A phosphorylation-based switch controls TAA1-mediated auxin biosynthesis in plants

Qian Wang, Guochen Qin, Min Cao, Rong Chen, Yuming He, Liyuan Yang, Zhejun Zeng, Yongqiang Yu, Yangtao Gu, Weiman Xing, W. Andy Tao & Tongda Xu

Auxin determines the developmental fate of plant tissues, and local auxin concentration is precisely controlled. The role of auxin transport in modulating local auxin concentration has been widely studied but the regulation of local auxin biosynthesis is less well understood. Here, we show that TRYPTOPHAN AMINOTRANSFERASE OF ARABIDOPSIS (TAA1), a key enzyme in the auxin biosynthesis pathway in Arabidopsis thaliana is phosphorylated at Thr 101 (T101). T101 phosphorylation status can act as an on/off switch to control TAA1-dependent auxin biosynthesis and is required for proper regulation of root meristem size and root hair development. This phosphosite is evolutionarily conserved suggesting post-translational regulation of auxin biosynthesis may be a general phenomenon. In addition, we show that auxin itself, in part via TRANSMEMBRANE KINASE 4 (TMK4), can induce T101 phosphorylation of TAA1 suggesting a self-regulatory loop whereby local auxin signalling can suppress biosynthesis. We conclude that phosphorylation-dependent control of TAA1 enzymatic activity may contribute to regulation of auxin concentration in response to endogenous and/or external cues.
The dynamics of hormone concentrations are essential for developmental processes in both plants and animals. As widely reported in multicellular eukaryotic systems, different concentrations of each hormone can trigger diverse developmental outputs. An imbalance in hormone levels can cause a range of illnesses in animals and abnormal development in plants. Auxin is a highly concentration-dependent hormone in plants that regulates all major aspects of plant development. It has been intensively reported that the dynamic auxin concentration gradients are established along with the plant organs, such as roots and that the local concentration of auxin determines distinct developmental fates. Furthermore, local auxin concentration adjusts quickly in response to environmental changes. Polar auxin transport plays an important role to modulate local auxin levels, but local auxin production determines the overall auxin content within a plant and also plays an essential role in plant development and in adaptation to environmental changes.

Indole-3-acetic acid (IAA), the primary natural auxin in plant, is synthesized by multiple pathways. The indole-3-pyruvic acid (IPA) pathway mediated by TAA1/TARs (Tryptophan Aminotransferase of Arabidopsis 1/ Tryptophan Aminotransferase Related proteins) and YUC (YUCCA) is a well-established auxin biosynthesis pathway that contributes the majority of free IAA production. IAA is tightly linked to the local concentration of auxin. Accumulation of auxin promotes cell division while lower auxin concentration triggers cell differentiation, which determines root meristem size. Auxin stimulates root hair development in the maturation zone as a way for plants to adapt to environmental changes. As previously reported, auxin biosynthesis mutants show strong defects in both root apical meristem and root hair development. Moreover, TAA/YUC-mediated auxin biosynthesis optimizes plant growth in response to a range of environmental changes. In Arabidopsis root development, the dynamically maintained meristem, elongation, and maturation zones are tightly linked to the local concentration of auxin. Accumulation of auxin promotes cell division while lower auxin concentration triggers cell differentiation, which determines root meristem size. Auxin stimulates root hair development in the maturation zone as a way for plants to adapt to environmental changes.

Results

Phosphorylation at T101 regulates ATAA1 enzymatic activity. To investigate the underlying regulatory mechanism of auxin biosynthesis at the non-transcriptional level, we used mass spectrometry (MS) to identify the potential protein modifications of auxin biosynthesis enzymes in Arabidopsis. First, we analysed TAA1, the key enzyme that converts tryptophan to IPA in the presence of the pyridoxal phosphate (PLP) cofactor. We generated pTAA1-TAA1-GFP transgenic plants, treated these with a phosphatase inhibitor to prevent protein dephosphorylation, and used immunoprecipitated TAA1-GFP proteins for mass spectrometric analysis (Supplementary Fig. 1). Interestingly, we identified a phosphorylation site at T101 within the in vivo immunoprecipitated TAA1 protein (Fig. 1a). According to the TAA1 protein structure, the T101 residue is located within the PLP binding pocket, indicating that phosphorylation of TAA1 at T101 may affect TAA1 enzymatic activity (Fig. 1b). To verify this, we mutated the T101 residue to aspartic acid (T101D) to mimic the phosphorylation state and tested its enzymatic activity in vitro. We purified various mutated TAA1 proteins from Escherichia coli and separated the proteins using a native gel, then stained the gel using a catalytic reaction buffer (Method section). In this way, the active TAA1 would catalyse transamination reaction then result in a dark colour in the gel. TAA1K217A protein was set as a control, as the K217 residue is reported to be required for PLP binding. In contrast to the TAA1WT protein, which displayed the colour of reaction products in the gel, TAA1T101D protein was not active in the assay suggesting that E. coli-purified TAA1T101D proteins completely lost their transaminase catalytic activity similar to the TAA1K217A proteins (Fig. 1c). We used another liquid assay and confirmed that the TAA1T101D proteins were unable to convert tryptophan to IPA (Fig. 1d). To investigate whether the non-phosphorylated form has the opposite effect on TAA1 enzymatic activity to the phosphorylated form, we mutated T101 to alanine (T101A). Interestingly, we found TAA1T101A proteins exhibited similar or even higher enzymatic activity compared with TAA1WT only when PLP cofactors were present in a sufficient amount in the assay (Fig. 1c, d; Supplementary Fig. 2a, b). Under PLP-deficient conditions, neither TAA1T101A or TAA1T101D proteins showed normal enzymatic activity due to an abnormal ability to capture PLP from E. coli which is distinct from E. coli-purified TAA1WT proteins that are already bound to PLP (Supplementary Fig. 2a–c). This indicates that the T101 residue is indeed important for PLP-dependent enzymatic activity of TAA1 and that the T101A mutation does not fully simulate non-phosphorylated T101.

To further investigate the function of T101 phosphorylation in vivo, we introduced the TAA1 promoter-driven TAA1WT, TAA1T101D, and TAA1T101A into the wei8-3 (TAA1 is also known as weak ethylene insensitive 8, WEI8) mutant which is a strong TAA1 loss-of-function mutant. Consistent with previous work, blocking TAA1 function in Arabidopsis by either genetic mutation or the chemical inhibitor L-kyurenine (L-Kyn) impairs the root apical meristem and root hair development, and this can be rescued by exogenous auxin application, thus providing a good system to study how auxin levels are controlled (Supplementary Fig. 3). Compared with TAA1WT, we found that TAA1T101D could not complement either the root meristem or the root hair phenotype in the wei8-3 mutant, indicating an abolished function of TAA1T101D in vivo (Fig. 1e, f; Supplementary Fig. 4). TAA1T101A only partially rescued the mutant phenotype and could not fully rescue root meristem phenotype of wei8-3;tar2-1 mutant, suggesting that TAA1T101A is not fully functional in vivo even though PLP slightly increased root meristem size of TAA1T101A, wei8-3 (Fig. 1e, f; Supplementary Fig. 4; Supplementary Fig. 5). These results are consistent with the biochemical results and indicate that TAA1 activity is switched off when T101 is in a phosphorylated state in plants.
endoplasmic reticulum-localized TAR2 have redundant functions in auxin biosynthesis. Interestingly, when overexpressing TAA1[T101D] in weib-3;tar2-1 mutants, we observed dominant negative effects on plant growth. 35S-TAA1[T101D]; weib-3;tar2-1 seedlings exhibited a more enhanced phenotype compared to weib-3;tar2-1 (Fig. 2a–h). Fifty four out of 342 35S-TAA1[T101D]; weib-3;tar2-1 seedlings (15.79%) showed a strong phenotype with the abnormal cotyledon and/or the root retardation similar to weib8;tar1;tar2 triple mutant, while only 8 of 451 (1.77%) of weib-3;tar2-1 seedlings showed this phenotype. This indicated that the function of TAR1 in 35S-TAA1[T101D]; weib-3;tar2-1 might be partially abolished.

**Fig. 1** Phosphorylation modification on T101 residue turns off TAA1 enzymatic activity. a Mass spectrometric analysis showing the phosphorylation modification at Thr101 site on TAA1. Structural simulation of the connection between T101 residue and PLP cofactor. In gel tryptophan aminotransferase activity (Trp-AT) of E. coli-puriﬁed GST-TAA1WT, GST-TAA1[T101A], and GST-TAA1[T101D]. GST-TAA1[K217A] was set as a control. Protein amount shown using Coomassie blue staining. E. coli-puriﬁed proteins were separated in a native gel followed by the catalytic reaction and further staining. The intensity of dyed bands in the gel represent catalytic activity. Lack of a dark band in the GST-TAA1[T101D] and GST-TAA1[K217A] lanes indicate complete loss of enzymatic activity. Three independent biological repeats showed similar results. Transaminase catalytic activity detected by measuring IPA production with Salkowski reagent (Methods section). Transaminase catalytic reactions were performed in the reaction reagent containing 2.5 μg recombinant protein with a His tag. The products were measured by reading the absorbance at 530 nm at different reaction time points. Values denoted the mean ± s.d. (n = three independent biological repeats). Representative pictures of root apical meristem from 5-day-old seedlings in Col-0, weib-3, and pTAA1-gTAA1 (WT, T101D, T101A); weib-3 complementation transgenic plants. White arrows show the meristem zone; Scale bar 50 μm. Quantification of root meristem size in e. Three independent lines of TAA1WT; weib-3 and TAA1[T101D]; weib-3, and 2/7 lines of TAA1[T101A]; weib-3 showed similar results. n denotes the number of independent seedlings; one-way ANOVA with Tukey multiple comparisons test. Different letters represent signiﬁcant difference between each other, P < 0.0001.
Interestingly, we found that TAA1 and TAR1 were able to form heterodimers as tested by coimmunoprecipitation assay (Supplementary Fig. 6). Together, these data suggest that phosphorylation of the TAA1 protein at the T101 site not only controls the on–off switch of TAA1 enzymatic activity but may also affect the functions of other TAA1 homologs by dimerization.

**T101 phosphorylation on TAA1 is evolutionarily conserved.** When we examined the TAA orthologs in other plant species, we discovered that the T101 residue is remarkably conserved across different plant species (Fig. 2i; Supplementary Fig. 7), including evolutionary distant species, such as *Marchantia Polymorpha*, in which auxin signalling is already present despite being ancestral to land plants.30–32 This finding implies that the T101-mediated switch of AtTAA1 may be conserved during evolution. To provide evidence for this hypothesis, we tested whether MpTAA was also phosphorylated by MS using 35S-MpTAA transgenic *Marchantia*. We found Thr224 on MpTAA protein, a conserved residue analogous to T101 of AtTAA1, was also phosphorylated in vivo (Fig. 2i; Supplementary Fig. 8). Then we tested the function of a phosphor-mimic mutation of MpTAA in transgenic *Marchantia*. As expected, the phosphor-mimic version (T224D) of MpTAA also affected the function of MpTAA (Fig. 2j, k). It has been reported that MpTAA mediates auxin biosynthesis and is essential for the development of *Marchantia*.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.33–35 Both the mutation of MpTAA and the treatment with the TAA inhibitor (L-Kyn) suppress thallus growth, while overexpression of MpTAA in wild-type *Marchantia* (Tak-1) can enhance its resistance to the L-Kyn treatment.
that TAA1 could be targeted and phosphorylated by protein kinases. To identify the potential kinases that regulate TAA1, we performed a mass spectrometric analysis of the TAA1 complex proteins in young Arabidopsis seedlings and found candidate protein kinases. We then used the unbiased yeast two-hybrid assay to screen the interactions between protein kinase candidates and TAA1. We found that TMK4, a member of the transmembrane kinase family proteins in the regulation of auxin signaling, specifically interacted with TAA1 in yeast (Fig. 3a). We then confirmed this interaction using an in vitro pull-down assay (Fig. 3b). In addition, we found that TMK4 was coimmunoprecipitated with TAA1 in vivo (Fig. 3c). Thus, although TMK4 proteins localized at the plasma membrane and TAA1 proteins distributed in the cytosol, which we observed in transgenic plants driven by native promoters (Supplementary Fig. 9), a portion of cytosolic TAA1 at the submembrane was targeted by TMK4.

Using an in vitro kinase assay, we showed that TMK4 phosphorylated TAA1 directly (Fig. 3d). To verify the phosphorylation sites targeted by TMK4, we detected the phospho-peptide of TAA1 by MS after the kinase reaction in vitro, and found the pT101-containing phospho-peptide indicating TMK4 could phosphorylate TAA1 at the T101 site (Supplementary Fig. 10a). By mass spectrometric analysis, we further quantified the phosphorylation level of T101 on TAA1 protein in vivo and

**Fig. 3 T101 of TAA1 is targeted by TMK4 kinase.** a Yeast two-hybrid assay of multiple protein kinases (bait) and TAA1 protein (prey). Three independent biological repeats. b In vitro pull-down assay showing GST-TMK4 kinase domain (KD) can be pulled down by TAA1-His proteins. His-Sumo proteins were used as a negative control. TMK4 KD was detected via western blot using an anti-GST antibody. TAA1-His and His-Sumo shown by Coomassie blue staining (CBB). Three independent repeats showed similar results. c TMK4 was coimmunoprecipitated with TAA1 in protoplasts. 35S-TMK4-HA and 35S-TAA1-Flag were transiently coexpressed in Arabidopsis protoplasts. Single transformation of 35S-TMK4-HA was used as control. Three independent repeats showed similar results. d In vitro kinase assay of TMK4 kinase domain (KD) on TAA1 protein. Three biological repeats. e Mass spectrometric analysis of the phospho-peptide intensity in Col-0 and tmk4-1. The intensity indicates the phosphorylation level of T101 containing phospho-peptide (93aa–116aa). Standardization and another two repeats were shown in Supplementary Fig. 10b, c. f Quantification of TAA1 enzymatic activity with or without TMK4KD phosphorylation. E. coli-purified His-sumo-TMK4KD and TAA1 proteins were mixed together to perform a phosphorylation reaction triggered by ATP before the in-gel transaminase catalytic reaction (Methods section). ATP+ indicated the phosphorylation reaction between TMK4KD and TAA1 with ATP, and ATP– indicated the abolished phosphorylation reaction without ATP. P values were shown as indicated (n represents five biological repeats); two-sided t-test. Centre line in box represents mean and whiskers show minimum to maximum.

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found that it was decreased in the tmk4-1 mutant compared to wild type (Fig. 3e; Supplementary Fig. 10b, c). Thus, TMK4 is one of the protein kinase candidates that targets the T101 site on the TAA1 protein.

Considering the phosphorylation status of T101 was able to determine the on–off switch of TAA1 enzymatic activity, we further verified whether TMK4 could regulate TAA1 enzymatic activity through phosphorylation at T101. We mixed both TMK4 and TAA1 proteins to perform a phosphorylation reaction triggered by ATP prior the in-gel transaminase catalytic reaction. As expected, we found that TMK4 significantly inhibited the catalytic activity of TAA1WT but not TAA1T101A (Fig. 3f and Methods section). The transaminase activity of total protein from the tmk4-1 mutant increased significantly compared to wild type, but was rescued in the tmk4-1;wei8-3 double mutant (Supplementary Fig. 11a), indicating TMK4 negatively modulates TAA1 enzymatic activity in vivo. Meanwhile, we found that TMK4 did not obviously affect the TAA1 gene transcription and/or TAA1 proteins subcellular localization (Supplementary Fig. 11b, c). Therefore, we have comprehensive evidence that TMK4 interacts and phosphorylates TAA1 at the T101 site to switch off the TAA1 enzymatic activity.

TMK4 negatively regulates auxin biosynthesis through TAA1.

To confirm whether TMK4 is involved in regulating TAA1-dependent auxin biosynthesis, we tested the tmk4-1 mutant phenotype during root development. The tmk4-1 mutant showed an enlarged root meristem and longer root hairs (Supplementary Fig. 12a–f), similar to the phenotype of seedlings treated with auxin. This phenotype was confirmed by another allele, tmk4-2, and was complemented by genomic TMK4 (Supplementary Fig. 12a–f). To further confirm the function of TMK4 in auxin biosynthesis, we directly measured levels of free IAA in the tmk4-1 mutant. Indeed, we found an increased IAA abundance in the tmk4-1 mutant that could be rescued by genomic TMK4 (Supplementary Fig. 12g). In addition, both DR5-GUS and DII-Venus auxin markers showed that auxin concentration in tmk4 mutants was dramatically increased (Fig. 4a, b; Supplementary Fig. 13a, b), consistent with the measurement of free IAA levels.

By introducing the TAA1 mutation to the tmk4 mutant or blocking TAA1 enzymatic activity by L-Kyn chemical in the tmk4 mutant, we found that the increased auxin concentration in tmk4 was restored when using DR5-GUS or DII-Venus as an auxin marker (Fig. 4a, b; Supplementary Fig. 13a, b). The increased levels of free IAA in tmk4-1 was also mostly restored with the introduction of TAA1 mutations (Fig. 4c). Similarly, both the root meristem and the root hair phenotype in the tmk4-1 mutant were mostly blocked by either the TAA1 mutation or L-Kyn treatment (Fig. 4d, e; Supplementary Fig. 13c). These genetic data strongly indicate that TAA1-based auxin biosynthesis contributes most of the overproduced auxin in tmk4 and corroborate the biochemical data to support the negative regulation of auxin biosynthesis by TMK4 through TAA1.

TMK4 belongs to a transmembrane kinase family which is a key regulator in auxin signalling. Interestingly, we found that the phosphorylation level of TMK4 proteins increased in the auxin overproduction mutant yuc1-D59 as observed using a gel shift assay, indicating that auxin may activate TMK4 (Supplementary Fig. 14a). It has been reported that auxin controls its own biosynthesis to balance its local concentration.40,41 When we treated the Arabidopsis protoplast with exogenous auxin, we also found an increase in phosphorylation on TAA1 proteins at T101 residue, suggesting a negative feedback regulation of auxin biosynthesis (Supplementary Fig. 14b). The auxin effect on the T101 phosphorylation level of TAA1 proteins was partially blocked in the tmk4 mutant (Supplementary Fig. 14b), suggesting the TMK4-TAA1 regulation module might be one of the self-regulation mechanisms to control auxin biosynthesis in plants.

Discussion

In plants, TAA/YUC-mediated auxin biosynthesis pathway is a well-established pathway to produce the majority of free IAA.4,14,15 In depth studies previously revealed that local auxin biosynthesis plays an indispensable role for plants in response to both developmental and environmental cues, and suggesting that accurate regulation of auxin biosynthesis at multiple levels is required.6,8,12,18 The spatial-temporal regulation of auxin at the transcriptional level has been described as an important mechanism for auxin biosynthesis regulation, yet little is known about the non-transcriptional regulation of this pathway. Here, we establish a phosphorylation-based regulatory mechanism that determines TAA1 enzymatic activity which further controls auxin biosynthesis. To our knowledge, this is the first identified post-translational regulatory mechanism of auxin biosynthesis in plants.

In this work, we identified an in vivo phosphorylation site at T101 residue on TAA1 protein by mass spectrometric analysis and provided further evidence of the mechanistic switch of TAA1 enzymatic activities using several experimental approaches. This regulatory mechanism of auxin biosynthesis is essential for maintaining the well-organized root development in Arabidopsis (Fig. 1; Supplementary Fig. 4). Importantly, the T101 phosphorylation site is evolutionarily conserved and functions in the ancient plant Marchantia, indicating the significance of this novel regulatory mechanism on auxin biosynthesis (Fig. 2i–k; Supplementary Fig. 8c). The T101 phosphorylation site is located in the PLP binding pocket, which is required for TAA1 enzymatic activity. Furthermore, using functional analysis of both phosphomimic TAA1T101D and non-phosphor TAA1T101A mutations, we found that the phosphorylation at the T101 residue might obstruct its binding to the PLP cofactor. Both mutations affected TAA1 binding with the coenzyme PLP, yet TAA1T101A proteins had restored enzymatic activity when supplied with sufficient PLP while the TAA1T101D mutant could not restore enzymatic activity despite the presence of sufficient PLP levels (Fig. 1c, d; Supplementary Fig. 2), suggesting the phosphorylation is one of the key regulatory mechanisms of the TAA1 enzyme. This work also initiates the future directions to identify other regulatory mechanisms, including other phosphorylation sites of TAA1 that regulate auxin concentration in plants.

Next, we identified TMK4, a protein kinase that targets TAA1 protein at the T101 site, as a negative regulator of auxin biosynthesis (Fig. 3; Fig. 4; Supplementary Fig. 10). TMKs have been implicated as key regulators in auxin signalling.37,38 Here, we found that TMK4 partially participates in auxin regulation of T101 phosphorylation on TAA1 proteins (Supplementary Fig. 14), suggesting that auxin may facilitate a feedback regulation of its biosynthesis that balances the local auxin concentration that is essential for distinctive developmental outputs.

Interestingly, our data indicated that other kinases may also be involved in regulating the TAA1 phosphorylation since TAA1 was still slightly phosphorylated in the tmk4 mutant (Fig. 3e). It would be interesting to identify additional kinases that can target TAA1 or its homologs. Identifying such kinases would provide evidence for a more comprehensive signalling network in control of auxin biosynthesis for other developmental or environmental responses. We also found that free IAA level and root phenotype in tmk4 was not fully suppressed by wei8-3 single mutant (Fig. 4c; Supplementary Fig. 13c). This indicated that, besides TAA1, TMK4 must have other targets involved in auxin
Fig. 4 T MK4 negatively modulates local auxin biosynthesis through TAA1. a GUS staining of 5-day-old Col-0, tmk4-1, wei8-3, and tmk4-1;wei8-3 crossed with DR5-GUS marker. Scale bar, 50 μm. Quantification data and statistical analysis were shown in Supplementary Fig. 13a. b DII-Venus signal in the roots of Col-0 and tmk4-2. Seedlings were grown on medium with DMSO or 1.5 μM L-Kyn for 5 days. Scale bar, 75 μm. Quantification data and statistical analysis were shown in Supplementary Fig. 13b. c Free IAA quantification of Col-0, tmk4-1, wei8-3, and tmk4-1;wei8-3. Six individual biological repeats (n) were performed using 8-day-old seedlings. P values were shown as indicated; two-sided paired t-test. Centre line in box represents mean and whiskers show minimum to maximum. d Representative pictures of root apical meristem from 5-day-old seedlings. White arrows show meristem zone. Scale bar, 50 μm. e Quantification of root meristem size in d. n denotes the number of independent seedlings; one-way ANOVA with Tukey multiple comparisons test. Different letters represent significant difference between each other, P < 0.05. f Working model showing the phosphorylation-dependent mechanism of the on-off switch of TAA1 in plants. This central mechanism may regulate auxin levels in response to various upstream cues, which further modulates plant development and/or helps to deal with environmental changes.

Methods

Plant materials and growth conditions. All plant materials were grown in plant growth chambers (PERVICAL AR-66L3) using long-day condition (16 h light/8 h dark) at 23 °C, unless indicated otherwise. For Arabidopsis, Col-0 was used as wild-type plant. tmk4-1 (GABI_348E01) and tmk4-2 (SAIL_597_G10) in Col-0 background were used for phenotypic analyses. wei8-3 (SALK_127890)18, tar2-1 (SALK_0211258), and tar2-2 (SALK_137800) were used for genetic crossing and TAA1 function analyses. To generate pTMK4-TMK4-GFP transgenic plants, 3.3 kb TMK4 promoter and TMK4 genomic coding region were amplified by polymerase chain reaction (PCR) and inserted into a pDONR-Zeo vector, subsequently transferred into a pGWB504 destination vector by LR reaction and then transformed into Arabidopsis. To generate pTAA1-TAA1WT-GFP transgenic plants, 2 kb promoter region of TAA1 and TAA1 genomic coding region were amplified by PCR and inserted into a pDONR-Zeo vector, subsequently transferred into a pGWB504 destination vector and then transformed into Arabidopsis.

Concentration regulation. It would be also interesting to know how auxin signalling modules auxin biosynthesis through multiple targets.

Altogether, our work illustrates a phosphorylation-dependent switch of TAA1 enzymatic activity that is essential for the regulation of auxin biosynthesis in plants (Fig. 4f). Our observations offer an option for a mechanism of how local auxin biosynthesis is tightly controlled and subsequently regulates the dynamic auxin concentration gradients along with plant tissues under certain developmental or environmental stimulus. Our findings also provide a new understanding on how plants cope with their comprehensive developmental processes, in part, by modulating auxin biosynthesis.

Identification of phosphorylation sites in vivo by LCMS/MS. Seven-day-old pTAA1-TAA1 GFP transgenic Arabidopsis seedlings grown in long-day conditions were collected and treated with 50 μM cantharidin (Sigma C7632) for 6 h. Total proteins were extracted with extraction buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 5 mM EDTA, 5% glycerol, 1% Triton-X-100 with protease inhibitor, phosphatase inhibitor, and PMSF). GFP-Trap agarose beads (GFP-Trap®_A, gta-20,
ChromoTek) were used to immunoprecipitate TAA1-GFP proteins. The proteins were separated on a sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) gel (throughout 30 min staining, and followed by liquid chromatography with tandem MS (LC/MS) analysis. For comparing T101 phosphorylation levels in Col-0 and tmk-1, PTA1-TAA1;Col-0, and PTA1-TAA1;tmk-1 transgenic plants were used for immunoprecipitating TAA1 proteins. The amount of TAA1-GFP proteins was standardized by two non-phosphorylated protein bands to detect auxin effects on T101 phosphorylation level. 355-ATA1-GFP plasmids were transformed into Col-0 and tmk-1 protoplasts incubated for 8 h. The protoplasts were collected after treated with EtOH (as mock) or 500 nM IAA for 10 mins. TAA1-GFP proteins were immunoprecipitated from the protoplasts further used for quantitative mass spectrometric analysis.

Protein peptides were reconstituted in 4 μL of 0.1% formic acid (FA) and then were injected onto a NanoAcquity UPLC system (Waters, USA) equipped with a C18 capillary trapping column (20 mm × 180 μm, Waters PN:186006527) and a C18 analytical column (75 μm × 150 mm, Waters PN:186003543), and separated over a 70-min gradient at 350 nL/min. This HPLC system was online with an Orbitrap Fusion MS (Thermo Fisher, USA) using high-energy collision-induced dissociation mode. MS survey scan was performed at a resolution of 120,000 over the m/z range of 300–1,800, and MS/MS was selected by data-dependent scanning on the ten most intense ions (automatic gain control (AGC) 1E5, maximum injection time 100 ms). Dynamic exclusion was set to a period of 60 s. MS/MS data were searched against the protein sequences of TAA1 using Mascot Daemon 2.5 (Matrix Science UK). The search parameters were restricted to tryptic peptides at a maximum of two missed cleavages. Cysteine carbamidomethylation was designated as a fixed modification. Oxidation of methionine and serine, threonine or tyrosine phosphorylation were considered as variable modifications. Mass tolerances were set to 10 ppm for MS ions and 0.5 Da for MS/MS fragment ions. Peptide assignments were filtered by an ion score cutoff of 15.

To monitor phosphorylation changes on TAA1 proteins, the parallel reaction monitoring (PRM) was applied for phospho-peptides quantification. Peptide samples were dissolved in 9 μL of 0.1% FA into an EASY-nLC 1200 HPLC system (Thermo Fisher, USA). Eluent was introduced into the mass spectrometer, using an in-house C18 capillary column packed with 2.2 μm C18 particles (Polymeric; 75 μm inner diameter and 12 cm of bed length). The mobile phase consisted of 0.1% FA in water (buffer A) and 0.1% FA in 80% (v/v) acetonitrile (buffer B). The gradient was set as 5–25% buffer B for 40 mins and 25–45% for 10 mins at 300 nL/min. The sample was acquired on an Orbitrap Fusion MS (Thermo Fisher, USA). Each sample was analysed under PRM with an isolation width of ±1.6 Da. In all experiments, a full mass spectrum at 120,000 resolutions (AGC target 2E5, 50 ms maximum injection time) was followed by PRM scans at 30,000 resolutions (AGC target 1E5, 200 ms maximum injection time). PRM data were manually curated within Skyline (version 4.1.0.18163)44.

A total of 600 μL samples, 2.5 μg TAA1-His proteins were used for each reaction then the absorbance was measured by Biotek cell imaging multimode reader (BioTek Instruments, Inc. Winooski, VT) at different time points.

IPA measurement via Salkowski reagent. To test TAA1 enzymatic activity in vivo, Salkowski reagent was used to quantify IPA production. This assay was performed as previous reported45,46 with slight modification. Briefly, –50 mg root tissues were ground up in liquid nitrogen, added into the same volume of extraction buffer (100 mM HEPES, pH 7.5, 250 mM sorbitol, 5 mM β-mercaptoethanol, 0.5% (v/v) Triton X-100, and 0.1% (w/v) PMSF), mixed well and kept on ice for 10 min. The extraction mixture was centrifuged at 10,000 × g for 10 mins at 4 °C. Fifty microliter total proteins were mixed with reaction buffer (100 mM phosphate buffer, pH 8.0, 10 mM L-tryptophan, 3 mM 2-oxoglutarate, 10 mM PPL, prewarmed at 37 °C for 3 mins). Proteins mixed with reaction buffer without adding a-ketoglutaric acid were used as a background control. The reactions were performed at 37 °C for 30 min at room temperature. The absorbance at 530 nm absorption was measured by microplate reader (Thermo Variaskan Flash). Concentration of total protein was determined using Coomassie brilliant blue G250 dye by detecting the absorbance at 595 nm. The relative enzymatic activity was shown as A1/200A0. The absorbance was also used to test the enzymatic activity of E. coli-purified proteins. TAA1 CDS was amplified and inserted into PET14 vector and expressed in E. coli. Proteins were further purified by Ni-NTA resin (QIAGEN cat. no. 30210). For the recombinant proteins, 2.5 μg TAA1-His proteins were used for each reaction then the absorbance was measured by Biotek cell imaging multimode reader (BioTek Instruments, Inc. Winooski, VT) at different time points.

PLP detection by measuring absorbance at 388 nm. PLP was reported to show maximum absorbance ~388 nm46,47. TAA1WT-His, TAA1T101A, His, TAA1T101D, His, and TAA1K217A-His proteins were purified from E. coli. A total of 600 μL proteins (2 μg/μL) were denatured at 95°C for 5 mins to release PLP from proteins. The supernatant containing PLP was obtained by centrifugation and used for reading absorbance at 388 nm. Different concentrations of PLP (Sigma, P9255) were set as positive controls.

Yeast two-hybrid assay. The C-terminals of TMK family proteins named TMK1C, TMK2C, TMK3C, and TMK4C48 were cloned and inserted into the pGBK7 vector as bait. The full-length CDS of D6PK, PINOID, and the kinase domains of BAK1, RAK1, and RAK2 were cloned and inserted into the pGWB7 vector. TAA1 CDS was cloned, inserted into PDON-Zreo then inserted into the pGWB vector by LR reaction as prey. The Golden Yeast strain was used for yeast two-hybrid assay. Yeast transformation was done by the transformation kit (Fro-zen-EZ Yeast Transformation II Kit®, T2001, ZYMO RESEARCH), and the interaction was tested on -Trp-Leu and -Trp-Leu-AdH-His medium.

Commmunoprecipitation assay in protoplast. The coding region of TMK4 and TAA1 were amplified, and then inserted into the transient expression vectors 35SS-HBT-HA-NOS and 355-HBT-Flag-NOS, respectively. TMK4-HA and TAA1-Flag were coexpressed in Col-0 protoplasts, while single expression was used as a negative control. Total proteins were extracted from the protoplasts with the extraction buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 5 mM EDTA, 5% glycerol, 1% TritonX-100 with protease inhibitor, phosphatase inhibitor, and PMSF), then incubated with anti-Flag beads (Sigma cat. #A2220) at 4°C for 4 h followed by two times of washing (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 5 mM EDTA, 5% glycerol, and 0.01% TritonX-100). Proteins were separated on SDS-PAGE gels and detected by anti-HH (HA-7, H6533, Sigma, 1:2000 dilution) or anti-Flag (M2008L, Abmart, 1:2000 dilution) antibodies via western blots. 355-TAA1-Flag and 355-TAR1-GFP were generated to detect interaction between TAA1 and TAR using similar assay. TAR1-GFP were detected by an anti-GFP antibody (HT801, TransGen Biotech, 1:2000 dilution) via western blot.

In vitro pull-down assay. TAA1-His and His-Sumo proteins were expressed in E. coli and captured by Ni-NTA resin. TMK4 KD (587aa-928aa) sequence was amplified and inserted into pGEX4T-2 vector. GST-TMK4 KD proteins were isolated from E. coli by Glutathione agarose (sigma G4510) and incubated with
TAA1-His or His-Sumo beads in binding buffer (20 μM HEPES, pH 7.5, 40 μM KC1, 1 mM EDTA, 1% glycerol, 0.1% TritonX-100 with PMSF) for 2 h at 4 °C. The beads were washed three times by washing buffer (20 μM HEPES, pH 7.5, 40 μM KC1, 1 mM EDTA, and 0.055% TritonX-100). GST-TMK4 KD proteins were detected by GST antibody (M20007, Abmart, 1:2000 dilution) via western blots, and His-TAA1 and His-Sumo proteins were detected by Coomassie blue staining.

In vitro kinase assay. TMK4 kinase domain (578aa–928aa) was inserted into His-sumo vector. His-sumo-TMK-KD proteins and TAA1-His proteins were mixed in the reaction buffer (50 mM HEPES, 10 mM MgCl2, 10 mM MnCl2, 1 mM DTT with 50 μM γ32P ATP, and 50 μM ATP). After incubation at 25 °C for 1 h, the reaction was stopped by adding 5x SDS loading buffer. Radioactive signals from TAA1-His were detected using the Typhoon imaging system (GE Healthcare TYPHOON FLA 9500).

To detect phosphorylation sites of TAA1 targeted by TMK4 in vitro, similar assay was performed using TAA1-His or GST-TAA1 as substrate. His-sumo-TMK4KD as kinase with 50 μM cold ATP adding in reaction system. After phosphorylation reaction, the substrates were sent for mass spectrometric analysis to identify phosphorylation sites of TAA1 in vitro. TAA1 proteins without kinases adding were used as a negative control group.

GUS staining and quantification. Five-day-old seedlings were fixed in 90% acetone in ice for 20 min, washed by the staining buffer (100 mM NaH2PO4 pH 7.0, 10 mM EDTA, 0.5 mM K3[Fe(CN)6]·3H2O, 0.5 mM K4[Fe(CN)6], 0.1% TritonX-100), and incubated in the staining buffer with 0.5 mg/mL 5-bromo-4-chloro-3-indolyl-b-d-glucuronic acid (x-Gluc) at 37 °C for 3 h. The reactions were stopped by adding 75% ethanol. Images were taken by Leica DM6B Microscope with roots immersed in clearing buffer (chloral hydrate: water: glycerol = 4:100:96) and incubated in the staining buffer with 0.5 mg/mL 5-bromo-4-chloro-3-indolyl-b-d-glucuronic acid (x-Gluc) at 37 °C for 3 h. The reactions were stopped by adding 75% ethanol. Images were taken by Leica DM6B Microscope with roots immersed in clearing buffer (chloral hydrate: water: glycerol = 4:100:96). The fluorescence absorbance (excitation: 365 nm, emission: 455 ± 5 nm) via microplate reader (Thermo Variaskan Flash). Relative GUS activity was standardized by total protein amount measured by Coomassie blue G250 dye.

Gel shift assay. Five-day-old seedlings were collected and ground up in liquid nitrogen. Proteins were extracted by 2x SDS loading buffer (100 μM Tris-HCl pH 6.8, 4% SDS, 12% glycerol, 0.02% bromophenol blue, and 2% β-mercaptoethanol) followed by 95 °C denaturation. Proteins were separated on 6% SDS–PAGE gel containing 50 μM phos-tag (Phos-tag TM Acrylamide AAL-107, WAKO) and 100 μM MnCl2 and detected by anti-TMK4 antibody (135-360aa, generated by Abclonal Biotechnology, 1:2000 dilution).

Structure analysis. Information of the TAA1 protein structure (PDB code: 3bwn) was obtained from the PDB database (http://www.rcsb.org)9. Pymol (Version 0.99rc6) software was used for structure analysis and image capture.

Phylogenetic analysis of TAA1 family. Information of protein sequences was obtained from the NCBI database (https://www.ncbi.nlm.nih.gov/) and the phytozone database (http://phytozone.jgi.doe.gov/pz/portal.html). Clustal Omega was used for sequence alignment (https://www.ebi.ac.uk/Tools/msa/clustalo/).

Statistical analysis. All box plots and bar graphs were generated by GraphPad Prism 7 software. To compare multiple groups, analysis of variance (ANOVA) one-way test was applied using the GraphPad Prism 7 software. All P values which is <0.0001 were shown as P < 0.0001, P > 0.05 indicates no significant difference. Lines in dot plots indicate median with s.d.

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

Data availability
The data supporting the findings in this study are available and described within the paper and its supplemental information. Source Data (gels and graphs) for Figs. 1–4 and Supplementary Figs. 2–6, 8, 10–14 are provided as a separate Source Data file.

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Author contributions

T.X. and Q.W. initiated this project and designed the experiments. Q.W. and G.Q. carried out most of the experiments. M.C. helped with the in vitro kinase assay. R.C. helped with phenotypic analysis. Y.H. assisted with TMK4 phosphorylation assay, L.Y. and Z.Z. generated MpTAA constructs, Y.Y. helped to make TMK4 antibodies, and Y.G. assisted with protein purification. W.X. helped with the computer analysis of the TAA1 structure, and A.T. helped with mass spectrometric analysis. T.X. and Q.W. wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Correspondence and requests for materials should be addressed to T.X.

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