Overexpression of *SlUPA-like* induces cell enlargement, aberrant development and low stress tolerance through phytohormonal pathway in tomato

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*upa20* induces cell enlargement and hypertrophy development. In our research, overexpression of *SlUPA-like*, orthologous to *upa20*, severely affected the growth of vegetative and reproductive tissues. Wilted leaves curled upwardly and sterile flowers were found in transgenic lines. Through anatomical analysis, palisade and spongy tissues showed fluffy and hypertrophic development in transgenic plants. Gene expression analysis showed that GA responsive, biosynthetic and signal transduction genes (e.g. *GAST1*, *SIGA20OXs*, *SIGA3OXs*, *SIGID1s*, and *SIPRE*s) were significantly upregulated, indicating that GA response is stimulated by overproduction of *SlUPA-like*. Furthermore, *SlUPA-like* was strongly induced by exogenous JA and wounding. Decreased expression of *PI-I* and induced expression of *SlJAZ* (including *SlJAZ2*, *SlJAZ10* and *SlJAZ11*) were observed in transgenic plants, suggesting that JA response is repressed. In addition, *SlUPA-like* overexpressed plant exhibited more opened stomata and higher water loss than the control when treated with dehydration stress, which was related to decreased ABA biosynthesis, signal transduction and response. Particularly, abnormal developments of transgenic plants promote the plant susceptibility to *Xanthomonas campestris pv. campestris*. Therefore, it is deduced from these results that *SlUPA-like* plays vital role in regulation of plant development and stress tolerance through GA, JA and ABA pathways.

*Xanthomonas* causes multiple of crop diseases, such as vascular wils, cankers, leaf spots, fruit spots and blights. The susceptible plant often display pustules on the abaxial surface of leaf, such as the enlargement of mesophyll cell. Palisade and spongy parenchyma cells are strongly enlarged along with the decreased starch contents in chloroplasts, which is similar to the phenotypes induced by AvrBs3. The *AvrBs3* family proteins are type III effectors delivered by *Xanthomonas*. The N-terminal and C-terminal portion of *AvrBs3* are highly conserved and their biological activities are related to the particular repetitive region. The conserved C-terminal contains potent nuclear localization signal (NLS). So, *AvrBs3* like plant transcription factor locates to the nuclear and modulates the transcriptional expressions of host genes. Marois, *et al.* reported that 22 induced and 2 repressed genes are identified, of which 13 genes are confirmed to be induced by *X. campestris pv. vesicatoria* (*Xcv*). The 13 genes were designed as *upal-1-13* (up-regulated by *AvrBs3*), which show high homology to auxin-induced genes, expansion-like genes and pectate lyase genes. Among them, a few gene sequences exhibit low homology to gibberellins (GA)-responsive genes. Therefore, the specific host-gene activation by *AvrBs3* is a complex signaling network, of which the function of phytohormones is indispensable.

Phytohormones regulate plant growth and development, but their signaling pathways are also captured by pathogens to defend against plant defenses. Some effectors or phytoxins secreted by pathogens have evolved to overcome pattern triggered immunity (PTI) of plants and establish the success infections by means of reprogramming plant hormonal pathway. Coronatine (COR) produced by *Pseudomonas syringae*, suppresses the...
SA-mediated defense in planta^{5,6}. Hop1, one of the effectors of P. syringae, disrupts HSP70 (HEAT SHOCK PROTEIN70) in chloroplasts and alters the biosynthesis of SA^{7,8}. Two effectors secreted by Xanthomonas, XopJ and XopD, suppress SA-dependent defense^{9,10}. Effectors also use the antagonistic effects between JA and SA to colonize host plants. HopX1 from Pseudomonas syringae, promotes the degradation of jasmonate ZIM-domain (JAZ) proteins through its central ZIM domain. The JA-induced defenses are activated to counteract SA-dependent defenses^{11}. In general, SA-mediated signaling pathway are involved in resistance to biotrophic pathogens and activation of JA response leads to defense against necrotrophic pathogens^{12}. In addition, a balance of both JA and GA sustains the antagonistic behavior of defense and plant growth. JA is associated with activation of defense against pathogen ingress and GA promotes plant growth at the expense of defense^{13}. GA was firstly identified in the pathogenic fungus Gibberella fujikuroi, the causal agent of excessive elongation of infected plant. Knockout or overexpression of *Elongated uppermost internode (Eui)* genes encoding a GA deactivating enzyme was susceptible and resistant to the infection of Xanthomonas and Magnaporthe oryzae in rice^{14}. However, no information is obtained so far that the effector proteins secreted by pathogens directly interfere with the GA signaling pathway.

Stomata are important entry sites for pathogens ingress. Pathogens have evolved effector proteins or phyto-toxins to counteract host stomal defenses by inhibiting stomatal closure or promoting stomatal opening^{15}. The phytoxoy COR can impede pathogen-associated molecular pattern (PAMP)-induced stomatal closure by the ABA-dependent pathway^{16}. HopX1 from virulent Pta11528 bacteria play an important role in maintaining stomatal aperture^{11}. Likewise, expression of HopP2 results in significantly wider stomatal aperture and insensitivity to PAMP-induced stomatal closure^{17}. Besides, it was reported that activation of ABA signaling pathway could inhibit stomatal opening induced by pathogens. Overexpression of RCAR3, one of ABA receptors, inhibits stomatal reopening during Pst DC3000 infection^{18}. Pst DC3000-induced stomatal closure is affected by the mutation of N Ced1 gene (*notabilis* (*not*) mutant)^{19}. Whether Xanthomonas or AvrBs3 acts on stomatal immunity still needs more evidence.

It is reported that bHLH family gene participates in regulation of GA and JA metabolisms or responses. *PHYTOCHROME-INTERACTING FACTOR3-LIKE5 (PIL5)* inhibits seed germination by repressing the expressions of GA biosynthetic genes in Arabidopsis^{10}. Phytochrome interacting factors (PIFs) control hypocotyl elongation in Arabidopsis seedlings^{21}. MYCs (MYC2, MYC3 and MYC4) act redundantly on the activation of JA response^{22}. But, *upa20*, another bHLH family gene, is not proved to be implicated in alteration of GA or JA response.

*upa20* is the direct target of AvrBs3 in pepper. Transient expression of *upa20* induces hypertrophy phenotypes and decreased starch content in chloroplasts. Hypertrophy of mesophyll cells is attributed to enlargement of cell size, whereas cell division in hypertrophic leaves is not mentioned. There are two homologous genes of *upa20* in tobacco and one in tomato genomes^{19}. Overexpression of *Nt upp-L* results in hyperplasia development. Palisade and spongy parenchyma cells of transgenic leaves are smaller but abundant in number compared with control. The molecular mechanism of hyperplasia development is due to acceleration of cell cycle from G1 phase to mitotic phase, leading to the promotion of cell division^{23}. From these results, *upa20* and *upp-L* in pepper and tobacco participate in different regulatory mechanism. The function of homologous gene is still undetermined in tomato.

Inducing visible cellular changes in transient expression of *avrBs3* selects the scope of the AvrBs3-specific effects to *solanaceous* plants, of which the tomato is selected as our research material. The function of *SIU-PA-like*, homologous gene to *Ca upa20*, was identified in tomato. Constitutive expression of *SIU-PA-like* induced cell enlargement and reduced cell numbers in leaves of wild type tomato, Ailsa Craig (*AC*+). Moreover, overexpression of *SIU-PA-like* improved GA response and susceptibility to pathogen infection together with repressed JA and ABA response.

**Results**

**Cloning and molecular characterization of *SIU-PA-like***. In four distinct subfamilies of Upp/Upla-like transcription factors by the phylogenetic analysis, the gene SL AW034575 from the group 1 of *solanaceous* plants is orthologous to *Ca upa20*, whereas not to *Nt upp-L*. We cloned the full-length cDNA of the SL AW034575 gene from the AC+ by rapid amplification of cDNA ends (RACE) and named it *SIU-PA-like* following the nomenclature of *upa* (upregulated by AvrBs3). The putative *SIU-PA-like* protein has 330 amino acids with a typical bHLH domain region from amino acid 156 to 214, and an estimated molecular mass of 36.3 kDa. Additionally, we searched DNA sequence of *SIU-PA-like* from tomato genome (GenBank accession No.NP_001266190.1). Alignment of the genomic DNA and cDNA showed that the *SIU-PA-like* contained seven exons and six introns.

**The expression profiles of *SIU-PA-like* in different tissues of wild type.** To clarify the role of *SIU-PA-like* in plant development, its transcript accumulation in various tissues was quantified by quantitative RT-PCR. As shown in Fig. 1A, *SIU-PA-like* was highly expressed in stem, mature leaves, flowers and immature fruits, whereas low in root, young leaves, senescent leaves, sepals, and ripe fruits. Besides, pistils and petals showed high *SIU-PA-like* accumulation compared to sepals and stamens in flower tissues (Fig. 1B). These results implied that *SIU-PA-like* is expressed in specific organ that grows larger via cell elongation or enlargement.

**Overexpression of *SIU-PA-like* induces cell enlargement by regulation of cell cycle.** To further analyze the biological function of *SIU-PA-like* in tomato, we generated transgenic tomato plant by overexpressing *SIU-PA-like*. A total of 8 independent transgenic lines were obtained and three of them were selected for further characterization. From Fig. 2A–D, the leaves of *SIU-PA-like*-overexpressed lines (*SIU-PA-like-OE*) showed fluffy and hypertrophic development in both palisade and spongy parenchyma tissues, which leads to thickened blades. From the anatomical analysis, palisade tissue was mostly destroyed in mature leaves in contrast to relative
integrity in young leaves of transgenic lines. Moreover, the mean area of adaxial and abaxial pavement cells were simultaneously enlarged in leaves of transgenic lines (Fig. 2E, Fig. S1). However, the length and width of the first leaflet were both significantly reduced compared to the control (Fig. 2F), although the ratio of length to width was almost unaffected (data not shown), indicating that cell numbers in SIUPA-L-OE leaves were decreased. Likewise, the number of palisade cell per centimeter was decreased in transgenic plants (Fig. 2G). The hypertrophy development is due to cell enlargement, which results from the abnormal expressions of E2F-response genes. In Arabidopsis, over-expressions of KRP1 and KRP2 lead to a compensatory increase in cell volume in response to severely reduced cell number and expressing of KRP1 and KRP4, driven by AtML1 promoter, induces leaves curling upward. In this assay, expression levels of cyclin-dependent kinase inhibitor genes KIP-RELATED PROTEIN1/INHIBITOR1 OF CDC2 KINASE (LeKRs) were determined by RT-PCR. Figure 2H showed that LeKRP3 and LeKRP4 were significantly induced in young leaves of transgenic plants, especially LeKRP4 that is orthologous to the AtKRP1 (Fig. S2). In mature leaves of transgenic plants, LeKRP4 level was still higher than control (Fig. 2I). Besides, expression level of three cyclin genes were investigated. CycA3;1, CycB1;1 and CycD2;1 were significantly repressed in young leaves of transgenic plants as well as CycD2;1 in mature leaves (Fig. 2J,K), indicating that overexpression of SIUPA-like represses the progress of cell cycle and promotes cell enlargement. Therefore, these results revealed that overexpression of SIUPA-like induces cell enlargement, which is due to the repressed progress of cell cycle.

**Overexpression of SIUPA-like interferes with the growth and development of plants.** In addition to inducing the cell enlargement, overexpression of SIUPA-like severely interfered with plant growth and development. At seedling stage, the expression of SIUPA-like in transgenic lines was 2 to 25-fold higher than control (Fig. 3A). Compound leaves of SIUPA-L-OE lines bear sparse leaflets, especially fewer intercalary leaflets. All of the leaflets were curled upwardly, leading to the whole plant looked slender (Fig. 3B,C). In adult plants, the transcripts of SIUPA-like in mature leaves of transgenic lines were 20 to 120-folds higher than AC (+) (Fig. 3A). We observed that SIUPA-L-OE lines cannot bloom and set up fruits (Fig. 3D). So, we could not get any seeds. When the flowers of transgenic plants were opened artificially, the petals, stamens and pistils were much smaller than control (Fig. 3E). The vegetative organs of transgenic lines also showed obvious alterations. The average height of transgenic lines was shorter than control as well as the average diameter (Fig. 3F). The average diameter of plants was measured at 10 cm above the ground. Furthermore, all mature leaves of transgenic lines were curled upwardly and wilted under the normal conditions (Fig. 3G), and the leaf vein was thicker and wider than wide type. From the cross-sectional view, the structure of vascular bundle in leaf veins was disrupted. Nearly all the cells were enlarged except a few phloem cells and even no apparent xylem cells were found (Fig. S3). Meanwhile, the contents of total chlorophyll, chlorophyll a and b were significantly reduced in mature leaves (Fig. 3H). Moreover, axillary buds furthest from the apex of plant could not form (Fig. S4), which resembles pro mutant carrying a point mutation in GRAS region.

**Excessive accumulated transcripts of SIUPA-like promote GA response.** Palisade and spongy parenchyma cells were strongly enlarged, which is similar to GA-induced phenotypes. To determine the reason of hypertrophy phenotype, the endogenous content of GA in control and transgenic lines were investigated. Figure 4A showed that total GA contents in young leaves of transgenic lines were 1.2- to 2-fold higher than control. To reveal the molecular mechanism of GA accumulation in transgenic lines, the transcripts of GA biosynthetic and catabolic genes were examined. Figure 4B showed that expressions of GA synthetic genes were significantly increased, especially SIGA3OX2, whereas no obvious change was found in GA-deactivation gene SIGA2OX1. In tomato genome, there are three putative GA receptor genes, GIBBERELLIN INSENSITIVE DWARF1s (GID1s), which were tentatively termed: SIGID1-A to C (Fig. S5). Two SIGID1s were expressed at extremely high level in young leaves of the transgenic plants, especially SIGID1-B (Fig. 4C), indicating activating GA response may be due...
to the over-accumulated GA receptors. The abundance of GAST1 mRNA (gibberellic acid stimulated transcript 1) is induced by GA3 treatment at transcriptional level. EXP8 is proposed to induce cell expansion and EXP8 expression is induced by exogenous GA. In this assay, the transcriptional levels of both GAST1 and EXP8 in young leaves were significantly increased (Fig. 4D), indicating that overexpression of SlUPA-like promotes GA response. It was reported that increased GA contents and two SlGID1s gene expressions facilitate the formation of GA-GID1-DELLA protein complex, which downregulates the repression of DELLA proteins. Therefore, hindering inhibitory effects of DELLA proteins may be one of strategies to enhance the GA response.

Paclobutrazol resistance (PRE1), a helix-loop-helix protein, alters various aspects of gibberellin-dependent response, such as elongation of petioles. In tomato, there are five putative PREs that were tentatively termed: SlPRE1–5 (Fig. S6). In our research, expression levels of two SlPREs in young leaves of transgenic lines were significantly higher than wild type (Fig. 4E), suggesting that over-accumulated transcripts of two SlPREs contribute to GA response.

Interestingly, endogenous GA contents in mature leaves of SlUPA-L-OE lines were reduced as well as the decreased expression of EXP8 (Fig. 5A,B), whereas GAST1 and four SlPREs (including SlPRE1–SlPRE4, particularly SlPRE4) were remarkably induced (Fig. 5B,C), suggesting that GA response was still stimulated in mature leaves.

**Overexpression of SlUPA-like represses JA response.** In order to characterize the function of SlUPA-like in hormonal response, its expression was investigated under five phytohormonal treatments. The transcripts of SlUPA-like were substantially accumulated within 1 hour under JA treatment (Fig. 6A). JA also has
an important role in wounding induction, which will rapidly increase within a few minutes\(^3^3\). Besides, the expressions of \(JAZ\) genes are simultaneously increased as primary responsive genes to JA treatment and mechanical wounding\(^3^4\). In our experiment, \(SlUPA-like\) expression was rapidly induced within 1 h and peaked at 8 h, followed by a decline but maintained at relatively high level until 24 h when treated with wounding (Fig. 6B), indicating that \(SlUPA-like\) is regarded as a primary responsive gene in response to JA. In transgenic plants, expression of \(PI-I\) was significantly repressed by 85~90% compared to control (Fig. 6C). It was reported that suppressed JA response is due to the \(JAZ\) proteins binding to MYC transcription factors\(^3^5–3^7\). In our experiment, \(SlJAZ2\), \(SlJAZ10\) and \(SlJAZ11\) were severely induced in young and mature leaves of transgenic lines, especially \(SlJAZ10\) and \(SlJAZ11\) (Fig. 6D–F). Phylogenetic analysis showed that At\(JAZ7\), At\(JAZ8\), S\(lJAZ9\), S\(lJAZ10\) and S\(lJAZ11\) belong to one clade and lack the conserved LPIAR motif, which makes At\(JAZ8\) stable in inhibition of JA response\(^3^8\). So, these results suggested that overexpression of \(SlUPA-like\) represses JA response by overexpressions of three \(SlJAZ\) genes.

However, the inhibitory effects of three \(SlJAZs\) were limited. The length of root was repressed by exogenous \(MeJA\) in dose-dependent manner, which was similar to wild type (Fig. 6G). Therefore, the repressed JA response induced by overexpression of \(SlUPA-like\) can be overpassed by the application of exogenous JA, which may be due to the functional redundancy between \(SlJAZs\).

Overexpression of \(SlUPA-like\) stimulates stomata opening under dehydration stress. Stomata controls gas exchange between leaves and environment along with water loss during transpiration. It is known that wide stomatal aperture leads to high water loss. In wild type plants, \(SlUPA-like\) was downregulated after 4 hours of dehydration treatment (Fig. 7A). Moreover, the leaves of \(SlUPA-L-OE\) lines exhibited higher water loss than control under dehydration stress (Fig. 7B). In general, stomatal behavior is the vital index for assessing water loss. Under dehydration stress, the percentage of opened stoma in control was obviously decreased after 2 hours, whereas the percentage was increased in transgenic lines (Fig. 7C), implying that \(SlUPA-like\) promotes stomatal opening and decreases water retention under dehydration stress.

ABA is known for regulation of stomatal closure in response to abiotic stress and pathogen-associated molecular pattern (PAMP)-triggered immunity\(^3^9\). In order to discover the reason of stomatal behavior in transgenic lines, the expression levels of ABA biosynthetic, signal-transduction and responsive genes were monitored.
**SlNCED1** was primarily involved in ABA biosynthesis in tomato and activation of SnRK2 by ABA can phosphorylate downstream effectors, leading to activation of ABA response. In mature leaves of SlUPA-L-OE plants, expression of **SlNCED1** was remarkably repressed as well as **SlSnRK2.3**, implying that ABA biosynthesis and signal-transduction may be both suppressed (Fig. 7D). Meantime, three ABA responsive gene expressions, **CAT1**, **GME2** and **LEA**, were obviously repressed by 54–92% in transgenic plants, particularly **LEA**, indicating that ABA response is suppressed in transgenic plants (Fig. 7E). So, stimulating stomatal open in transgenic lines may be due to the inhibition of ABA response.

**Overexpression of SlUPA-like improves plant susceptibility to pathogens.** We have shown that GA response was excessively improved, instead JA response was repressed, and the percentages of opened stoma of SlUPA-L-OE plants were increased under stress condition. Given that GAs negatively regulate disease resistance of host plants and deficient JA level positively regulates the susceptibility of plant to *X. campestris*, stomatal opening provides access for infections of *Xcv*, we logically speculate that SlUPA-like overexpressed plants may be susceptible to pathogens. To test this hypothesis, leaflets of transgenic and control plants were inoculated with *Xanthomonas campestris pv. campestris* (*Xcc*). As expected, the leaflets of transgenic lines showed...
yellow spot after 3 days of inoculation and spread out most parts of the blades after 8 days, whereas the control leaflets were unaffected all the time (Fig. 8). To further determine \( Xcc \) infection, a 253 bp DNA fragment from genome of \( Xcc \) was amplified in leaves of both \( AC^{++} \) and transgenic lines. The PCR products were only accumulated in transgenic lines (Fig. S7). Additionally, Fig. S8 showed that \( SlUPA-L \)-OE plants were more susceptible to plant disease than \( AC^{++} \) when exposed to infected plants. These data demonstrated that overexpression of \( SlUPA-like \) improves the susceptibility of tomato plants to pathogens.

**Discussion**

GA-GID1-DELLA protein complex may contribute to the cell enlargement of young leaves. Hypertrophy development is attributed to cells enlargement along with decreased contents of starch\(^2\), which are similar to phenotypes of excessive GA accumulation. Loss-of-function of \( eui \) mutant shows elongated internode and reduced development of starch granule\(^4\). Experiment confirmed that overexpression of \( SlUPA-like \) induced cell enlargement and GA response (Figs 2 and 5). \( LeKRP \) genes were remarkably increased in both young and mature leaves of transgenic plants, together with the suppressed expressions of cell cycle related genes (Fig. 2), suggesting that overexpression of \( SlUPA-like \) facilitates the DNA endo-replication and inhibition of cell cycle. Meantime, both transcripts of \( SlGID1-A \) and \( SlGID1-B \) were significantly increased along with increased GA contents (Fig. 4C). Through phylogenetic analysis, \( AtGID1-b, SlGID1-A \) and \( SlGID1-B \) belong to the same clade (Fig. S5). \( AtGID1b \)-overexpressions can rescue GA response by forming the GA-GID1-DELLA protein complex\(^3\). In young leaves of transgenic plants, the key elements for formation of GA-GID1-DELLA protein complex coexisted. However, in mature leaves of transgenic lines, GA content were decreased (Fig. 5A), implying that this kind of protein complexes is probably destroyed. Therefore, the GA-GID1-DELLA protein complex probably contributes to the enhanced GA response and cell enlargement in young leaves of transgenic lines.

Expressions of \( GAST1 \) are probably induced by overexpression of \( SYPRE \) independent of GA synthesis. In the young leaves of transgenic lines, expression levels of \( SYPRE3 \) and \( SYPRE4 \) were significantly increased compared to wide type (Fig. 4E). Similarly, transcripts of four \( SYPRE \), \( SIPRE1-SIPRE4 \), were upregulated in the mature leaves and particularly the expression of \( SIPRE4 \) was further magnified compared to young leaves (Fig. 5C). Meanwhile,
GAST1 expression was both increased in young and mature leaves, especially in mature leaves (Figs 4 and 5), indicating that expression of GAST1 was sequentially induced by SlPREs. However, the expression of EXP8 was suppressed in mature leaves (Fig. 5B), implying that the GAST1 was regulated by different upstream signaling pathways. It is reported that EXP8 does not respond to ABA and water, but specially responds to GA30. By contrast, expression of GAST1 was response to both GA and ABA 29. These results suggested that GAST1 locates further downstream in GA signaling pathway. Moreover, it was proved that GA stimulates cell elongation through activation of PRES, which locate downstream point of GA biosynthesis48. Therefore, continuous GA response in transgenic lines was attributed to overexpression of SlPRE, which may be independent of GA biosynthesis.

Repressed JA response may not result in female sterility of flowers in transgenic lines. Plant protease inhibitors (PIs) can be regarded as defensive proteins to minimize the adverse effects from the attack of phytophagous insects or pathogens. In our experiment, the expression of PI-I, JA induced gene, was strongly repressed in SlUPA-like-overexpressed lines (Fig. 6C), suggesting that JA response is suppressed. However, the expression of PI-II was unaffected by overexpression of SlUPA-like (Fig. 6C), although its expression can be induced by JA 49. Therefore, it need further exploration that JA response can be monitored by PI-I expression.

jasmonic acid–insensitive1 (jai1) mutant is defective in JA signaling of tomato, exhibiting female sterility50. jai1 mutants can set up fruit upon pollination, although the stigmas of flowers protrude from the anther cone during flower development51. That is, the anther filaments are not capable of reaching the position above the stigma at the time of flower blooming51. However, flowers in SlUPA-L-OE lines were sterile and their phenotypes were not similar to jai1 (Fig. 3E). Moreover, the sterile flowers of transgenic lines didn’t have the ability to parthenocarpy when treated with 2. 4-D (data not shown). The reason of infertility of SlUPA-L-OE lines needs further research.

SlUPA-like may directly regulate the expressions of SlJAZs, SIGIIDs, SlPREs, SlJAZs and GAST1 were extremely induced by SlUPA-like (Figs 4–6). It was hypothesis that part or all of these genes contain the binding site for bHLH family is
CANNTG (where N can be any nucleotide), in which the canonical sequence is CACGTG motif. Within 1.5-kb upstream of the initiator-ATG, the numbers of putative binding site for bHLH were counted (Table S1). The numbers of binding sites in promoters of SIGIDs and SIPREs were not accordance with their expression levels, indicating that overexpression of two SIGIDs and four SIPREs may be indirectly regulated by SIUPA-like. The expression patterns of three SlJAZs (SlJAZ2, SlJAZ10, and SlJAZ11) largely accorded with the distribution of binding sites in their promoters (Table S1), revealing that these genes may be direct targets of SIUPA-like. Besides, the expression of SlJAZ9, homolog to SlJAZ11, is not induced by SIUPA-like. Previous works proved that G-rich DNA element is capable of formatting secondary structure and functions as repressor elements. Within the promoter of SlJAZ9, Poly (G) motif from −1453 to −1733 bp makes it hard to be bonded by transcription factors. In addition, the promoter of GAST1 contained no classical binding site, implied that excessive expression of GAST1 may be mediated by other factors. Nevertheless, all these results from bioinformatics analysis were not sufficient to make conclusion, which need to be validated by chromatin immunoprecipitation.

SIUPA-like may participate in the crosstalk between GA and JA response. In transgenic plants, expression levels of three SlJAZs were extremely up-regulated in transgenic leaves along with repressed JA response (Fig. 6). Meantime, SIPREs were significantly induced in transgenic plants with the activation of GA response. These data suggested that SIUPA-like may mediate the crosstalk of GA and JA response. Overexpression of AtJAZs has GA-hypersensitivity phenotypes, particularly AtJAZ9 whose overaccumulation could be resistant to exogenous paclobutrazol. In our research, extremely upregulated expressions of SlJAZs were found in transgenic plants. Moreover, overexpression of SIPREs shows GA response (Figs 4 and 5). So, stimulated GA response in transgenic plant may be due to the over-accumulation of both JAZ and PRE proteins. It was reported that JAZ9 interferes with DELLA-PIF3 interactions and competes with DELLA targets. So, overexpression of SlJAZ9/10 may destroy the inhibitory effect of DELLA proteins. However, DELLA proteins don’t interact with AtJAZ2, AtJAZ7 and AtJAZ8 in vitro. Whether the interactions between three SlJAZs and DELLA proteins mediated by other factors still needs to be tested. In addition, ILI1 and PRE1 inactivate the inhibitory function of IBH1 (another bHLH protein) through hetero-dimerization. Therefore, it was possible that SIUPA-like protein is another target of PREs in tomato.

Stimulating stomatal opening may be mediated by ABA response in transgenic lines. Locating the epidermis of leaves, stoma is primary organ to lose water during transpiration and is also the key channel for pathogen infection. Plant have evolved stomatal defense to fight against the infection of pathogens. Certainly, plant pathogens secreted toxin or effectors to counteract stomatal defense to promote the stomatal reopening. In this assay, when treated with severe water loss, most of stomatal pores closed in control plant, whereas more stoma in transgenic plants reopened (Fig. 7C). Meanwhile, expressions of ABA biosynthetic, signal-transduction and responsive genes were significantly suppressed in transgenic lines (Fig. 7D,E). To some extent, the stomatal behavior under infection of pathogen could be reproduced through dehydration treatments. Thus, stimulated stomatal opening by overexpression of SIUPA-like may be mediated by suppressed ABA response.

Aberrant development facilitates the infections of pathogens in transgenic lines. Increased GA response, suppressed JA response and stimulated stomatal opening give us clues why overexpression of SIUPA-like leads to aberrant development (including curled and wilted leaves, sterile flowers and so on). It was reported that alterations of hormonal pathways improved plants susceptibility to Xanthomonas, which infection was mediated by stomatal behaviors. Plant pathogens firstly gain entry into plant tissue via wounds or natural openings such as stomata. In order to counteract stomatal defense of host plant, pathogens can evade host immunity by suppressing the stomatal closure. For these reasons, stomatal opening promoted by overexpression of SIUPA-like together with reprogrammed GA and JA response enhanced infection efficiency (Fig. 8), although the Xcc is not specific pathogen for solanaceous plants. This study revealed that SIUPA-like can be regard as a landmark gene in monitoring the disease tolerance of plants.

**Methods**

**Plant Materials and Growth Conditions.** In our experiment, *Solanum lycopersicum* Mill. cv. Ailsa Craig (AC++) a near-isogenic tomato line, was used as model plant grown in greenhouse (16-h-day/8-h-night cycle, 25 °C/18 °C day/night temperature, 80% humidity, and 250 μmol m⁻² s⁻² light intensity). Plants of the first generation (T0) came from tissue culture. Tissue culture plantlets from transgenic lines and control with no construct were mixed, and stored at −80 °C until RNA extraction. Each treatment repeated with two times. (b). Dehydration treatments. (c) Phytohormones treatments. 35-day-old AC++ seedlings were used for treatments. Tomato seedlings were sprayed with different concentration of phytohormones (50 μM IAA, 50 μM GA3, 50 μM MeJA, 100 μM ABA and 50 μM ACC) respectively until the water dripped, and then immediately enclosed in transparent bags with some small holes (about 0.5 cm in diameter) on the surface. After treatments for 0, 1, 2, 4, 8, 12 and 24 hours, the leaves of seedlings were harvested for analysis. All the samples were immediately frozen and stored at −80 °C until RNA extraction. Each treatment repeated with two times. (b). Dehydration treatments.

**Sequence, structure and phylogenetic analysis.** The exons and introns of genomic DNA were analyzed by GENESCAN (http://genes.mit.edu/GENESCAN.html). The theoretical molecular weight was calculated with the Expasy compute Mw tool (http://expasy.org/tools/protparam.html). Multiple sequence alignment was conducted using the ClustalX 1.81 and DNAMAN 5.2.2 programs.

**Plant treatments.** (a) Phytohormones treatments. 35-day-old AC++ seedlings were used for treatments. Tomato seedlings were sprayed with different concentration of phytohormones (50 μM IAA, 50 μM GA3, 50 μM MeJA, 100 μM ABA and 50 μM ACC) respectively until the water dripped, and then immediately enclosed in transparent bags with some small holes (about 0.5 cm in diameter) on the surface. After treatments for 0, 1, 2, 4, 8, 12 and 24 hours, the leaves of seedlings were harvested for analysis. All the samples were immediately frozen and stored at −80 °C until RNA extraction. Each treatment repeated with two times. (b) Dehydration treatments.
35-day-old AC\(^{++}\) seedlings were carefully pulled out from soil, washed with water and left on a piece of dry filter paper with 60% humidity at 25 ± 1 °C. After 0, 1, 2, 4, 8, 12, 24, 48 and 72 hours, the leaves were collected, frozen in liquid nitrogen and stored at −80 °C. The transgenic plants and wild type were planted in greenhouse and each line contains 3–5 plants. When the control set up fruits, mature leaves of transgenic lines and control were harvested from top, middle and lower part of each plant, and left on a piece of dry filter paper with 60% humidity at 25 ± 1 °C. After different hours, the statistical analysis of opened stoma in mature leaves was performed. (c). Mechanical wounding treatments. 35-day-old AC\(^{++}\) seedlings were used for mechanical wounding treatment. The leaflets were cut with scalpels into small pieces and left on a piece of wet filter paper in sealed pots for 0, 1, 2, 4, 8, 12 and 24 hours, then the pieces of leaves were collected, respectively.

**Construction of SIUPA-like overexpression vector and plant transformation.** Total RNA of AC\(^{++}\) leaf was extracted using Trizol (Invitrogen) according to the manufacturer’s instructions. Then, 0.5 μg of total RNA was used to synthesize first-strand cDNA through reverse transcription-PCR using Moloney murine leukemia virus reverse transcriptase (Takara) with tailed oligo(dT)\(_{18}\) primer. 2 μL of cDNA was used to clone the full-length cDNA with primers of SIUPA-like-F and d'T-R through high-fidelity PCR (Prime START mix DNA polymerase; Takara). The amplified products were tailing by using the DNA-Tailing kit (Takara) and linked with pMD18-T vector (Takara). Positive clones were picked out via *Escherichia coli* full-length cDNA with primers of SIUPA-like described protocols\(^\text{58}\). Positive transgenic plants were selected on kanamycin medium and screened by polymerase chain reaction analysis with primers NPTII-F and NPTII-R. Then, T0 plants from each transgenic line were propagated through apex culture in vitro propagation.

**The percentages of open stomata under dehydration stress.** Adult plants of AC\(^{++}\) and SIUPA-like overexpressed lines were planted in field. At 10:00 am, the mature leaves were sampled at the same position from the apex of compound leaves and immediately abaxial stoma of detached leaves were inspected with microscopy. The percentage of open stomata was calculated. Then, the detached leaves were placed in wet filter paper. After 1, 2, 3 and 4 hours of dehydration treatments, the percentage of opened stomata was calculated, respectively. All experiments were repeated 3 to 5 times.

**The root growth inhibition by MeJA.** The aseptic axillar buds with approximately same size from AC\(^{++}\) and transgenic lines were planted in rooting MS medium. The different concentration of methyl jasmonate (MeJA), 0 μM, 10 μM, 20 μM and 50 μM, was added into rooting medium, respectively. After 25 days of cultivation, the lengths of roots were calculated. Three biological replicates were used for each treatment.

**Quantitative real-time PCR analysis.** Total RNAs from different tissues of AC\(^{++}\) and transgenic lines were extracted. Quantitative real-time PCR was performed using the SYBR Premix Ex Taq II kit (Promega) in a 10 μL total sample volume (5.0 μL of 2 × SYBR Premix Ex Taq, 0.5 μL of primers, 1.0 μL of cDNA, and 3.5 μL of deionized water) under the following conditions: 95 °C for 2 min, followed by 40 cycles of 98 °C for 15 s and annealing temperature for 30 s. To remove the interference of genomic DNA and the template from the environment, no-template control and no-reverse transcription control experiments were performed. Additionally, three replications for each sample were used, and standard curves were run simultaneously. The tomato *SICAC* and *SIEF1a* genes were used as internal standards\(^\text{59}\). Primers used for quantitative RT-PCR are shown in supplementary Table S2.

**Microscopy.** The detached leaves sampled from AC\(^{++}\) and transgenic lines were immediately fixed by formalin–acetic acid–alcohol fixative (FAA). Then, steps of dehydration, fixation, sectioning and dewaxing were operated on the fixed materials. Finally, the treated materials were observed under the microscope.

**Chlorophyll quantification in tomato leaves.** For chlorophyll measurement, leaves were sampled from control and transgenic plants at the same height and simultaneously weighed. Samples were grinded into powder with liquid nitrogen, extracted with 10 mL 80% aqueous acetone (v/v). The extracts were centrifuged at 4, 000 g for 5 min and the absorbances of the supernatant were measured at 646 and 663 nm, using a lambda 900 scanning spectrophotometer (PerkinElmer). Total chlorophyll contents were calculated by the following equation: Chl\(_{a}\) (mg/mL) = 20.29(OD645) + 8.02(OD663), Chl\(_{a}\) (mg/mL) = 12.7(OD663) – 2.69(OD645), and Chlb (mg/mL) = 22.9 (OD645) – 4.677 (OD663). OD represents the absorbance, while 645 and 663 represent the wavelength. Chl, Chl\(_{a}\) and Chlb represent total chlorophyll, chlorophyll a and chlorophyll b, respectively.

**Quantification of Gas.** In this assay, quantity of endogenous GAs was determined by GA ELISA Kit (Sangon Biotech, Shanghai, China). The fresh samples including young and mature leaves from AC\(^{++}\) and SIUPA-L-OE lines were grinded in liquid nitrogen. About 1 g treated sample was extracted with 80% methanol and centrifuged at 10,000 g for 15 min. The supernatant was partitioned against ethyl acetate and purified by Sephadex chromatography and C18 cartridges\(^\text{60}\). Standards or samples were then added to the appropriate microtiter plate wells with a biotin-conjugated polyclonal antibody preparation specific for GA and Avidin conjugated to Horseradish Peroxidase (HRP) was added to each microplate well and incubated. Then a TMB substrate solution was added to each well. Only those wells that contain GA, biotin-conjugated antibody and enzyme-conjugated Avidin will exhibit a change in color. The enzyme-substrate reaction was terminated by the addition of a sulphuric acid.
solution and the color change was measured spectrophotometrically at a wavelength of 450 nm ± 2 nm by Multimode Reader. The concentration of GA in the samples was determined from the standard curve.

**Infection experiments.** In consideration of infectivity of *Xanthomonas campestris pv. campestris* (Xcc), mechanical injuries were caused on leaflets of AC− and transgenic lines by toothpick. Immediately, 1 µl inoculums were placed into wounds. Then, the leaflets were sealed by one transparent plastic bag. A few days later, bacterial growth in leaflets was monitored. After 3 and 8 days of inoculation, the inoculated leaves were sampled for DNA extraction by CTAB. A 253-bp genomic DNA fragment of Xcc was amplified with primers (Table S2). A 253-bp fragment of M. oryzae 28S rDNA was amplified from fungal genomic DNA using the primers 5′–TCGCCCTACCGAGAAATCCC–3′ and 5′–CCGAACCTTGGACCGCAT–3′, to monitor the infections of Xcc and the products were detected by agarose gel electrophoresis.

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**Acknowledgements**

This work was supported by National Natural Science Foundation of China (no. 31572129), and the Natural Science Foundation of Chongqing of China (cstc2015jcjA80026), and the Fundamental Research Funds for the Central Universities (No. 106112015CD)JZR235504).

**Author Contributions**

G.C. and Z.H. designed and managed the research work and improved the manuscript. B.C., J.H., Y.Z., W.Y., Z.Z. and Y.F. performed the experiments. B.C. wrote the manuscript.

**Additional Information**

Supplementary information accompanies this paper at http://www.nature.com/srep

**Competing financial interests:** The authors declare no competing financial interests.

**How to cite this article:** Cui, B. et al. Overexpression of SIUPA-like induces cell enlargement, aberrant development and low stress tolerance through phytohormonal pathways in tomato. *Sci. Rep.* **6**, 23818; doi: 10.1038/srep23818 (2016).