HEARTBREAK Controls Post-translational Modification of INDEHISCENT to Regulate Fruit Morphology in *Capsella*

Graphical Abstract



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In Brief

Dong et al. reveal post-translational modification as a so-far-undisclosed driver of morphological diversity. They show that the SUMO protease HEARTBREAK is required to de-SUMOylate a key regulator of fruit-shape determination in *Capsella*, thereby initiating a transduction pathway leading to local auxin biosynthesis and anisotropic cell growth.

Highlights

- *HTB* encodes a SUMO protease required for fruit shape in *Capsella*
- Anisotropic cell growth is suppressed in the fruit valves of the *htb* mutant
- HTB stabilizes CrIND through de-SUMOylation to facilitate local auxin biosynthesis







Report

HEARTBREAK Controls Post-translational Modification of INDEHISCENT to Regulate Fruit Morphology in *Capsella*

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SUMMARY

Morphological variation is the basis of natural diversity and adaptation. For example, angiosperms (flowering plants) evolved during the Cretaceous period more than 100 mya and guickly colonized terrestrial habitats [1]. A major reason for their astonishing success was the formation of fruits, which exist in a myriad of different shapes and sizes [2]. Evolution of organ shape is fueled by variation in expression patterns of regulatory genes causing changes in anisotropic cell expansion and division patterns [3-5]. However, the molecular mechanisms that alter the polarity of growth to generate novel shapes are largely unknown. The heart-shaped fruits produced by members of the Capsella genus comprise an anatomical novelty, making it particularly well suited for studies on morphological diversification [6–8]. Here, we show that post-translational modification of regulatory proteins provides a critical step in organ-shape formation. Our data reveal that the SUMO protease, HEARTBREAK (HTB), from Capsella rubella controls the activity of the key regulator of fruit development, INDEHISCENT (CrIND in C. rubella), via de-SUMOylation. This post-translational modification initiates a transduction pathway required to ensure precisely localized auxin biosynthesis, thereby facilitating anisotropic cell expansion to ultimately form the heart-shaped Capsella fruit. Therefore, although variation in the expression of key regulatory genes is known to be a primary driver in morphological evolution, our work demonstrates how other processes—such as post-translational modification of one such regulatoraffects organ morphology.

RESULTS AND DISCUSSION

The *heartbreak* (*htb*) Mutant Has Valve Growth Defects and Reduced Cell-Growth Anisotropy

Organs in multicellular organisms have evolved into specific shapes that are critical for their function. Accordingly, little diversity is observed in organ morphology between individuals of the same species, with organs consistently and robustly developing into specific shapes [9]. By contrast, major variation in organ shape can exist between closely related species, as observed for fruits, leaves, insect wings, or the outer ears of mammals [7, 10–12]. Changes in the expression pattern of key regulatory genes is a major driver of such morphological diversity, ultimately giving rise to changes in cell division patterns and cell expansion [13, 14]. We have shown that sequence variation in regulatory domains of the fruit-tissue identity gene,

INDEHISCENT (IND) (*CrIND* in *Capsella*), is responsible for augmentation of its expression domain in the heart-shaped fruits from *Capsella rubella*. In turn, CrIND induces expression of auxin biosynthesis genes required for growth of the shoulders of the heart [8].

To identify genetic factors controlling this process and required for the formation of heart-shaped fruits in *Capsella*, we carried out a forward genetic screen of an ethyl methanesulfonate (EMS)-induced *Capsella rubella* (Cr22.5) mutant population. One mutant, *heartbreak* (*htb*), was isolated because of its strong defects in fruit development with compromised outgrowth of the fruit shoulders (Figures 1A, 1B, and 1D). Moreover, compared with wild type (WT), the *htb* mutant exhibits defects throughout both vegetative and reproductive development (Figures S1A–S1J). This demonstrates that the *HTB* gene regulates multiple developmental processes.



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Figure 1. The htb Mutant Produces Fruits with Defective Fruit Shape and Reduced Anisotropic Cell Growth

(A–C) Fruit morphology of WT (A), htb-1 (B), and rescue line of htb-1 transformed with pHTB:HTB:GFP (C) at stage 17.

(D) Shoulder index measurements of fruits from WT, htb-1, and $htb-1^{re}$ (pHTB:HTB:GFP htb-1) plants.

(E–G) Scanning electron microscopy (SEM) images of fruits from WT at developmental stages 12 (E), 13 (F), and 14 (G), showing fruit-shoulder growth after pollination.

(H–J) SEM images of fruits from htb-1 at stages 12 (H), 13 (I), and 14 (J), showing compromised development of the fruit shoulders.

(K–N) Time-lapse live imaging of developing fruits from stages 12 to 13 and 13 to 14 in WT (K and M) and *htb-1* mutant (L and N). Cells are outlined by RFP signal of the clonal sectors derived from heat-shock treatment of *pHS:CRE/BOB-lox* line. The heatmaps represent the anisotropy (K and L) and the overall cell area ratio (M and N).

Scale bars, (A–C) 5 mm; (E–N) 100 μm. Error bars in (D) represent SD of 30 individual fruits. **p < 0.01 (Student's t test). See also Figure S1.

In WT *Capsella*, the heart-shaped fruit develops from a discformed (ovate spheroid) gynoecium soon after pollination [6] (Figure 1E). From stage 13 onward, directional outgrowths of the apical parts of the valves found formation of the heart shape by stage 14 (Figures 1F and 1G; developmental stages defined in [8]). Comparative ontogenetic analysis revealed no defects between WT and *htb* during early gynoecium development (Figures 1E and 1H). In contrast to WT, however, the outgrowth of the *htb* valve apex is significantly suppressed from stage 13 (Figures 1F, 1G, 1I, and 1J).

During postfertilization development, anisotropic cell expansion drives fruit growth toward the final size and shape [7, 15]. To assess the cellular basis underlying the *htb* phenotype, we traced the cell growth dynamics by time-lapse imaging of developing fruits [16]. We chose three specific stages: stage 12 (immediately preceding the initiation of shoulder outgrowth); stage 13 (outgrowth begins); and stage 14 (shoulders are clearly formed; Figures 1E–1G). In stage-13 WT fruits, cells in the apical part of the valve grew anisotropically along the medio-lateral axis (Figures 1K and S1K). At stage 14, most of the cells in the apical part of the WT fruit had become highly anisotropic, growing toward the developing fruit shoulders, although cells in the basal part of the fruit remained largely isotropic from stages 12 to 14 (Figure 1K). In WT, the overall cell expansion rate was similar between apical and basal parts of the fruit (Figures 1M and S1L). In contrast to WT, cells in the valves of the *htb* mutant grew isotropically throughout all the stages studied here, leading to reduced growth rate in the shoulders (Figures 1L and S1K). Also, in comparison to WT, the *htb* mutant displayed a decreased overall cell expansion rate in the apical part of the fruit.





Figure 2. Molecular Cloning and Expression Analysis of HTB

(A) Cloning of the *htb-1* allele identified a G-to-A mutation in the acceptor site of the first intron of Carubv10008238, which disrupts the splicing of the first intron and results in a 7-bp deletion in the second exon, generating a premature stop codon in exon 2. The *htb-2^{ge}* allele was generated by CRISPR with a single-base-pair deletion in the exon 2, resulting in a frameshift giving rise to a 77-amino-acid (aa) protein. The guide RNAs and PAM sequences were indicated by red and blue characters, respectively.

(B–G) GUS staining of *pHTB:GUS* line showing the dynamic expression of *HTB* during fruit development. Uniform expression of *HTB* is detected in inflorescence tissue (B) and in the gynoecium at stage 11 (C) and 12 (D). A stronger *HTB* expression is detected in the developing fruit shoulders in stages 13 (E) and 14 (F). At stage 15, only residual *HTB* expression is observed in the fruit (G).

(H and I) Subcellular localization of HTB:GFP protein in the roots of pHTB:HTB:GFP line.

Scale bars in (B)-(I) represent 100 µm.

(J) Comparative analysis of SUMO conjugates in total protein extracts from leaf, inflorescence (inflo.), and stage-13 (S13) and stage-15 (S15) fruits between WT and *htb-1*. The α-tubulin was immunoblotted as a loading control.

See also Figure S2.

(Figures 1N and S1L). These data demonstrate that the *HTB* locus functions to promote anisotropic cell growth in the fruit valves.

The *HTB* Gene Encodes a Nuclear-Localized Protein Annotated as a SUMO Protease

The *htb* mutation segregates as a single-locus recessive trait (Figure S2B). By whole-genome sequencing and associative mapping, we identified two candidate mutations in the predicted genes, *Carubv10012951* and *Carubv10008238*. A synonymous mutation in the first exon of *Carubv10012951* precluded it for

further consideration. Instead, a potential causal mutation in the acceptor site of the first intron of a predicted gene, Carubv10008238, was investigated further. This gene encodes a putative small ubiquitin modifer (SUMO) protease, a member of the ULP2 subfamily of cysteine proteases, and is orthologous to the *Arabidopsis SPF1/ASP1* gene (Figure S2A) [17, 18]. The mutation disrupts the splicing of the first intron, which instead occurs after an alternative site 7 bp into the second exon, resulting in a frameshift and premature stop codon (Figures 2A and S2C). We will refer to this mutant allele as *htb-1*. Verification of the causality of this mutation on fruit shape was confirmed as follows: (1)

expression of Carubv10008238 driven by its native promoter fully complemented the *htb-1* mutant (Figures 1C and 1D); (2) a knockout line of Carubv10008238 using CRISPR-Cas9, leading to a single-base deletion in the second exon (*htb-2^{ge}*), phenocopied the *htb-1* fruit character alongside other developmental defects (Figures 2A and S2D–S2F); and (3) F₁ plants of *htb-1* crossed with the *htb-2^{ge}* mutant show the same phenotype as *htb-1* (Figure S2G). Together, these experiments show that the developmental defects observed in the *htb-1* mutant are caused by loss of the Carubv10008238 gene, which we henceforth refer to as *HEARTBREAK* (*HTB*), encoding a putative SUMO protease.

In agreement with the wide range of developmental defects of the *htb-1* mutant, we found a *pHTB:GUS* reporter line to be expressed throughout plant development, including vascular tissue of cotyledons and roots and in root tips of seedlings (Figures S2H and S2I). *pHTB:GUS* signal seemed uniformly distributed in the inflorescences and young gynoecia (Figures 2B–2D). Notably, in the developing fruit, stronger *HTB* promoter activity is detected in the shoulders from stage 13 to stage 14, when the heart shape starts to develop, although at stage 15, only residual *HTB* expression is detected (Figures 2E–2G). *HTB* expression therefore correlates spatially and temporally with fruit growth in agreement with its role in promoting anisotropic cell growth in the valves.

The SPF1/ASP1 protein is located in the nucleus of *Arabidopsis* cells [18, 19]. To test the subcellular localization of HTB, we used a *pHTB:HTB:GFP* reporter line, which fully complements the *htb-1* mutant (Figures 1C and 1D). Strong GFP signal was seen specifically within the nucleus but excluded from the nucleolus in root cells (Figures 2H and 2I). A similar nuclear localization pattern was observed using transient overexpression of an HTB-GFP fusion protein in WT leaf protoplasts (Figure S2J). These data suggest that HTB exerts its function on fruit-shape formation by affecting the activity of nuclear proteins.

SUMO Conjugate Levels Are Elevated in the *htb-1* Mutant

The SUMOylation of proteins is a dynamic process with reversibility in conjugation and deconjugation [20]. SUMO proteases falling into the class of ubiquitin-like proteases (ULPs) belong to the cysteine protease family and are able to mediate SUMO maturation as well as SUMO deconjugation from protein targets through their endopeptidase and isopeptidase activity, respectively [21]. In order to determine whether HTB affects SUMO-conjugation levels, we compared the SUMOylation profiles between WT and htb-1 by western blotting using specific anti-SUMO1 antibodies. Compared with WT, high-molecularweight SUMO conjugates constitutively accumulated in totalprotein extracts from the htb-1 mutant. This was particularly evident in inflorescence tissue and stage-13 fruits (Figure 2J), suggesting that the developmental defects observed in the htb-1 mutant (Figures S1A-S1J) is due to over-SUMOylation of proteins that are targets of the HTB SUMO protease.

HTB Controls Fruit Development by Regulating Auxin Biosynthesis

SUMO proteases have been reported to control SUMOylation levels of transcription factors, chromatin remodeling factors,



and/or transcriptional co-repressors [18, 22-24]. In order to understand the relationship between the transcriptional profile and HTB function in fruit development, we performed a comparative transcriptomic analysis of stage-13 fruits between WT and htb-1, when the developmental difference started to emerge (Figures 1F and 1I). The RNA-profiling analysis generated a total of 605 significant differentially expressed genes (DEGs) between WT and htb-1. Among them, 190 were upregulated and 415 were downregulated (Data S1A and S1B). Gene Ontology (GO) and pathway-enrichment analyses identified enrichment of DEGs in processes such as oxidation-reduction, protein phosphorylation, responses to light stimulus, and cell wall organization and modification (Figures S3A and S3C; Data S2A and S2B). Intriguingly, genes involved in hormone response were over-represented in the DEGs, especially among the downregulated genes (Figures S3B and S3D; Data S2A and S2B). Among the 26 downregulated DEGs associated with hormone response, 11 were associated with auxin response, pinpointing a possible role of HTB in regulating auxin dynamics during fruit-shape determination (Figures S3D and S3E; Data S2B).

We recently reported that the development of the heart-shaped Capsella fruit requires an auxin maximum in the fruit shoulders ensured by local expression of auxin biosynthesis genes, CrTAA1 and CrYUC9 [8]. Hence, we analyzed whether auxin dynamics was disrupted in the htb-1 fruits compared to WT. To visualize the auxin signaling pattern in the fruit valves, we used the pDR5v2:GUS reporter whose expression marks and precedes shoulder growth and introduced it into htb-1. In stage-14 WT fruit, a gradient of auxin signaling was observed in the valves with a maximum in the fruit shoulders (Figure 3A). In contrast, in the htb-1 mutant, the auxin maxima in the shoulders were lost, signifying a reduction of auxin response in the *htb-1* fruits (Figure 3B). We next asked whether the lack of auxin maxima in the htb-1 fruit shoulders was due to low auxin levels. Direct measurements of both the predominant natural auxin, indole-3-acetic acid (IAA), and its precursor, indole-3-pyruvate (IPA), showed a significant reduction in the shoulders of htb-1 compared to WT (Figures 3C and 3D). In correlation with reduced IPA and IAA levels, we found that expression of CrTAA1 and CrYUC9 was lower in the htb-1 fruit shoulders compared to WT (Figures 3E-3J). These data suggest that the decrease in auxin response observed in htb-1 fruits can be attributed to low levels of auxin biosynthesis, resulting from reduced CrTAA1 and CrYUC9 expression. Rescue of the htb-1 phenotype by exogenous application of IAA and valveshoulder-specific expression of a bacterial auxin biosynthesis gene (pCrIND:iaaM) provided further evidence that auxin biosynthesis is a downstream output of HTB activity required for fruitshape formation (Figures 3K and 3O).

HTB Controls CrIND Function by De-SUMOylation

In *Capsella*, shoulder-specific expression of *CrTAA1* and *CrYUC9* is regulated by the basic-helix-loop-helix (bHLH) transcription factor, CrIND [8]. In the *crind-1*^{ge} mutant, the fruit shoulders fail to fully expand due to depletion of auxin in the fruits compared to WT [8]. The *htb-1* mutant exhibits a similar phenotype as *crind-1*^{ge}, and lack of an obvious exacerbation of the single mutant phenotypes in the *htb-1 crind-1*^{ge} double mutant suggests that *HTB* and *CrIND* function in the same pathway (Figure 4A). To explore this possibility further, we crossed *htb-*





Figure 3. HTB Regulates Fruit Growth via Fine-Tuning Auxin Homeostasis

(A and B) Auxin signaling visualized by *pDR5v2:GUS* in stage-14 fruits of WT (A) and *htb-1* (B).

(C and D) Measurements of IPA (C) and IAA (D) in fruit shoulders of WT and htb-1 in stage-14 fruits.

(E–H) Expression of CrTAA1 and CrYUC9 shown by GUS staining of the pCrTAA1:GUS and pCrYUC9:GUS reporter lines at developmental stage 14 in WT (E and G) and htb-1 (F and H).

(I and J) Expression analysis of CrTAA1 (I) and CrYUC9 (J) in fruit shoulders of WT, htb-1, and htb-1 pCrIND:CrIND^{K124R}:GFP at stage 14.

(K-N) Fruit morphology of WT (K), IAA mock (L), or IAA (M) treatment on htb-1 and htb-1 pCrIND:iaaM (N) at stage 17.

(O) Shoulder index measurements of fruits from WT, htb-1 ± IAA treatment, and htb-1 pCrIND:iaaM plants.

(P–S) Fruit morphology of WT (P), *crful-1* (Q), *crful-1;crind-1^{ge}* (R), and *crful-1;htb-1* (S) at stage 17. Red dots indicate the location from where SEMs were taken in (T)–(W).

(T–W) SEM images of valve epidermal cells of WT (T), crful-1 (U), crful-1;crind-1^{ge} (V), and crful-1;htb-1 (W) at stage 17.

(X) Expression analysis of *CrIND* in stage-14 fruits of WT and *htb-1*. n.s. indicates no statistically significant difference from WT.

Scale bars in (A), (B), and (E)–(H), 150 μ m; (K)–(N) and (P)–(S), 5 mm; and (T)–(W), 50 μ M. Error bars in (C), (D), (I), (J), and (X) represent SD of three biological replicates and in (O) represent SD of 30 individual fruits. **p < 0.01 (Student's t test). See also Figure S3 and Data S1 and S2.

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1 with the *crful-1* mutant previously shown to be partially rescued by mutations in the *CrIND* gene [8] (Figures 3P–3R, 3T–3V, and S3F). The *htb-1* mutant also partially rescues the strong growth defect of *crful-1*, although to a lesser extent than *crind-1*^{ge} (Figures 3S, 3W, and S3F). It is therefore possible that the absence of HTB partially overturns the effect of ectopic *CrIND* expression previously reported to occur in *crful-1* [8].

Interestingly, expression of *CrIND* was unchanged in *htb-1* compared to WT (Figure 3X). This led us to test whether CrIND function is regulated post-translationally by HTB through SUMOy-lation. In plant cells, SUMOylation occurs through an isopeptide-bond formation between the di-glycine at the C-terminal of the SUMO peptide and the accessible lysyl ε -amino group within the targets [25]. Cumulative SUMO target datasets suggest a consensus ψ -K-X-D/E canonical SUMOylation motif (ψ , hydrophobic amino acid; X, any amino acid) [26]. Searching the CrIND sequence identified a consensus SUMO motif in amino acid



Figure 4. HTB Stabilizes CrIND by De-SU-MOylation

(A–C) Fruit morphology of htb-1; $crind-1^{ge}$ (A), htb-1pCrIND:CrIND:GFP (B), and htb-1 pCrIND: $CrIND^{K124R}:GFP$ (C) at stage 17.

(D) Shoulder index measurements of fruits from WT, *htb-1*, *htb-1 pCrIND:CrIND:GFP*, and *htb-1 pCrIND:CrIND*^{K124R}:GFP plants.

(E) SUMOylation status of CrIND protein using *pLhGR>>CrIND:FLAG* and *pLhGR>>CrIND^{K124R}: FLAG* lines. Immunoprecipitation experiments were conducted using anti-FLAG beads. Immunoblots were probed with anti-FLAG or anti-SUMO1 antibodies.

(F) Chromatin immunoprecipitation (ChIP) analysis of CrIND/CrIND^{K124R} associated with the *CrYUC9* and *CrTAA1* promoters.

(G) Model for the control of heart-shape fruit development by CrIND and HTB in *Capsella*. Precise production of auxin in the tips of fruit shoulders controlled by CrIND induces anisotropic cell growth in the valves in a direction toward the shoulder tips. CrIND protein is de-SUMOylated by HTB, whereas in the *htb-1* mutant, CrIND is SUMOylated and destabilized, thereby reducing its ability to activate expression of auxin biosynthesis gene (*CrTAA1* and *CrYUC9*).

Scale bars, (A–C) 5 mm. Error bars in (D) represent SD of 30 individual fruits and in (F) represent SD of three biological replicates. n.s. indicates no statistically significant difference from *htb*-1, *p < 0.05 and **p < 0.01 (Student's t test). See also Figure S4.

positions 123–126 (AKMD) with lysine in position 124 (K124) as a potential SUMOconjugation residue. To investigate the functional relevance of K124 in CrIND with HTB, we produced a mutant variant of CrIND, in which K124 is mutated to the related but unSUMOylatable amino acid, arginine (R), and compared the function of CrIND and CrIND^{K124R} in *htb-1* background. The K124R mutation did not change the protein function, as both

pCrIND:CrIND:GFP and *pCrIND:CrIND*^{K124R}:GFP fully rescued the *crind* mutant (Figures S4A–S4D). In the *htb-1* background, however, we observed a different behavior of these two proteins. Although *pCrIND:CrIND:GFP* failed to complement the *htb-1* mutant, *pCrIND:CrIND*^{K124R}:GFP effectively rescued the fruit defects of *htb-1*, developing fully heart-shaped fruits (Figures 4B–4D). This implies that post-translational modification of the K124 residue in CrIND is the primary cause of the defect in *htb-1* fruit-shoulder growth and suggests that HTB functions to de-SU-MOylate CrIND on this residue.

We then tested whether CrIND SUMOylation status depends on HTB. To this end, overexpression of FLAG-tagged versions of *CrIND* and *CrIND^{K124R}* in WT and the *htb-1* mutant was achieved using a two-component dexamethasone (DEX)-inducible system (Figure S4E). A pull-down experiment of FLAG-tagged CrIND/CrIND^{K124R} detected a high-molecular-weight version of SUMOylated CrIND only in the *htb-1* mutant background



(Figure 4E). Moreover, western blotting with FLAG antibody revealed low abundance of CrIND in *htb-1* compared to WT, whereas no reduction was observed with the CrIND^{K124R} version in *htb-1* (Figure 4E). These data demonstrate that HTB positively controls CrIND levels through de-SUMOylation, suggesting that SUMOylation on K124 of CrIND leads to its destabilization.

In agreement with reduced stability of CrIND in the htb-1 mutant, chromatin immunoprecipitation (ChIP) assays in htb-1 revealed that promoter regions of CrTAA1 and CrYUC9 were less enriched with CrIND-GFP compared to CrIND^{K124R}-GFP (Figure 4F). On the other hand, the binding affinities to CrTAA1 and CrYUC9 promoters were not significantly different between CrIND-GFP and CrINDK124R-GFP when ChIP assays were carried out in the crind-1^{ge} background (Figure S4F). Furthermore, CrTAA1 and CrYUC9 expression in the fruit shoulders is restored in htb-1 carrying the pCrIND:CrIND^{K124R}:GFP transgene (Figures 3I and 3J). Together, these biochemical and genetic data demonstrate that HTB acts directly on CrIND, leading to local expression of auxin biosynthesis genes. Although the effect of SUMOylation can vary widely between proteins, our results align with observations in both plants and animals that SUMOylation of transcription factors affects their stability and activity toward target genes [22-24, 27].

Our analyses did not identify any morphological differences between fruits from *crind* plants expressing *pCrIND:CrIND:GFP* and *pCrIND:CrIND*^{K124R}:*GFP* (Figures S4C and S4D). Although we cannot rule out subtle defects in other processes under CrIND control, this suggests that SUMOylation-resistant CrIND functions identically to wild-type CrIND, thus raising the question as to the purpose of the SUMOylation motif. We have previously shown that ectopic expression of *IND* in *Arabidopsis* can lead to profound developmental defects [28, 29]. As the expression domain of *CrIND* expanded to the tip of fruit valves in *Capsella*, we speculate that SUMOylation provides an additional regulatory layer to control CrIND activity to prevent deleterious effects.

HTB Protein Function Is Conserved between Capsella and Arabidopsis

In Arabidopsis, mutations in SPF1/ASP1, the ortholog of HTB, result in delayed flowering and abnormal floral and ovule development, although no fruit defect was described [17-19]. We therefore asked whether the HTB function in relation to fruit development is unique to Capsella. To this end, we transformed htb-1 with constructs of SPF1/ASP1 genomic sequences driven by the native HTB promoter (pHTB:AtSPF1/ASP1). The pHTB:AtSPF1/ASP1 construct fully complemented the htb-1 fruit defects in a similar manner to the pHTB:HTB construct, showing that HTB and SPF1/ASP proteins are functionally conserved (Figures S4G–S4J). This scenario is further supported by a population genetics analysis on the HTB locus in C. grandiflora, which is an out-crossing member of the Capsella genus [30]. In C. grandiflora, the protein sequence of HTB has been subjected to purifying selection (ω [dN/dS] < 1) [31], signifying no evidence for neo-functionalization of the HTB proteins (Figure S4K). Therefore, the difference in fruit shape between Capsella and Arabidopsis is not caused by functional diversification of the HTB protein itself. Rather, HTB is more likely to have been recruited specifically in the Capsella genus to modulate CrIND protein function, leading to precise auxin production



and specific anisotropic cell expansion to form the heart-shaped *Capsella* fruit (Figure 4G).

Concluding Remarks

In this study, we showed that the SUMO protease HTB targets the bHLH transcription factor CrIND for de-SUMOylation on lysine residue, K124. Removal of CrIND SUMOylation by HTB is required to stabilize CrIND and allow local activation of auxin biosynthesis genes in the fruit valves (Figure 4G). This, in turn, leads to stimulation of anisotropic cell expansion and formation of the heart-shaped *Capsella* fruit (Figure 4G). Although variation in the expression of key regulatory genes is known to be a primary driver in controlling morphological evolution, we demonstrated here how a post-translational modification of one such regulator, CrIND, affects organ morphology.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j. cub.2020.07.055.

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AUTHOR CONTRIBUTIONS

Y.D. and L.Ø. designed the research. Y.D. performed all the experiments with assistance from Ł.Ł., T.E., and N.S. M.M. and R.S.S. conducted the cell growth analysis. J.Š. and K.L. carried out IAA and IPA measurements; R.H. and T.S. did the population genetics analysis of *HTB* locus. A.K.S. and A.S. were involved in the SUMOylation test of CrIND. Y.D. and L.Ø. outlined and wrote the manuscript, and L.Ø. supervised the project. All authors participated in the discussion of the data and in the production of the final version of the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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REFERENCES

- Soltis, P.S., Folk, R.A., and Soltis, D.E. (2019). Darwin review: angiosperm phylogeny and evolutionary radiations. Proc. Royal Soc. B: Biol. Sci. 286, 20190099.
- Seymour, G.B., Østergaard, L., Chapman, N.H., Knapp, S., and Martin, C. (2013). Fruit development and ripening. Annu. Rev. Plant Biol. 64, 219–241.
- Carroll, S.B. (2000). Endless forms: the evolution of gene regulation and morphological diversity. Cell 101, 577–580.
- Carroll, S.B. (2005). Evolution at two levels: on genes and form. PLoS Biol. 3, e245.
- Wray, G.A. (2007). The evolutionary significance of cis-regulatory mutations. Nat. Rev. Genet. 8, 206–216.
- Eldridge, T., Łangowski, Ł., Stacey, N., Jantzen, F., Moubayidin, L., Sicard, A., Southam, P., Kennaway, R., Lenhard, M., Coen, E.S., and Østergaard, L. (2016). Fruit shape diversity in the Brassicaceae is generated by varying patterns of anisotropy. Development 143, 3394–3406.
- Łangowski, Ł., Stacey, N., and Østergaard, L. (2016). Diversification of fruit shape in the Brassicaceae family. Plant Reprod. 29, 149–163.
- Dong, Y., Jantzen, F., Stacey, N., Łangowski, Ł., Moubayidin, L., Šimura, J., Ljung, K., and Østergaard, L. (2019). Regulatory diversification of *INDEHISCENT* in the *Capsella* genus directs variation in fruit morphology. Curr. Biol. 29, 1038–1046.e4.
- 9. Moyroud, E., and Glover, B.J. (2017). The evolution of diverse floral morphologies. Curr. Biol. 27, R941–R951.
- Runions, A., Tsiantis, M., and Prusinkiewicz, P. (2017). A common developmental program can produce diverse leaf shapes. New Phytol. 216, 401–418.
- Salcedo, M.K., Hoffmann, J., Donoughe, S., and Mahadevan, L. (2019). Computational analysis of size, shape and structure of insect wings. Biol. Open 8, bio040774.
- Coleman, M.N., and Ross, C.F. (2004). Primate auditory diversity and its influence on hearing performance. Anat. Rec. A Discov. Mol. Cell. Evol. Biol. 281, 1123–1137.
- Huu, C.N., Kappel, C., Keller, B., Sicard, A., Takebayashi, Y., Breuninger, H., Nowak, M.D., Bäurle, I., Himmelbach, A., Burkart, M., et al. (2016). Presence versus absence of CYP734A50 underlies the style-length dimorphism in primroses. eLife 5, e17956.
- 14. Sicard, A., Kappel, C., Lee, Y.W., Woźniak, N.J., Marona, C., Stinchcombe, J.R., Wright, S.I., and Lenhard, M. (2016). Standing genetic

variation in a tissue-specific enhancer underlies selfing-syndrome evolution in *Capsella*. Proc. Natl. Acad. Sci. USA *113*, 13911–13916.

- Ripoll, J.J., Zhu, M., Brocke, S., Hon, C.T., Yanofsky, M.F., Boudaoud, A., and Roeder, A.H.K. (2019). Growth dynamics of the *Arabidopsis* fruit is mediated by cell expansion. Proc. Natl. Acad. Sci. USA *116*, 25333– 25342.
- Barbier de Reuille, P., Routier-Kierzkowska, A.-L., Kierzkowski, D., Bassel, G.W., Schüpbach, T., Tauriello, G., Bajpai, N., Strauss, S., Weber, A., Kiss, A., et al. (2015). MorphoGraphX: a platform for quantifying morphogenesis in 4D. eLife 4, 05864.
- Kong, X., Luo, X., Qu, G.P., Liu, P., and Jin, J.B. (2017). Arabidopsis SUMO protease ASP1 positively regulates flowering time partially through regulating FLC stability. J. Integr. Plant Biol. 59, 15–29.
- Castro, P.H., Santos, M.Â., Freitas, S., Cana-Quijada, P., Lourenço, T., Rodrigues, M.A.A., Fonseca, F., Ruiz-Albert, J., Azevedo, J.E., Tavares, R.M., et al. (2018). Arabidopsis thaliana SPF1 and SPF2 are nuclearlocated ULP2-like SUMO proteases that act downstream of SIZ1 in plant development. J. Exp. Bot. 69, 4633–4649.
- Liu, L., Jiang, Y., Zhang, X., Wang, X., Wang, Y., Han, Y., Coupland, G., Jin, J.B., Searle, I., Fu, Y.-F., and Chen, F. (2017). Two SUMO proteases SUMO PROTEASE RELATED TO FERTILITY1 and 2 are required for fertility in *Arabidopsis*. Plant Physiol. *175*, 1703–1719.
- Mukhopadhyay, D., and Dasso, M. (2007). Modification in reverse: the SUMO proteases. Trends Biochem. Sci. 32, 286–295.
- Yates, G., Srivastava, A.K., and Sadanandom, A. (2016). SUMO proteases: uncovering the roles of deSUMOylation in plants. J. Exp. Bot. 67, 2541–2548.
- 22. Gill, G. (2005). Something about SUMO inhibits transcription. Curr. Opin. Genet. Dev. 15, 536–541.
- Orosa-Puente, B., Leftley, N., von Wangenheim, D., Banda, J., Srivastava, A.K., Hill, K., Truskina, J., Bhosale, R., Morris, E., Srivastava, M., et al. (2018). Root branching toward water involves posttranslational modification of transcription factor ARF7. Science 362, 1407–1410.
- 24. Srivastava, M., Srivastava, A.K., Orosa-Puente, B., Campanaro, A., Zhang, C., and Sadanandom, A. (2020). SUMO conjugation to BZR1 enables Brassinosteroid signaling to integrate environmental cues to shape plant growth. Curr. Biol. 30, 1410–1423.e3.
- Miura, K., and Hasegawa, P.M. (2010). Sumoylation and other ubiquitinlike post-translational modifications in plants. Trends Cell Biol. 20, 223–232.
- Rytz, T.C., Miller, M.J., McLoughlin, F., Augustine, R.C., Marshall, R.S., Juan, Y.T., Charng, Y.Y., Scalf, M., Smith, L.M., and Vierstra, R.D. (2018). SUMOylome profiling reveals a diverse array of nuclear targets modified by the SUMO ligase SIZ1 during heat stress. Plant Cell *30*, 1077–1099.
- Rosonina, E., Akhter, A., Dou, Y., Babu, J., and Sri Theivakadadcham, V.S. (2017). Regulation of transcription factors by sumoylation. Transcription 8, 220–231.
- Sorefan, K., Girin, T., Liljegren, S.J., Ljung, K., Robles, P., Galván-Ampudia, C.S., Offringa, R., Friml, J., Yanofsky, M.F., and Østergaard, L. (2009). A regulated auxin minimum is required for seed dispersal in *Arabidopsis*. Nature 459, 583–586.
- Moubayidin, L., and Østergaard, L. (2014). Dynamic control of auxin distribution imposes a bilateral-to-radial symmetry switch during gynoecium development. Curr. Biol. 24, 2743–2748.
- 30. Guo, Y.L., Bechsgaard, J.S., Slotte, T., Neuffer, B., Lascoux, M., Weigel, D., and Schierup, M.H. (2009). Recent speciation of *Capsella rubella* from *Capsella grandiflora*, associated with loss of self-incompatibility and an extreme bottleneck. Proc. Natl. Acad. Sci. USA *106*, 5246–5251.
- McDonald, J.H., and Kreitman, M. (1991). Adaptive protein evolution at the Adh locus in Drosophila. Nature 351, 652–654.
- Liu, H., Ding, Y., Zhou, Y., Jin, W., Xie, K., and Chen, L.L. (2017). CRISPR-P 2.0: an improved CRISPR-Cas9 tool for genome editing in plants. Mol. Plant 10, 530–532.





- Jeanmougin, F., Thompson, J.D., Gouy, M., Higgins, D.G., and Gibson, T.J. (1998). Multiple sequence alignment with Clustal X. Trends Biochem. Sci. 23, 403–405.
- Tamura, K., Peterson, D., Peterson, N., Stecher, G., Nei, M., and Kumar, S. (2011). MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol. Biol. Evol. 28, 2731–2739.
- 35. Slotte, T., Hazzouri, K.M., Ågren, J.A., Koenig, D., Maumus, F., Guo, Y.L., Steige, K., Platts, A.E., Escobar, J.S., Newman, L.K., et al. (2013). The *Capsella rubella* genome and the genomic consequences of rapid mating system evolution. Nat. Genet. 45, 831–835.
- Novák, O., Hényková, E., Sairanen, I., Kowalczyk, M., Pospíšil, T., and Ljung, K. (2012). Tissue-specific profiling of the Arabidopsis thaliana auxin metabolome. Plant J. 72, 523–536.
- Bray, N.L., Pimentel, H., Melsted, P., and Pachter, L. (2016). Near-optimal probabilistic RNA-seq guantification. Nat. Biotechnol. 34, 525–527.

- Love, M.I., Huber, W., and Anders, S. (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 15, 550.
- Huang, W., Sherman, B.T., and Lempicki, R.A. (2009). Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. Nat. Protoc. 4, 44–57.
- Yoo, S.D., Cho, Y.H., and Sheen, J. (2007). *Arabidopsis* mesophyll protoplasts: a versatile cell system for transient gene expression analysis. Nat. Protoc. 2, 1565–1572.
- Steige, K.A., Laenen, B., Reimegård, J., Scofield, D.G., and Slotte, T. (2017). Genomic analysis reveals major determinants of cis-regulatory variation in *Capsella grandiflora*. Proc. Natl. Acad. Sci. USA *114*, 1087– 1092.
- Stoletzki, N., and Eyre-Walker, A. (2011). Estimation of the neutrality index. Mol. Biol. Evol. 28, 63–70.



STAR***METHODS**

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---|-------------------------|------------------|
| Antibodies | | |
| Anti-GFP monoclonal antibody | Roche | RRID: AB_390913 |
| Mouse monoclonal [M2] anti-FLAG-HRP antibody | Abcam | RRID: AB_869428 |
| Rabbit polyclonal anti-SUMO1 antibody | Abcam | RRID: AB 2198088 |
| Rabbit anti-mouse IgG-HRP secondary antibody | Abcam | RRID: AB_955440 |
| Goat anti-rabbit IgG-HRP secondary antibody | Abcam | RRID: AB_955447 |
| Mouse monoclonal Anti- α -tubulin antibody | Sigma | RRID: AB_477579 |
| Bacterial Strains | 0.9.1.4 | |
| DH5-alpha competent <i>E. coli</i> | NEB | C29871 |
| Agrobacterium tumefaciens strain LBA4404 | N/A | N/A |
| Biological Samples | | |
| Capsella rubella (WT, 22.5) | [6] | N/A |
| pHS:CRE/BOB-lox | [6] | N/A |
| pDR5v2:GUS | [8] | N/A |
| pCrTAA1:GUS | [8] | N/A |
| pCrYUC9:GUS | [8] | N/A |
| pCrIND:iaaM | [8] | N/A |
| crind-1 ^{ge} pCrIND:CrIND:GFP | [8] | N/A |
| htb-1 | This paper | N/A |
| htb-2 ^{ge} | This paper | N/A |
| crind-1 ^{ge} | [8] | N/A |
| crful-1 | [6] | N/A |
| crful-1; crind-1 ^{ge} | [8] | N/A |
| htb-1; crind-1 ^{ge} | This paper | N/A |
| htb-1; crful-1 | This paper | N/A |
| Chemicals Peptides, and Recombinant Proteins | | |
| Phusion High-Fidelity DNA polymerase | NEB | M0530L |
| Dnasel | QIAGEN | 79254 |
| In-Fusion Cloning Recombinase | Clontech | 638909 |
| Protease Inhibitor Cocktail | Roche | 11836170001 |
| PMSF | Roche | 10837091001 |
| Dexamethasone | Sigma-Aldrich | D4902 |
| Indole-3-acetic acid (IAA) | Sigma-Aldrich | 15148 |
| Ethyl methanesulphonate | Sigma-Aldrich | M0880 |
| N-Ethylmaleimide | Sigma-Aldrich | 04259 |
| DMSO | Sigma-Aldrich | D8418 |
| Gibberellin | Sigma-Aldrich | G7645 |
| Hygromycin | Roche | 10843555001 |
| DL-phosphinothricin | Duchefa | P0519.0250 |
| Formaldehyde | Sigma-Aldrich | F8775 |
| Cellulase R10 | Yakult | 190517 |
| | | |
| Macerozyme R IU | Yakult | 131126 |
| Macerozyme R10 Tween-20 | Yakult Sigma-Aldrich | 131126 P9406 |
| Tween-20 | Sigma-Aldrich | P9406 |
| - | | |

(Continued on next page)

CellPress

Current Biology Report

| Continued | | |
|--|---------------|---------------------------------------|
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
| Critical Commercial Assays | | |
| QIAprep Spin MiniPrep Kit | QIAGEN | 27104 |
| DNeasy Plant Mini Kit | QIAGEN | 69104 |
| QIAquick PCR Purification Kit | QIAGEN | 28104 |
| RNeasy Plant Mini Kit | QIAGEN | 74904 |
| Pierce Protein G Magnetic Beads | ThermoFisher | 19958500 |
| Pierce ECL Western Blotting Substrate | ThermoFisher | 32209 |
| Anti-FLAG [M2] Magnetic Beads | Sigma-Aldrich | M8823 |
| SuperScript IV First-Strand Synthesis System | ThermoFisher | 18091050 |
| SYBR Green JumpStart Taq ReadyMix | Sigma-Aldrich | S4438 |
| Recombinant DNA | | |
| pHTB:GUS | This Paper | N/A |
| pHTB:HTB | This Paper | N/A |
| pHTB:HTB:GFP | This Paper | N/A |
| pHTB:AtSPF1/ASP1 | This Paper | N/A |
| pCrIND:CrIND ^{K124R} :GFP | This Paper | N/A |
| pLhGR>>CrIND:FLAG | This Paper | N/A |
| pLhGR>>CrIND ^{K124R} :FLAG | This Paper | N/A |
| Software and Algorithms | | |
| MorphoGraphX | [16] | https://www.mpipz.mpg.de/MorphoGraphX |
| CRISPR-2.0 | [32] | http://crispr.hzau.edu.cn/CRISPR2/ |
| CLUSTAL-X | [33] | http://www.clustal.org/clustal2/ |
| MEGA5 | [34] | https://www.megasoftware.net/ |
| Other | | |
| PVDF membrane | GE Healthcare | 10600021 |
| Miracloth | Merck | 475855 |
| X-ray film | Kodak | 4741019289 |

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Lars Østergaard (lars.ostergaard@jic.ac.uk).

Materials Availability

• Plasmids and germplasm generated in this study is available upon request.

Data and Code Availability

• Data from genome sequencing and RNA-Seq have been deposited with the European Bioinformatics Institute (EBI): BioProject ID: PRJEB39302

Title: ena-STUDY-John Innes Centre-08-07-2020-15:50:15:918-1289 Release date: 2020-09-08, or until publication

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Plant materials, EMS-induced mutagenesis and growth condition

All *Capsella rubella* materials used in the study were in *Cr22.5* ecotype background. The *pHS:CRE/BOB-lox* line was described in [6], the *pDR5v2:GUS*, *pCrYUC9:GUS* and *pCrTAA1:GUS* reporter lines were previously described [8]. All these reporters were introgressed into *htb-1* mutant by crossing.

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For mutant screening, wild-type (WT) *Cr22.5* seeds were incubated with ethyl methanesulphonate (EMS, Sigma) at a concentration of 0.25% by volume in 0.02% Tween-20 (Sigma) rotating for 16 hours followed by 12 washes in 0.02% Tween-20 - in water. The seeds were germinated on soil in long-day (16 hr light/8 hr dark) conditions at 22°C and harvested to generate the M2 population. The *htb-1* mutant was discovered in the M2 segregation population. The *htb-1* mutant was backcrossed to WT three times to wash the genetic background and used for further studies.

The seeds were germinated on MS medium 1% sucrose and 0.8% agar containing 10 µM Gibberellin (Sigma) at 22°C. The 10-dayold seedlings were then transplanted into soil in a controlled environment room at 22°C, 16 hr light/8 hr dark conditions.

METHOD DETAILS

Plasmids construction and plant transformation

For the construction of the *pHTB:GUS* reporter plasmid, ~1.6kb promoter of Carubv10008238 was isolated and inserted upstream of *GUS* gene of pCambia1301 vectors. For the construction of *pHTB:HTB:GFP* plasmid, the genomic sequence from the Carubv10008238 locus (~7.9kb) was isolated and inserted into the pCambia1302 vectors. For *pHTB:AtSPF1/ASP1* and *pHTB:HTB* plasmid, the full length of genomic DNA of *AtSPF1/ASP1* (At1g09730,~6.2kb) or *HTB* (~6.5kb) was inserted downstream of the native *HTB* promoter in pCambia1302 vectors. The *pCrIND:CrIND^{K124R}:GFP* plasmid was domesticated from the *pCrIND:CrIND:GFP* plasmid described in [8]. For construction of *pLhGR>>CrIND:FLAG* plasmids, the *CrIND* and *CrIND^{K124R}* coding sequence is fused with 3X FLAG and inserted downstream of GR-inducible promoter to generate the *pGAL6:CrIND:FLAG* and *pGAL6:CrIND^{K124R}*. *FLAG* plasmids. The resultant plasmids were recombined with *p35S:GVG:GR* plasmid and phosphinothricin selection marker using golden-gate cloning methods to produce the binary vectors. For construction of the CRISPR/Cas9 genome editing plasmids, the DNA sequences encoding gRNAs adjacent to the PAM sequences (NGG) were designed using the CRISPR-P 2.0 software [32] that target two specific sites in the first exons of *Carubv10008238*. The gRNAs (Table S1) were synthesized as oligonucleotides with golden-gate cloning adapters and then were insert downstream of U6 promoters. The resulting gRNAs plasmid were then recombined with *pRPS5a:Cas92:E9t* and hygromycin selection marker using golden-gate cloning methods to produce the binary vectors are using golden-gate cloning methods to produce the binary vectors. All vectors were verified by sequencing and introduced into *Agrobacterium tumefaciens* strain LBA4404 by electroporation.

Transformation of *Capsella* followed the floral dipping method previously described [8]. The transformants were screened on MS plants with 1% sucrose and 0.8% agar containing 40mg/L hygromycin (Roche) or 25mg/L DL-phosphinothricin (Duchefa). For each construct, at least 10 independent transformants were obtained for further analysis.

Genome sequencing and association mapping

Leaf materials were collected from the BC_3F_2 segregation population of *htb-1* and WT crossing. Samples were pooled as WT and mutant (Mu) based on the fruit phenotypes, with each pool containing ~90 individuals. Nuclear DNA was extracted and fragmented and the sequencing libraries were prepared according to the manufacturer's instructions (Illumina). Sequencing reactions were processed on Illumina NextSeq500 platform generating paired-end reads with 100 X in depth. The SNPs were extracted by aligning the sequencing results with the v.1.0 reference genome of *Capsella rubella* [35]. For association mapping, we filter the SNPs between the WT and Mu samples by three criteria: (1) only consider the G to A and C to T SNPs as these are the mutations induced by EMS; (2) the mutation is heterozygous in the WT pool and homozygous in the Mu pool; (3) the mutated SNP frequency in WT pool is 33.3% and 100% in the Mu pool. From such screen, we identified two candidates, *Carubv10012951* and *Carubv1008238*. The G to A mutation happened in the first exon of *Carubv10012951* generates a synonymous mutation that preclude it for further consideration.

Phenotyping and Scanning Electron Microscopy (SEM)

The shoulder index value was calculated with the anti-trigonometric function θ = Arctan((L1-L2)/W) using the parameters described previously [8]. For whole-mount fruit photos, stage 17 fruits of each genotype were collected and photographed using Nikon D610 camera with a 105mm prime lens. For Scanning Electron Microscopy (SEM), the inflorescences of each genotype were fixed in FAA and dissected. The samples were critically-point dried in CO₂ and spotter-coated with gold. The samples were subsequently examined using a Zeiss Supra 55VP field Scanning Electron Microscope with an acceleration voltage of 3.0 kV.

Live imaging and cell growth analysis

For live imaging, *pHS:CRE/BOB-lox* lines were grown on soil in a glass house under long-day conditions until bolting (22°C, 16 hr light/8 hr dark). Then, the inflorescences were dipped into water bath at 38°C for 20 min and plants were grown for next 7 days. The fruits at stage 12 were dissected and transferred onto Petri dishes containing 1/2 MS medium including vitamins (Duchefa) supplemented with 1% sucrose. Half of the fruit epidermis was imaged with RFP signal at 48-h intervals using a Leica SP5 upright laser confocal microscope with a water immersion objective (x25/0.95). Excitation wavelengths and emission windows were 514 nm and 529-545 nm. Confocal stacks were acquired at 1024x1024 resolution, with less than 0.5-μm distance in Z-dimension. Between imaging, samples were kept in a growth chamber under long-day condition (22°C, 16 hr light/8 hr dark). The acquired images were stitched and analyzed using MorphoGraphX [16]. In order to calculate the cell area ratio and growth anisotropy, cells showing fluorescence were segmented and cell relations were indicated manually between successive time points. If cells divided in the subsequent time points, the daughter cells were merged. Heat-maps between two time-points are shown on the later time-point





(e.g., heatmap for fruit stage 12-13 is shown on the fruit stage 13). Representative growth tracking series were collected from a single growth tracking experiment and 3 time-lapse series were performed for wild-type and *htb-1*.

Auxin treatment and auxin metabolite quantification

To quantify auxin metabolite levels in the fruit, shoulder tissues of stage 14 fruit of WT and *htb-1* fruits were dissected under a light microscope and immediately fixed in liquid nitrogen. Extraction, purification and the LC-MS/MS analysis of endogenous IAA and specific IAA metabolites was carried out according to the method described previously [36].

RNA extraction, comparative transcriptomic sequencing and expression analysis

Either the whole fruit of stage 13 or the fruit shoulder samples from stage-14 fruits of WT and *htb-1*, respectively, were immediately fixed in liquid nitrogen. Total RNA was isolated from the samples using the RNeasy Plant Mini Kit (QIAGEN). 500ng of total RNA was reverse transcribed into cDNA in a 10 μ L reaction with the SuperScript IV First-Strand Synthesis System (ThermoFisher) according to the manufacturer's instructions.

For RNA-sequencing, poly (A) mRNA was purified from total RNA prepared from stage-13 fruits and fragmentated. Double-strand cDNA was synthesized, followed by sequencing adaptor ligation, electrophoresis purification and PCR amplification to generate the libraries using mRNA-Seq 8 sample prep kit (Illumina) according to the manufacturer's instructions. The libraries were paired-end sequenced on the Illumina NextSeq sequencer. The clean reads generated by trimming the adapters were mapped and annotated against the *Capsella rubella* v.1.0 genomic sequence using the Kallisto version 0.44.0 [37]. Read counts were generated using Kallisto version 0.44.0 [37]. Differentially expressed genes were identified as those with a fold change \geq 2 and a *p* value < 0.05 using DESeq2 software in R environment [38] (Data S1). The enrichment of the DEGs in the biological pathways were analyzed with DAVID Bioinformatics Resources 6.8 [39] (Data S1). Two biological replicates of RNA-seq for each sample were conducted.

For real-time qPCR, gene specific primers were designed (Table S1), and verified by PCR and sequencing. The efficiency of the primers (95% to 105%) was determined by creating a standard curve. The SYBR Green JumpStart Taq Ready Mix (Sigma) was used to perform real-time qPCR with ROX as a reference dye on a BioRad CFX96 Q-PCR System (BioRad). The CT value of each gene was determined by normalizing the fluorescence threshold. The relative expression level of the target gene was determined using the ratio $= 2^{-\Delta CT}$ method, and *CrUBQ10* was used as an internal control.

For the subcellular localization of the proteins in protoplast. Protoplast preparation and transformation was followed the protocol described in *Arabidopsis* [40] with minor modifications. Briefly, protoplasts were prepared from fully expanded leaves of 3-week old seedlings under short-day growth condition (22°C, 10 hr light/14 hr dark) using enzyme buffer [20mM MES (pH 5.7); 1.5% (wt/vol) cellulase R10 (Yakult); 0.4% (wt/vol) macerozyme R10 (Yakult); 0.4M mannitol; 20mM KCl; 10mM CaCl₂ and0.1% BSA]. A total of 10 μ g plasmid was transformed into 200 μ L protoplast containing 2-4 × 10⁴ cells using PEG-mediated transformation. The cells were cultured in W5 buffer [2mM MES (pH 5.7); 154mM NaCl; 125mM CaCl₂ and 5 mM KCl] in dark condition at 22°C over-night and then subjected to confocal microscope (Leica SP5 laser scanning microscope) examination.

GUS histochemical assay were performed as previously described [8].

Chromatin immunoprecipitation (ChIP) and Western Blot (WB)

Stage 13-16 fruits from *htb-1/crind-1^{ge} pCrIND:CrIND^{K124R}:GFP*, *htb-1* and *crind-1^{ge}* plants were collected and fixed in 1 x PBS buffer containing 1% formaldehyde under vacuum for 15min. Approximately 3.0 g of tissue was ground in liquid nitrogen and nuclear was isolated by filtering with two layer of miracloth (Merck), chromatin fragments were prepared by sonication. After sonication, a 1/20 sample was taken out as DNA Input. The remaining samples underwent immunoprecipitation. GFP-tagged protein together with the associated DNAs were immunoprecipitated by using Pierce Protein G Magnetic Beads (ThermoFisher) coated with monoclonal anti-GFP antibody (Roche) at 4°C for 2 hr. Beads were washed two times with the immunoprecipitation buffer followed by two washes with TE buffer. Reverse crosslinking was done by boiling the beads at 65°C for 12 hours in presence of 1% SDS followed by Proteinase K treatment at 45°C for 1 hour. DNA was ethanol precipitated following phenol/chloroform extraction. qPCR was performed using SYBR Green JumpStart Taq ReadyMix (Sigma) on a BioRad CFX96 Q-PCR System (BioRad).

For detection of the SUMO-conjugation in the cells, ~0.5-g samples of leaf, inflorescence, stage-13 and stage-15 fruit tissues from WT and *htb-1* were fixed in liquid nitrogen and grinded. Total protein was extracted using extraction buffer [50 mM Tris; 150 mM NaCl; 0.2% (v/v) Triton X-100] supplemented with 1X Complete Protease Inhibitor Cocktail (PI, Roche), 20 mM N-Ethylmaleimide (NEM, Sigma) at 4°C for 1 hour. The supernatants were recovered after two steps of centrifuge at 13000 rpm, 4°C for 15 mins. Equal amounts of protein extracts were loading on a standard SDS-PAGE 10% (w/v) acrylamide gel and separated by electrophoresis. Protein was transferred onto a PVDF membrane (GE Healthcare) using a Mini Trans-Blot Cell (Bio-Rad). The membrane was blocked in blocking solution [5% (w/v) dry milk powder in TBST (1X TBS+0.1% Tween 20)] at 4°C for 4 hr. The primary antibody anti-SUMO1 (Abcam, 1:1000) or anti- α -Tubulin (Sigma, 1:5000) was added and incubated over-night at 4°C. The membrane was washed three times with TBST for 10 mins each step, and then incubated with the secondary anti-rabbit antibody (SUMO1, Abcam, 1:10000) or anti-mouse (α -Tubulin, Abcam, 1:10000) in blocking solution for 2 hr. The membrane was washed as described above and exposed to a film (Kodak) using a chemiluminescence reaction with Pierce ECL Western Blotting Substrate (ThermoFisher).

For detection of the SUMOylation of CrIND, the 7-day old *pLhGR>>CrIND*^{K124R}:*FLAG* seedlings were treated with 10 μM dexamethasone (DEX) (Sigma) in liquid MS medium supplemented with 1% sucrose at 22°C for 12 hr. ~1.0-g samples of each genotype were fixed and grinded into a fine powder in liquid nitrogen. The proteins were extracted using GTEN buffer [10% Glycerol; 25 mM



Tris; 1 mM EDTA; 150 mM NaCl; 0.1% (v/v) NP-40] supplemented with 1X PI, 20 mM NEM, 1 mM PMSF (Roche) and 10 mM DTT at 4°C for 1 hr. The supernatants were collected after two steps of centrifuge at 13000 rpm, 4°C for 15 mins and 50 μ L sample was taken out as Input. The remaining lysates were subjected to immune-precipitation using anti-FLAG M2 magnetic beads (Sigma) at 4°C for 2 hr. The beads were then washed with IP buffer [GTEN buffer; 1X PI; 20 mM NEM; 1 mM PMSF and 100 μ M DTT] four times at 4°C for 5 mins each step. 10 μ L Input and 5 μ L IP samples were loaded into a standard 10% acrylamide SDS-PAGE gel. The western blot was conducted according to the aforementioned protocol using either anti-SUMO1 or anti-FLAG (Abcam, 1:5000) antibody.

Population genetics, selection test and phylogeny

To test for evidence of selection at the *HTB* gene, we used polymorphism data for 20 individuals from one *Capsella grandiflora* population and double-checked the results with 13 samples collected in different populations [41]. We conducted a McDonald-Kreitman test (dN/dS) to compare the ratio of synonymous (4-fold degenerate) and non-synonymous (0-fold degenerate) polymorphisms (Pn/Ps) in the coding sequence of *CgHTB* within *Capsella* to fixed differences (Ka/Ks) between *Capsella* and *Arabidopsis* [31]. To assess whether the observed values of Ka/Ks and Pn/Ps were unusual, we compared the observed ratios to Ka/Ks and Pn/Ps at the *CgHTB* gene to ratios of genes in genomic regions with comparable recombination rates, gene densities (in 50-kb windows) and similar expression levels. *P*-values of a two-sided test for a difference between observed Ka/Ks and Pn/Ps and expected Ka/Ks and Pn/Ps were calculated based on the distribution of observed Ka/Ks and Pn/Ps of the comparable genes. In addition, we used the direction of selection (DoS) statistic which describes the direction and extent of selection with positive values indicating positive selection and negative values purifying selection, respectively [42].

For the phylogenetic analysis of ULP family of Cysteine Proteases from *Arabidopsis* and *Capsella*, the full-length protein sequences were downloaded from phytozome database and aligned with Clustal X software [33]. The Neighbor-Jointing (NJ) tree with bootstrap support value was generated based on Protein sequence using MEGA5 software [34].

QUANTIFICATION AND STATISTICAL ANALYSIS

All statistics were calculated in Microsoft Excel. All measured data are presented as means \pm SD specified along with sample sizes (n) in the methods and in figure legends. Comparisons between groups for the analysis of qRT-PCR and fruit characters was performed with Microsoft Excel Student's t test, and significance levels are marked as: *p < 0.05, **p < 0.01.