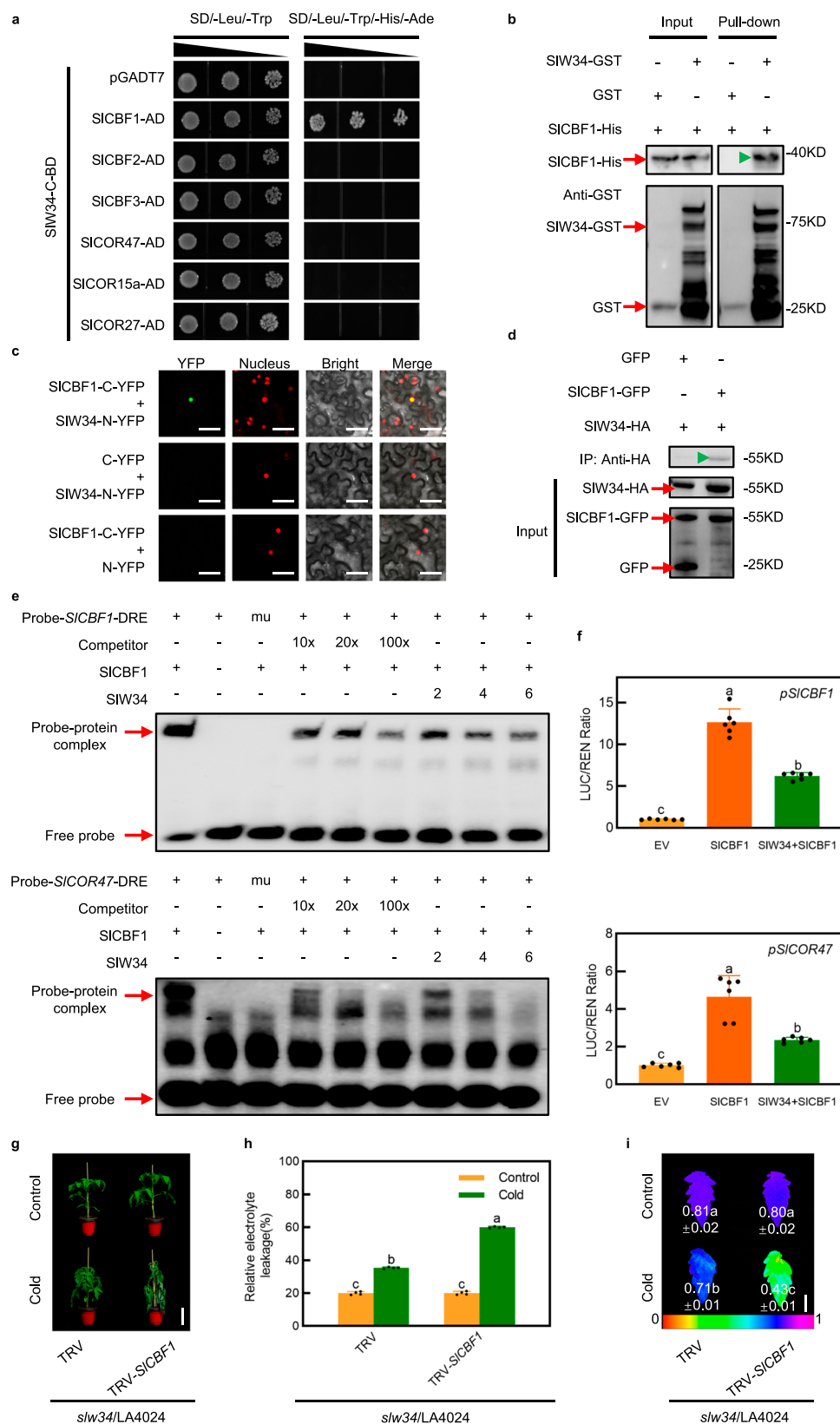
Moreover, we identified many W-box elements in *SICBFs* and *SICOR47* promoters (Supplementary Fig. 18a). Next, we conducted Y1H and ChIP-qPCR assays, respectively. Y1H results showed that *SIWRKY34* could directly bind to the promoters of *SICBFs* and *SICOR47* in yeast cells (Supplementary Fig. 18b). ChIP-qPCR analysis showed that *SIWRKY34* could directly bind to the promoters of *SICBFs* and *SICOR47* in vivo under cold stress (Supplementary Fig. 18c). Furthermore, dual-LUC results indicated that *SIWRKY34* could transcriptionally repress the expression of *SICBFs* and *SICOR47* (Supplementary Fig. 18d). Thus, *WRKY34* also directly suppresses gene transcription in the CBF-COR pathway.

### Discussion

The tomato likely originated in the Andes mountains of South America and adjacent tropical regions<sup>37</sup>. The diverse range of climatic and ecological conditions present across these areas has been instrumental in driving the diversification of tomatoes and their botanical relatives. Phylogenetic analyses have classified wild tomatoes into several groups: “*Lycopersicon* group” (*S. pimpinellifolium*, *S. cheesmaniae*, and

*S. galapagense*), “Arcanum group” (*S. arcanum*, *S. chmielewskii*, and *S. neorickii*), “*Eriopersicon* group” (*S. habrochaites*, *S. huaylasense*, *S. corneliomulleri*, *S. peruvianum*, and *S. chilense*), “*Neolycopersicon* group” (*S. pennellii*); and two outgroups: Section *Juglandifolia* (*S. juglandifolium* and *S. ochranthum*) and Section *Lycopersicoides* (*S. lycopersicoides* and *S. sitiens*)<sup>38</sup>. Each group adapts to specific altitudes and average temperatures, reflecting the influence of environmental factors on their evolutionary paths<sup>38</sup>. In this study, we found a significant correlation between a 60 bp InDel in the *WRKY34* promoter and *WRKY34* expression under cold stress (Fig. 3). Specifically, the *WRKY34* promoter in cultivated tomatoes, cherry tomatoes and “*Lycopersicon* group” of wild tomatoes exhibited this 60 bp deletion, while over 90% of other wild tomato groups contained the 60 bp insertion (Fig. 3d, e). Heatmap analysis revealed that the expression of *WRKY34* variant with the 60 bp deletion showed no significant change post cold stress, whereas *WRKY34* variant with the 60 bp insertion exhibited a marked decrease under cold stress (Supplementary Fig. 10). Notably, two wild species, *S. cheesmaniae* and *S. galapagense*, did not contain the 60 bp insertion (Fig. 3e and Supplementary Fig. 10), possibly due to their warm growing environment in low-altitude areas<sup>39,40</sup>. The *S. habrochaites* introgression line LA3942, which contains a single introgression fragment where *ShWRKY34* replaces *SIWRKY34*, shows better cold tolerance than its recurrent parent *S. lycopersicum* LA4024 (Fig. 2). Thus, the full 60 bp InDel can be introduced into cultivated and cherry tomatoes through backcrossing and other breeding technologies to improve their cold tolerance.

Unlike changes in coding sequences, variations in *cis*-regulatory regions can alter gene expression in response to environmental cues and developmental processes without changing the protein they encode<sup>6</sup>. During evolution and domestication, certain variations in *cis*-regulatory regions can confer advantageous traits, such as enhanced yields, stress tolerance, or nutrient content. For instance, research on cotton (*Gossypium hirsutum*) has identified changes in *cis*-regulatory regions that have significantly altered gene expression during domestication, contributing to the development of desirable fiber traits<sup>41</sup>. One specific study identified a 26 bp InDel in the 5' UTR of *ZmGLK36*, which modulated its expression and thus the plant's resistance to maize rough dwarf virus (RBSDV), highlighting a key genetic adaptation for crop improvement in the face of disease challenges<sup>42</sup>. Previously, we found that a key W-box single nucleotide polymorphism (SNP) affects the self-transcriptional regulation and protein accumulation of *WRKY33* under cold stress in cultivated tomato, thus contributing to the cold sensitivity of cultivated tomatoes compared with wild tomatoes<sup>7</sup>. Here, we found that the *WRKY34* promoter in cold-tolerant wild tomato species contains a 60 bp insertion, which directly causes its chromatin to open and recruits the transcriptional suppressor GATA29 under cold stress, thereby diminishing *WRKY34* expression and enhancing cold tolerance. Conversely, the absence of this 60 bp segment in the *WRKY34* promoter of cultivated tomatoes leads to a reduced response to cold stress, contributing to the cold sensitivity observed in these domesticated tomato varieties. Both of our studies discovered SNP or InDel variations within the promoter regions of *WRKY* family transcription factors, leading to changes in gene expression levels and thereby affecting cold tolerance in tomatoes. Specifically, the mutation in *WRKY33* resulted in the loss of a promoter *cis*-element during tomato evolution, while the mutation in *WRKY34* involved the deletion of a fragment within the promoter. Although the functions and mechanisms of *WRKY33* and *WRKY34* in response to cold stress are completely different, variations in their promoters both result in decreased cold tolerance in cultivated tomato. Therefore, natural variation in the multiple genes related to cold tolerance may have occurred during the evolution of cultivated tomatoes and were preserved during domestication, resulting in an overall phenotype of cold sensitivity in cultivated tomatoes. Future research should explore how technologies such as gene editing can



restore these natural variations and improve cold tolerance in cultivated tomatoes without altering other traits.

The role of SWIB/MDM2 domain-containing proteins in chromatin remodeling is increasingly recognized<sup>43,44</sup>. Typically found in SWI/SNF chromatin remodeling complexes, the SWIB/MDM2 domain is known to regulate gene transcriptional activity by altering nucleosome

positioning<sup>43</sup>. This regulation can either be specific, affecting certain genes, or broadly influencing chromatin states across the genome. However, our study reveals a more direct and precise mechanism of SWIB/MDM2 domain proteins in gene regulation. We observed that under normal conditions, SWIBs were inactive, not opening chromatin or recruiting transcription factors, resulting in the low expression of

**Fig. 7 | SIWRKY34 interferes with the transcriptional activity of SICBF1, thus reducing cold tolerance in tomato seedlings.** **a** Yeast two-hybrid assay (Y2H) testing the interactions of SIWRKY34 with SICBF1 in yeast. SIW34-C, C terminal of SIWRKY34 protein. **b** Pull-down assays show that GST-tagged SIWRKY34 physically interacts with His-tagged SICBF1. The green triangle indicates the target band of SICBF1-His pulled down by SIW34-GST. The combination of GST and SICBF1-His was used as a negative control. **c** Bimolecular fluorescence complementation assay (BiFC) showing the interactions between SIWRKY34 and SICBF1 in tobacco. Bar: 50  $\mu$ m. **d** Co-IP showing the interactions between SIWRKY34 and SICBF1. The green triangle indicates the SIW34-HA target band immunoprecipitated with SICBF1-GFP. The combination of GFP and SIW34-HA was used as a negative control. **e** EMSA show that SIWRKY34 interferes with the binding of SICBF1 to DRE elements on downstream SICBF1 and SICOR47 promoters. Unlabeled wild type probes were used as cold probes. mu, mutated probes in which the DRE elements (5'-G/ACCGAC-3') were replaced with 5'-AAAAAA-3'. **f** Transient dual-luciferase (dual-

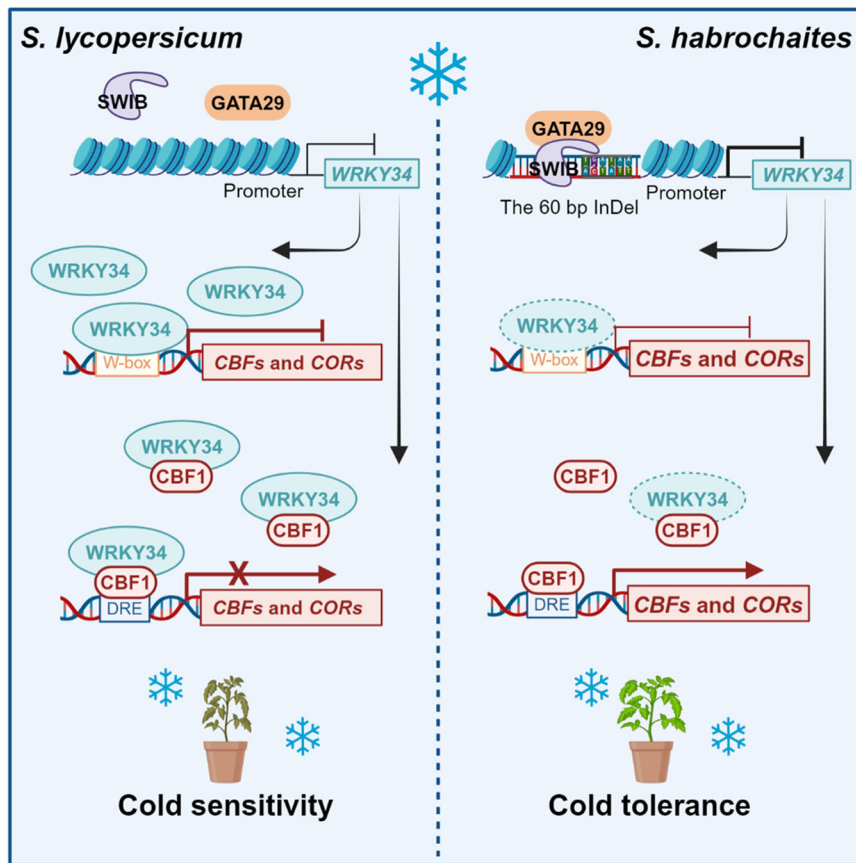
LUC) assay in tobacco leaves. Empty vector (EV) was included as control. Data are presented as means  $\pm$  SD of six biological replicates. **g** Phenotypes of *slurky34* mutant in LA4024 background (*slw34/LA4024*) and its *SICBF1*-silenced seedlings at 7 d after cold treatment. Eight plants of each genotype and treatment were tested with similar results. Bar: 10 cm. Relative electrolyte leakage (REL, **h**) and maximum photochemical efficiency of photosystem II ( $Fv/Fm$ , **i**) in *slurky34* mutant of LA4024 background (*slw34/LA4024*) and its *SICBF1*-silenced seedlings at 7 d after cold treatment. The false color code depicted at the bottom of the images ranges from 0 (black) to 1 (purple). Bar: 2 cm. Data are presented as means of four biological replicates  $\pm$  SD (**h**) or means of eight leaflets from independent plants (**i**). Experiments in (**a–e**) were repeated twice, and in (**f–i**), three times, all yielding similar results. Different letters above bars indicate significant differences ( $P < 0.05$ ) determined using one-way ANOVA with Duncan's multiple range test. TRV, non-silenced seedlings; TRV-*SICBF1*, *SICBF1*-silenced seedlings. Source data are provided as a Source Data file.

*WRKY34*. Nevertheless, under cold stress, SWIBs not only opened chromatin and recruited the transcription factor GATA29, but also directly bound to specific sites in the *WRKY34* promoter, thereby precisely inducing chromatin opening in specific regions and recruiting transcription factor GATA29 to bind to the GATA-box within the 60 bp region, further suppressing the expression of the *WRKY34* gene (Figs. 4–6). Even more excitingly, through protein structure prediction and experimental validation, we have demonstrated that three evolutionarily conserved amino acids in SWIBs are involved in DNA binding, with the key DNA binding site being TGATAA. We hypothesize that proteins of this family can recognize the TGATAA motif and then form a helical tripod structure to wrap around DNA, thereby further exerting their function. Therefore, the discovery about SWIBs extends beyond the traditional understanding of their role in altering nucleosome positions through chromatin remodeling and recruiting transcription factors. It reveals that these proteins can also directly bind to specific DNA sites through three evolutionarily conserved amino acids. This direct DNA binding capacity allows SWIB/MDM2 domain proteins to be key players in activating genes in chromatin-closed regions, enhancing the precision of specific gene expression control. Moreover, it demonstrates the ability of SWIB/MDM2 domain proteins to respond to specific environmental signals and regulate gene expression by acting directly on specific DNA elements, further emphasizing their versatility and importance in gene regulation. Beyond direct impacts on gene transcription, SWIB/MDM2 domain proteins are closely related to histone modifications and epigenetic regulation<sup>45</sup>. It has been reported that SWIB domain proteins might directly recognize and bind to specific histones or their modified forms, thereby altering nucleosome stability or regulating interactions with other chromatin-associated proteins. For example, under normal conditions, SWP73A repressed NLR (NOD-like receptor) gene *RPS2* through H3K9me2 modification, while this repression reduced or eliminated following pathogen infection, facilitating gene transcription and the activation of plant innate immunity<sup>46</sup>. However, the association of the SWIB/MDM2 domain proteins in our study with histone modifications warrants further investigation.

Research on *WRKY34* in *Arabidopsis* indicates that it is specifically expressed in pollen, negatively regulates the cold tolerance of mature pollen, and may be involved in the *CBF* signal cascade in mature pollen<sup>47</sup>. In our study, *WRKY34* also negatively impacts cold tolerance in tomato seedlings, primarily by interfering with the classic *CBF*-*COR* cold response pathway at both transcription and protein levels (Fig. 7 and Supplementary Fig. 18). *WRKY34* can directly bind to W-box elements in the promoters of *CBFs* and *CORs*, transcriptionally repressing their expression. Additionally, *WRKY34* interacts with *CBF1*, disrupting its transcriptional activation of itself and downstream *CORs*. Notably, while knocking out *WRKY34* enhanced cold tolerance, the *wrky34* mutants exhibited developmental defects, such as smaller fruits and

fewer seeds per fruit than WT (Supplementary Fig. 7). Moreover, the expression and protein accumulation of *WRKY34* were the highest in roots, followed by flowers and buds, with lower expression and protein accumulation in leaves and fruits (Supplementary Fig. 8). This emphasizes *WRKY34*'s necessity under normal conditions and suggests that complete functional loss isn't a desirable improvement approach for cold tolerance. Gene functions are multifaceted. Although knocking out or overexpressing genes can achieve desired traits, it may also disrupt other characteristics, like growth and development. Precisely regulating gene transcription through specific promoter control is vital in crop breeding, as this approach effectively balances the enhancement of desired traits with the overall health and growth of the plant. For example, a recent study utilized a gene-editing strategy targeting the *SIP1F4* binding motif in the *SICOMT2* promoter, effectively enhancing melatonin levels in tomato fruit during the ripening stage without impacting other developmental phases. In addition, this targeted approach achieved higher melatonin content and no growth defects compared to *pif4* knockout mutants, demonstrating the efficacy of precise genetic modulation in crop development<sup>48</sup>. Here, we identified a 60 bp insertion in the *WRKY34* promoter that diminished its expression under cold stress in *Solanum* species, thus boosting cold tolerance. Through multiple generations of backcrossing, we have also developed the *ShWRKY34* introgression tomato line LA3942, which exhibits cold tolerance without impacting other traits. Additionally, considering that variations in *cis*-regulatory regions typically exert subtler phenotypic impacts and circumvent the adverse effects of coding region mutations, we advocate employing gene-editing techniques to incorporate this 60 bp sequence into the *WRKY34* promoter of *Solanum* plants lacking this sequence.

Accessibility is generally positively correlated with expression, but examples of increased chromatin accessibility and decreased gene expression have been reported. For example, by investigating chromatin modifications and accessibility, a study suggests that although type A ARF exhibits an open chromatin configuration, it is regulated by a network of transcriptional repressors<sup>49</sup>. Therefore, the relationship between chromatin accessibility and gene expression is complex and influenced by multiple factors. Here, our results demonstrate one of these mechanisms and we thus propose a working model of *WRKY34*-mediated cold tolerance in wild and cultivated tomatoes (Fig. 8). Under cold stress, the presence of a 60 bp insertion in the *WRKY34* promoter of wild tomato *S. habrochaites* leads to the binding of chromatin remodeling factor SWIBs, thereby opening chromatin in the nearby region and recruiting transcriptional repressor GATA29 to bind to the GATA-box within the 60 bp, resulting in repression of *WRKY34* expression. *WRKY34* interferes with *CBF1*-induced expression of itself and *CORs* by interacting with *CBF1*. Furthermore, *WRKY34* directly binds to the promoters of downstream *CBFs* and *CORs* and represses their expression



**Fig. 8 | Proposed model of WRKY34-mediated cold tolerance in tomato.** The presence of the 60 bp InDel in the *WRKY34* promoter of wild tomato *S. habrochaites* leads to the binding of chromatin remodeling factor SWIBs, thereby opening chromatin in the nearby region, and recruiting more transcriptional repressor GATA29 to bind to the GATA-box within the 60 bp, resulting in repression of *WRKY34* expression. In addition, *WRKY34* interferes with the transcriptional regulation of CBF1 to itself and *CORs* by interacting with CBF1. On the other hand,

*WRKY34* directly binds downstream *CBFs* and *CORs* promoters and represses their expression under cold stress. However, the deletion of the 60 bp DNA fragment in the *WRKY34* promoter of cultivated tomatoes results in its inability to bind SWIBs under cold stress, preventing chromatin opening and recruitment of GATA29, and thus failing to suppress *WRKY34* expression and contributing to the cold sensitivity of these tomatoes. Figure 8 Created with BioRender.com released under a Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International license.

under cold stress. However, the deletion of the 60 bp DNA fragment in the *WRKY34* promoter of cultivated tomatoes results in its inability to bind SWIBs under cold stress, preventing chromatin opening and recruitment of GATA29, and thus failing to suppress *WRKY34* expression and contributing to the cold sensitivity of these tomatoes. Three additional points deserve to be mentioned. Firstly, ATAC-Seq data indicates that wild tomato LA1777 exhibits more chromatin opening under cold stress than cultivated tomato AC, suggesting potential functional differences in chromatin remodeling factors other than SWIBs between wild and cultivated tomatoes, which may impact cold tolerance. Secondly, in addition to the regulatory differences caused by non-coding regions, the differences in coding regions between wild and cultivated tomatoes and their potential impact on resistance traits warrant further investigation.

## Methods

### Plant materials and growth conditions

Wild tomato (*S. habrochaites* accession LA1777) and cultivated tomato (*S. lycopersicum* cv. Ailsa Craig, AC) were used for RNA-Seq and ATAC-Seq. The *S. habrochaites* introgression line LA3942, containing a single introgression fragment with *ShWRKY34* replacing *SlWRKY34*, along with its recurrent parent *S. lycopersicum* LA4024 and donor parent LA1777, was used for virus-induced gene silencing (VIGS) of *WRKY34* genes. LA4024 and LA3942 were selected for genetic transformation.

A total of 376 tomato accessions were collected from various sources, including Tomato Genetics Resource Center (TGRC), United State Department of Agriculture (USDA), University of Florida, and European Union Solanaceae Project (EU-SOL). These accessions include 63 wild tomato accessions (*3. S. cheesmaniae*, *2. S. galapagense*, *8. S. arcanum*, *3. S. chmielewskii*, *9. S. neorickii*, *3. S. huaylasense*, *4. S. corneliomulleri*, *4. S. peruvianum*, *12. S. chilense*, *10. S. habrochaites*, *4. S. pennellii* and *1. S. sitien*), *58. S. pimpinellifolium*, *74. S. lycopersicum* var. *cerasiforme*, and *181. S. lycopersicum* accessions (Supplementary Data 6).

Seeds were germinated on moistened filter paper at 28 °C in the dark and subsequently sown in 72-cell plastic flats filled with a mixture of peat and vermiculite (3:1, v:v). Upon reaching the two-leaf stage, seedlings were transplanted into plastic pots (10 cm × 10 cm in height × diameter, one seedling per pot) or 32-cell plastic flats containing the same medium. The plants were cultivated in a growth room under a 12 h photoperiod, with temperature of 25/20°C (day/night), and a photosynthetic photon flux density (PPFD) of 600  $\mu\text{mol m}^{-2} \text{s}^{-1}$ . The relative humidity was maintained at 70%, and plants were irrigated with 1/2 strength Hoagland's nutrient solution every 3 d.

### Cold stress treatment and cold tolerance evaluation

For cold stress treatment, tomato seedlings at the five-leaf stage or tobacco plants expressing reporter vectors were transferred to a cold artificial growth chamber set at 4 °C, maintaining the same conditions

as in the growth room. Each biological repeat contained eight seedlings from each tomato genotype, with three biological repeats per treatment. After 7 d of cold treatment, tomato seedlings were photographed. Then, relative electrolyte leakage (REL) was measured based on electrical conductivity and the maximum photochemical efficiency of photosystem II ( $F_v/F_m$ ) was measured using an Imaging-PAM Chlorophyll Fluorometer equipped with a computer-operated PAM-control unit (IMAG-MAXI; Heinz Walz, Effeltrich, Germany), as previously described<sup>50</sup>. Survival rate assays were conducted on 20-day-old seedlings (at the three-leaf stage) grown in the growth room, which were subjected to 4 °C treatment for the specified duration before being returned to normal conditions (25 °C) for 1 week of recovery. During this process, the survival rate (percentage of green plants recovered after cold treatment) was calculated. The survival rates of the seedlings were calculated with three independent replicates for each genotype.

#### RNA-Seq libraries preparation and data analysis

RNA-Seq was performed as previously described<sup>7</sup>. Briefly, tomato leaves of AC and LA1777 were collected under normal conditions or after 6 h of cold stress, respectively, and used for RNA extraction. RNA-Seq library preparation and paired-end sequencing were performed on an Illumina Novaseq™ 6000 sequence platform by LC Sciences (Hangzhou, China). Approximately 4 Gb of high-quality paired-end reads were generated from each library. Clean data (clean reads) were obtained by removing reads containing adapters, poly-N sequences and low-quality reads from raw data using Trimmomatic version 0.36. These clean reads were then aligned to the tomato genome (<https://solgenomics.net>, SL4.0) using the Hisat2 mapping tool. Genes with FPKM's  $P < 0.05$  and an absolute  $\log_2$  fold change  $\geq 1$  were considered as differentially expressed genes (DEGs).

#### Nuclei extraction and purification

Samples were prepared using sucrose sedimentation as previously reported<sup>51</sup> but with slight modifications. Briefly, young leaves of AC and LA1777 were collected under normal conditions or after 6 h of cold stress, respectively, and ground to fine powder in liquid nitrogen. For each sample, 0.2 g of frozen tissue powder was homogenized in pre-chilled 1 ml lysis buffer (15 mM Tris-HCl pH7.5, 20 mM NaCl, 80 mM KCl, 0.5 mM spermine, 5 mM 2-ME, 0.2% TritonX-100), and the nuclear fraction was purified as described<sup>52</sup>. The nuclei pellet was resuspended in 1 ml cold lysis buffer. For ATAC-Seq and ATAC-qPCR, a nuclei aliquot (25  $\mu$ l) was stained with DAPI (10  $\mu$ l of 1  $\mu$ g  $\text{ml}^{-1}$ ) and counted using a haemocytometer. Approximately 50,000 nuclei were used for each ATAC-Seq or ATAC-qPCR reaction.

#### ATAC-Seq libraries preparation and data analysis

ATAC-Seq was carried out as previously described<sup>53</sup> with some minor modifications. Briefly, nuclei were extracted and purified from samples, and the nuclei pellet was resuspended in the Tn5 transposase reaction mix. The transposition reaction was incubated at 37 °C for 30 min. Equimolar Adapter1 and Adapter2 were added after transposition, and PCR was then performed to amplify the library. After the PCR reaction, libraries were purified with AMPure beads (Beckman, A63881) and library quality was assessed with Qubit (Thermo Fisher, Q32854). The clustering of the index-coded samples was performed on a cBot Cluster Generation System using TruSeq PE Cluster Kit v3-cBot-HS (Illumina) according to the manufacturer's instructions. After cluster generation, the library preparations were sequenced on an Illumina platform at Novogene (Beijing, China) and 150 bp paired-end reads were generated.

Raw data was processed using fastp (version 0.20.0) to obtain clean reads, excluding adapters, poly-N, and low-quality sequences, while calculating Q20, Q30, and GC content. The reference genome and annotation were downloaded (<https://solgenomics.net>, SL4.0),

and its index was built with BWA (version 0.7.12) for alignment of clean reads. Reads from mitochondria and chloroplast DNA, improperly paired, and PCR duplicates were excluded. Peak calling was done with MACS2 (version 2.1.0). By default, peaks with q-value threshold of 0.05 were carried out for all datasets. Peaks of different groups were merged using 'bedtools merge'. We calculated the mean RPM of each group in the merge peak. Only peaks with an absolute  $\log_2$  fold change of  $\text{RPM} \geq 1$  and  $P < 0.05$  were considered as differential peaks. Genes associated with different peaks were identified using ChIPseeker. ChIPseeker was also used for gene and genomic region annotation<sup>54</sup>. GO enrichment analysis and KEGG pathway analysis were performed<sup>55</sup>. Differential peaks were identified with fold change of RPM more than 2. Genes associated with different peaks were identified using ChIPseeker. Peaks were visualized using the Integrative Genomics Viewer (version 2.12.2).

#### ATAC-qPCR

ATAC-qPCR was performed using the SYBR Green PCR Master Mix Kit (Takara, Shiga, Japan) on a Light Cycler 480 II detection system (Roche, Basel, Switzerland). Primers used for this analysis are shown in Supplementary Table 1. The relative accessibility and standard errors were determined using the  $2^{-\Delta\Delta\text{CT}}$  method<sup>56</sup>.

#### VIGS

Complementary DNA (cDNA) fragments of target genes were amplified using gene-specific primers containing EcoRI and BamHI restriction sites (Supplementary Table 2). Purified PCR products were cloned into the TRV2 vector. The plasmids were then transformed into *Agrobacterium tumefaciens* strain GV3101. Fully expanded cotyledons of tomato seedlings were infiltrated with a mixture of *A. tumefaciens* strain carrying the helper vector TRV1 mixed at 1:1 with the strain carrying either TRV2 (empty vector control, TRV) or TRV2-target gene vectors<sup>57</sup>. The infiltrated plants were maintained in the growth chambers, and the silencing efficiency of the targeted genes was determined by RT-qPCR (Supplementary Fig. 19).

#### Constructs for genetic transformation

For overexpression constructs, the full-length coding sequences (CDS) of *SlWRKY34*, *SlGATA29* and *SlSWIBb* were amplified from LA4024 cDNA and the CDS of *ShWRKY34* was amplified from LA1777 cDNA using specific primers (Supplementary Data 8). For generating transgenic overexpressing lines, the *SlWRKY34*, *ShWRKY34* and *SlGATA29* CDS were inserted into a pFGC1008-3HA binary plasmid vector behind the *CaMV 35S* promoter. The *SlSWIBb* CDS was inserted into a pAC402-GFP binary plasmid vector behind the *CaMV 35S* promoter. All vectors were transformed into *A. tumefaciens* strain GV3101 for plant transformation. The resulting *SlWRKY34* and *ShWRKY34* overexpression plasmids were introduced into cultivated tomato LA4024, while both LA4024 and LA3942 were used as transgenic recipient materials to transform *SlGATA29* and *SlSWIBb* overexpression plasmids. The transgenic overexpressing lines were further identified by RT-qPCR (Supplementary Fig. 20). The homozygous T2 transgenic lines were used in subsequent studies.

For the CRISPR/Cas9 constructs, single-guide (sgRNAs) or two-guide RNAs containing 20-bp targeting sequences were designed using the CRISPR-P web tool (<http://cbi.hzau.edu.cn/crispr/>) (Supplementary Data 8). The synthesized sequences were annealed and inserted into the BbsI site of the AtU6-sgRNA-AtUBQ-Cas9 vector as previously described<sup>58</sup>. The resulting plasmids were digested by HindIII and KpnI and then inserted into the pCAMBIA1301 binary vector digested by the same restriction enzymes. All resulting plasmids were transformed into *A. tumefaciens* strain GV3101 and infected into tomato cotyledons. First-generation transgenic plants were genotyped with specific primers surrounding the target sites (Supplementary Table 3). The homozygous F2 mutant lines without Cas9 were selected

and used for further study. As shown in Supplementary Fig. 21a, the *slwrky34-4* mutants harbored a 1 bp insertion and *slwrky34-5* a 2 bp deletion in the *SIWRKY34* coding region, leading to early translation termination. The protein bands of WRKY34 are barely observable in *wrky34* mutants (Supplementary Fig. 21b). The *slswibab* double mutants in LA4024 background harbored a 4 bp deletion in the *SISWIBa* coding region and a 1 bp deletion in the *SISWIBb* coding region, both leading to early translation termination (Supplementary Fig. 21c). The *slswibab* double mutants in LA3942 background harbored a 1 bp deletion in the *SISWIBa* coding region and a 4 bp deletion in the *SISWIBb* coding region, both leading to early translation termination (Supplementary Fig. 21c). The *slgata29* mutants in LA4024 background harbored a 2 bp deletion in the *SIGATA29* coding region and the *slgata29* mutants in LA3942 background harbored an 8 bp deletion in the *SIGATA29* coding region, both leading to early translation termination (Supplementary Fig. 21d).

### RNA isolation and RT-qPCR

Tomato leaves were collected under normal conditions or after indicated hours of cold stress. Other environmental conditions (e.g., humidity, lighting) of the growth chambers were kept consistent during sample collection. Total RNA was extracted from tomato leaves using an RNAprep Pure Plant Kit (Tiangen, DP419) following the manufacturer's instructions. DNase I-treated extracted RNA (2 µg) was reverse-transcribed using a ReverTra Ace qPCR RT Kit (Vazyme, R223). For RT-qPCR, quantitative PCR was performed using the SYBR Green PCR Master Mix Kit (Vazyme, Q711) on a Light Cycler 480 II detection system (Roche, Basel, Switzerland). Tomato housekeeping genes *Actin2* and *Ubiquitin3* were used as internal references. Relative gene expression was calculated as previously described<sup>56</sup>. Primers used for RT-qPCR are listed in Supplementary Data 9.

### DNA extraction, PCR and sequencing

Genomic DNAs of different tomato varieties were extracted using the TIANamp Genomic DNA Kit (Tiangen, DP304) and stored at -80°C. The nucleotide sequences of the 60 bp InDel in *WRKY34* promoters of different tomato varieties were amplified with the following general primers: F: 5'-TGATATGAAACATTCAAGTTGA-3'; R: 5'- TAGGTC GTGAAAATGAGGTACATA-3'. The amplified products were sequenced by Sanger sequencing using an ABI 3730xl instrument by Youkang Biotechnology (Hangzhou, China). The sequencing results are shown in Supplementary Data 6.

### Transient expression assays in tobacco leaves

Transient expression assays in tobacco leaves were performed as previously described in ref. 7. For promoter activity assays, promoters of *pSIW34*, *pShW34*, *pSIW34<sup>+60bp</sup>*, *pShW34<sup>mW-box</sup>*, *pShW34<sup>mGATA-box</sup>* and *pSIW34<sup>+30bp</sup>* were inserted into pGreenII 0800-LUC vectors as reporter genes. Renilla luciferase (*REN*) gene driven by *CaMV 35S* promoter in pGreenII 0800-LUC was used as an internal control to quantify transformation efficiency. Then, the above constructs were transformed into *A. tumefaciens* strain GV3101 and infiltrated into three-week-old tobacco leaves. After inoculation for 36 h, tobacco plants were treated at 4 °C for 6 h and proteins were extracted using Dual Luciferase Reporter Assay Kit (Vazyme, DL101).

For dual-luciferase (LUC) transcription activity assays, full-length CDSs of *SIGATA29*, *ShGATA29*, *SISWIBa/b*, *ShSWIB*, *SlCBF1* and *SIWRKY34* were inserted into pGreenII 0029 62-SK vectors as effectors. The empty vector SK was used as a control. Promoters of *pSIW34*, *pShW34*, *pSIW34<sup>+60bp</sup>*, *pShW34<sup>mW-box</sup>*, *pShW34<sup>mGATA-box</sup>*, *pSIW34<sup>+30bp</sup>*, *pSlCBF1* and *pSlCOR47* were inserted into pGreenII 0800-LUC vectors as reporter genes. Then, all the constructs were transformed into *A. tumefaciens* strain GV3101. The tobacco leaves were transfected with different combinations of vectors (*A. tumefaciens* strain carrying the

pGreenII 0800-LUC vector or pGreenII 0029 62-SK vector in a 1:10 ratio) for 36 h, then collected and lysed for the detection of dual luciferase activity (Vazyme, DL101) according to the manufacturer's recommendations.

The activities of firefly LUC and renilla luciferase REN were measured using the Glomax 96 microplate luminometer (Promega, Fitchburg, USA). The measured levels were normalized by calculating the LUC/REN ratio. Primers used for plasmid construction are listed in Supplementary Data 8.

### Y1H assay

Y1H assays were performed as previously described<sup>59</sup>. The 60 bp InDel was amplified and cloned into the pAbAi vector to generate pAbAi-60bp, while the 60 bp InDel containing mutant GATA-box or SWIB-mu4 was also amplified and cloned into the pAbAi vector to generate pAbAi-mut<sup>GATA-box</sup> or pAbAi-mut<sup>SWIB-mu4</sup>. The promoters of *SlCBF1/2/3* and *SlCOR47* were amplified and cloned into the pAbAi vector to generate pAbAi-SlCBF1, pAbAi-SlCBF2, pAbAi-SlCBF3 and pAbAi-SlCOR47. Full-length CDSs of *SIGATA29*, *ShGATA29*, *SISWIBa*, *SISWIBb*, *ShSWIB* and *SIWRKY34* were amplified and cloned into the pGADT7 vector as prey plasmids. The mutant *SISWIBa/b* CDSs (*SISWIBa<sup>R6A</sup>*, *SISWIBa<sup>L46A</sup>*, *SISWIBa<sup>K88A</sup>*, *SISWIBa<sup>allmut</sup>*, *SISWIBb<sup>R6A</sup>*, *SISWIBb<sup>L44A</sup>*, *SISWIBb<sup>K86A</sup>*, and *SISWIBb<sup>allmut</sup>*) were synthesized and cloned into the pGADT7 vector by Youkang Biotechnology (Hangzhou, China). The linearized pAbAi constructs were transformed into Y1HGold yeast strain as bait strains, and screened with different Aureobasidin A (AbA) concentrations to detect background *AbAr* expression of bait strains. Then, prey plasmids were transformed into bait strains, and the transformed yeast cells were selected on selective plates (SD-Leu) supplemented with 150 ng ml<sup>-1</sup> or 200 ng ml<sup>-1</sup> AbA. The empty pGADT7 was used as the negative control. Primers used for plasmid construction are listed in Supplementary Data 8.

### Recombinant proteins and electrophoretic mobility shift assays (EMSA)

Full-length CDSs of *SIGATA29*, *ShGATA29*, *SISWIBa*, *SISWIBb*, *ShSWIB*, *SlCBF1* and *SIWRKY34* were PCR amplified and cloned into the pET-28a vector (Supplementary Data 8). All recombinant vectors were transformed into *Escherichia coli* strain BL21 (DE3) and expressed at 37°C until OD<sub>600</sub> reached 0.6, and then induced by 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG, SIGMA, 092M4001V) at 16 °C for 14 h. The recombinant His-fusion proteins were purified according to the instructions provided with the Novagen pET purification system. To carry out EMASAs, oligonucleotide probes (Supplementary Table 4) were biotin-labelled using the Biotin 3'-End DNA Labeling Kit (Thermo Fisher Scientific, 89818) according to the manufacturer's protocol and annealed to double-stranded DNA. EMASAs were performed using the Light Shift Chemiluminescent EMSA kit (Thermo Fisher Scientific, 20148) according to the manufacturer's instructions<sup>60</sup>. Briefly, purified recombinant proteins were incubated with biotin-labelled probes at 28 °C for 30 min in 20 µl binding buffer (10× binding buffer, 50% glycerol, 25 ng µl<sup>-1</sup> poly-dL-dC, 1% NP-40). For competition assays, 10-, 20- or 100-fold non-labelled competitor DNA was added to the reaction. The reaction products were resolved on a 6% polyacrylamide gels in 0.5 × TBE at 100 V for 1-2 h on ice. Probe-protein complexes and free probes were transferred to a charged Hybond-N membrane and detected by western blotting with 1:5000 diluted anti-biotin antibodies (Abcam, ab53494).

### Microscale thermophoresis (MST) assay

The mutant *SISWIBa/b* CDSs (*SISWIBa<sup>R6A</sup>*, *SISWIBa<sup>L46A</sup>*, *SISWIBa<sup>K88A</sup>*, *SISWIBa<sup>allmut</sup>*, *SISWIBb<sup>R6A</sup>*, *SISWIBb<sup>L44A</sup>*, *SISWIBb<sup>K86A</sup>*, and *SISWIBb<sup>allmut</sup>*) were synthesized and cloned into the pET-28a vector by Youkang Biotechnology (Hangzhou, China). All recombinant His-proteins were

induced and purified as described above. The Cy5-labeled double-stranded DNA was synthesized and diluted by MST buffer containing 50 mM Tris-HCl (pH 8.0), 150 mM NaCl and 0.05% tween-20. 16 micro reaction tubes were set up, with tube 1 containing His-tag *SISWIBa/b* proteins and tubes 2 to 16 filled with MST buffer. A dilution process was initiated by transferring a sample from tube 1 to tube 2. A serial dilution was obtained by repeating 15 times and remove 10  $\mu$ l from tube number 16 after mixing. 10  $\mu$ l of Cy5-labeled DNA were mixed with 10  $\mu$ l of purified proteins and incubated at room temperature for 10 minutes, then loaded into silica capillaries (Polymicro Technologies, TSP010150). Binding reactions were measured using a Monolith NT.115 instrument (NanoTemper Technologies) at 25 °C, 40% MST power and 20% LED power. The Kd values were calculated using the mass action equation via the NanoTemper MO. Affinity Analysis software (NanoTemper Technologies).

#### ChIP-qPCR

ChIP experiments were performed using the EpiQuiK Plant ChIP kit (Epigentek, 50-109-6154) as described in the manufacturer's protocol<sup>7</sup>. Approximately 1.5 g of leaf tissue was harvested from transgenic plants and WT plants under normal conditions or after 6 h of cold stress, respectively. The harvested tissues were crosslinked in 1× PBS buffer containing 1% formaldehyde for 15 min under vacuum. Fixation was stopped by adding 1× PBS buffer containing 0.125 M glycine under vacuum for 5 min. After washing three times with cold sterilized water, the tissues were homogenized, dried and ground into powder in liquid nitrogen, followed by isolation and sonication of chromatin. Sonicated chromatin fragments were immunoprecipitated with either anti-HA antibody or anti-GFP antibody, while a goat anti-mouse IgG antibody was served as the negative control. The enriched DNA was amplified by qPCR using specific primers (Supplementary Table 5). Relative enrichment was calculated by comparing the percentage of anti-HA- or anti-GFP-immunoprecipitated DNA to the percentage of IgG-immunoprecipitated DNA.

#### Y2H assay

The CDSs of *SISWIBa*, *SISWIBb*, *ShSWIB* and the C-terminal fragment of *SIWRKY34* and *ShWRKY34* were amplified and cloned into the pGBKT7 vector. The CDSs of *SIGATA29*, *ShGATA29*, *SICBF1/2/3*, *ShCBF1/2/3*, *SICOR47*, *SICOR15a*, *SICOR27*, *ShCOR47*, *ShCOR15a* and *ShCOR27* were amplified and cloned into the pGADT7 vector. The resulting constructs were co-transformed in various combinations into *Saccharomyces cerevisiae* strain AH109 according to the manufacturer's instruction (Yeastmaker™ Yeast Transformation System 2). The transfected yeast cells were grown on SD-Leu-Trp plates at 28 °C for 3 d and then transferred to selective plates (SD-Leu-/Trp-/His-/Ade) at 28 °C for 4 d. Primers used for plasmid construction are listed in Supplementary Data 8.

#### Pull-down assay

Full-length CDSs of *SISWIBa*, *SISWIBb*, *ShSWIB* and *SICBF1* were amplified and cloned into the pET-28a vector. Full-length CDSs of *SIGATA29*, *ShGATA29* and *SIWRKY34* were amplified and cloned into the pGEX-4T-3 vector. The recombinant vectors were transformed into *E. coli* BL21 (DE3). The recombinant His-fusion proteins were purified according to the instructions provided with the Novagen pET purification system. To carry out pull-down assays, GST and GST-fusion proteins were extracted with extraction buffer and kept immobilized on Glutathione Sepharose 4B beads (Cytiva, 17075601)<sup>61</sup>. Glutathione beads containing the GST and GST-fusion proteins were incubated with equal amounts of different His-fusion proteins at 4 °C for 3 h and then were washed five times with PBS buffer (containing 0.1% tween-20). The proteins were detected by immunoblotting with anti-His antibody and anti-GST antibody. Primers used for plasmid construction are listed in Supplementary Data 8.

#### Bimolecular fluorescence complementation (BiFC)

For BiFC assay, full-length CDSs of *SISWIBa*, *SISWIBb*, *ShSWIB* and *SIWRKY34* were amplified and cloned into the binary vector p2YN. Full-length CDSs of *SIGATA29*, *ShGATA29* and *SICBF1* were amplified and cloned into the binary vector p2YC. They were then transformed into *A. tumefaciens* strain GV3101. For transient expression, different combinations of *A. tumefaciens* carrying different constructs at OD<sub>600</sub> = 0.8 were co-infiltrated into four-week-old *Nicotiana benthamiana* leaves. Nucleus-located H2B-mCherry was used as a nucleus marker. After 36 h of infiltration, the fluorescence signals of the infiltrated leaves were observed under a confocal laser scanning microscope (Zeiss LSM 780, Oberkochen, Germany) using preset settings of YFP (Ex: 488 nm, Em: 520-540 nm) and mCherry (Ex: 561 nm, Em: 610-630 nm). Primers used for plasmid construction are listed in Supplementary Data 8.

#### Co-immunoprecipitation (Co-IP)

Full-length CDSs of *SISWIBa*, *SISWIBb*, *ShSWIB* and *SICBF1* were amplified and cloned into GFP tag vector, while full-length CDSs of *SIGATA29*, *ShGATA29* and *SIWRKY34* were amplified and cloned into HA tag vector. They were then transformed into *A. tumefaciens* strain GV3101. For transient expression, different combinations of *A. tumefaciens* carrying different constructs at OD<sub>600</sub> = 0.8 were co-infiltrated into four-week-old *N. benthamiana* leaves. After 36 h of infiltration, the proteins co-expressed in the infiltrated leaves were extracted using Co-IP buffer and analyzed by immunoblotting with anti-HA antibody and anti-GFP antibody<sup>60</sup>. Extracts of equal total proteins were incubated with anti-GFP Magnetic Beads (Chromotek) for 3 h with gentle rotation at 4 °C. Beads were washed five times with the Co-IP buffer. The immunoprecipitated proteins were analyzed by immunoblotting with anti-HA antibody. Primers used for plasmid construction are listed in Supplementary Data 8.

#### Protein extraction and western blotting

Protein extraction and western blotting were performed as described<sup>60,62</sup>. Proteins were separated by SDS-PAGE using 10% (w:v) acrylamide gels and then transferred onto nitrocellulose membranes. WRKY34 protein was detected with anti-WRKY34 polyclonal antibody (No.230608037). Anti-WRKY34 polyclonal antibody was customized by the Laboratory Animal Center of Zhejiang University.

#### Statistical analysis

Data were subjected to statistical analysis of variance using the SPSS package (SPSS 19.0). Data were represented as the mean  $\pm$  SD. The difference between the two groups was assessed by two tailed Student's *t*-tests. Statistically significant differences among multiple groups were evaluated by one-way ANOVA followed by a Duncan's multiple range test. Details of each statistical test are indicated in the figure legends.

#### Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

#### Data availability

The RNA-Seq data generated in this study have been deposited in the NCBI Sequence Read Archive database under BioProject accession PRJNA825093. The ATAC-Seq data generated in this study have been deposited in the NCBI GEO data libraries under accession GSE254893. All cultivated tomato genes involved in this study can be found at the Sol genomics network with the following accession numbers: *SIWRKY34* (Solyc05g055750 [<https://solgenomics.net/locus/25311/view>]), *SICBF1* (Solyc03g026280 [<https://solgenomics.net/locus/4512/view>]), *SICBF2* (Solyc03g124110 [<https://solgenomics.net/locus/76752/view>]), *SICBF3* (Solyc03g026270 [<https://solgenomics.net/locus/17440/view>]), *SICOR47* (Solyc04g082200 [<https://solgenomics.net/locus/8351/view>]), *SICOR15a*

(Solyc06g083920 [<https://solgenomics.net/locus/28101/view>]), *SICOR27* (Solyc04g078880 [<https://solgenomics.net/locus/22587/view>]), *SIGATA29* (Solyc12g008830 [<https://solgenomics.net/locus/40868/view>]), *SISWIBa* (Solyc08g075400 [<https://solgenomics.net/locus/32384/view>]), *SISWIBb* (Solyc08g005590 [<https://solgenomics.net/locus/30733/view>]). All wild tomato genes involved in this study can be found at the NCBI or our RNA-Seq data with the following accession numbers or gene IDs: *ShWRKY34* (g49463), *ShCBF1* (ACB45087.1), *ShCBF2* (ACB45080.1), *ShCBF3* (ACB45078.1), *ShCOR47* (AHB20199.1), *ShCOR15a* (g53092), *ShCOR27* (g44172), *ShGATA29* (g19325), and *ShSWIB* (g55). Source data are provided with this paper.

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## Author contributions

M.G., J.Y. and J.Z. conceived and designed the experiments. M.G., F.Y., L.Z., L.W., Z.L. and Z.Q. performed the experiments. M.G. and J.Z. analyzed the data. V.F. provided the critical discussion. M.G. and J.Z. wrote the article. All authors reviewed and revised the article.

## Competing interests

The authors declare no competing interests.

## Additional information

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